



Center For The Evaluation Of Risks To Human Reproduction

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**NTP-CERHR REPORT on the
REPRODUCTIVE and DEVELOPMENTAL
TOXICITY of BISPHENOL A**

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1 **PREFACE**

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3 To be added after the Expert Panel meeting

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5 Reports can be obtained from the website (<http://cerhr.niehs.nih.gov>) or from:

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19 This report is prepared according to the Guidelines for CERHR Panel Members established by
 20 NTP/NIEHS. The guidelines are available from the CERHR web site (<http://cerhr.niehs.nih.gov/>). The
 21 format for this report follows that of CERHR Expert Panel Reports including synopses of studies
 22 reviewed, and an evaluation of the Strengths/Weaknesses and Utility (Adequacy) of the study for a
 23 CERHR evaluation. Statements and conclusions made under Strengths/Weaknesses and Utility
 24 evaluations are those of the CERHR Scientists and are prepared according to the NTP/NIEHS guidelines.
 25 In addition, the report includes comments or notes limitations of the study in the synopses. Bold, square
 26 brackets are used to enclose such statements. As discussed in the guidelines, square brackets are used to
 27 enclose key items of information not provided in a publication, limitations noted in the study, conclusions
 28 that differ from authors, and conversions or analyses of data conducted by CERHR. **The findings and**
 29 **conclusions of this report are those of the Expert Panel and should not be construed to represent**
 30 **the views of the National Toxicology Program.**
 31

1 ABBREVIATIONS

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µg	microgram(s)
µM	micromolar
ADA	American Dental Association
ANCOVA	analysis of covariance
ANOVA	analysis of variance
atm	atmosphere
AUC	area under the time-concentration curve
AUC _{BPA}	area under the time-concentration curve for bisphenol A
BMD _{1 SD}	benchmark dose, 1 control standard deviation
BMD ₁₀	benchmark dose, 10% effect level
BMDL	benchmark dose 95 th percentile lower confidence limit
BrdU	bromodeoxyuridine
bw	body weight
cAMP	cyclic adenosine monophosphate
CAS RN	Chemical Abstracts Service registry number
CFR	Code of Federal Regulations
CHO	Chinese hamster ovary
CI	confidence interval
C _{max}	maximum plasma concentration
CNS	central nervous system
cytochrome P	CYP
DMSO	dimethylsulfoxide
DNA	deoxyribonucleic acid
EC ₁₀	10% effective concentration
EC ₅₀	median effective concentration
ELISA	enzyme-linked immunosorbent assay
eq	equivalent(s)
ER	estrogen receptor
EROD	7-ethoxyresorufin O-deethylase
ERK	extracellular signal-regulated kinase
FDA	Food and Drug Administration
fM	femtomolar
FSH	follicle stimulating hormone
GABA	γ-aminobutyric acid
GC/MS	gas chromatography/mass spectrometry
GD	gestation day(s)
GLP	Good Laboratory Practices
GST	glutathione-S-transferase
hCG	human chorionic gonadotropin
HPLC	high performance liquid chromatography
hprt	hypoxanthine phosphoribosyl transferase
IARC	International Agency for Research on Cancer
IC ₅₀	median inhibitory concentration
IgG	immunoglobulin G
ip	intraperitoneal(ly)
IU	international unit
im	intramuscular
iv	intravenous(ly)

k_{el}	elimination constant
kg	kilogram(s)
K_m	rate constant
L	liter(s)
LC	liquid chromatography
LD_{50}	median lethal dose
LH	luteinizing hormone
LOD	limits of detection
LOQ	limits of quantification
m	meter(s)
M	molar
MAPK	mitogen activated protein kinase
mCi	millicurie(s)
MDL	minimum detection limit
mg	milligram(s)
mL	milliliter(s)
mM	millimolar
mol	mole(s)
mRNA	messenger ribonucleic acid
MS	mass spectrometry
MURST	Italian Ministry for Universities and Scientific and Technological Research
ng	nanogram(s)
NADPH	reduced nicotinamide adenine dinucleotide phosphate
NCTR	National Center for Toxicological Research
ND	not detected
NHANES	National Health and Nutrition Examination Survey
NICHHD	National Institute of Child Health and Human Development
NIEHS	National Institute of Environmental Health Sciences
NIH	National Institutes of Health
NIOSH	National Institute of Occupational Safety and Health
NOAEL	no observed adverse effect level
NOEL	no observed effect level
nM	nanomolar
NTP	National Toxicology Program
PBPK	physiologically based pharmacokinetic model
PBS	phosphate buffered saline
PCR	polymerase chain reaction
pM	picomolar
PND	postnatal day(s)
ppb	parts per billion
PCNA	proliferating cell nuclear antigen
pg	picogram
ppm	parts per million
ppt	parts per trillion
RIA	radioimmunoassay
RNA	ribonucleic acid
RT	reverse transcriptase
PVC	polyvinylchloride
sc	subcutaneous(ly)
SD	standard deviation
SDN-POA	sexually dimorphic nucleus in the preoptic area of the hypothalamus

SDS-PAGE	sodium dodecyl sulfate polyacrylamide electrophoresis
SEM	standard error of the mean
sst ₃	somatostatin subtype 3
T _{1/2}	half-life
tk	thymidine kinase
T _{max}	time to maximum plasma concentration
TUNEL	terminal deoxynucleotidal transferase-mediated dUTP nick-end labeling
TWA	time-weighted average
UDPGT	uridine diphosphate glucuronosyltransferase
US	United States
USEPA	United States Environmental Protection Agency
V _{max}	maximum velocity
WEEL	workplace environmental exposure level

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1.0 CHEMISTRY, USE, AND HUMAN EXPOSURE

1.1 Chemistry

Section 1 is initially based on secondary review sources. Primary study reports are addressed by the Expert Panel if they contain information that is highly relevant for determining the effect of exposure on developmental or reproductive toxicity or if the studies were released subsequent to the reviews.

1.1.1 Nomenclature

The CAS RN for bisphenol A is 80-05-7. Synonyms for bisphenol A listed in Chem IDplus (1) include: 2-(4,4'-Dihydroxydiphenyl)propane; 2,2-Bis(4-hydroxyphenyl)propane; 2,2-Bis(hydroxyphenyl)propane; 2,2-Bis(p-hydroxyphenyl)propane; 2,2-Bis-4'-hydroxyfenylpropan [Czech]; 2,2-Di(4-hydroxyphenyl)propane; 2,2-Di(4-phenylol)propane; 4,4'-(1-Methylethylidene)bisphenol; 4,4'-Bisphenol A; 4,4'-Dihydroxydiphenyl-2,2-propane; 4,4'-Dihydroxydiphenyldimethylmethane; 4,4'-Dihydroxydiphenylpropane; 4,4'-Isopropylidene diphenol; 4,4'-Isopropylidenebisphenol; 4,4'-Isopropylidene diphenol; Biphenol A; Bis(4-hydroxyphenyl) dimethylmethane; Bis(4-hydroxyphenyl)dimethylmethane; Bis(4-hydroxyphenyl)propane; Bisferol A [Czech]; Bisphenol. Bisphenol A; DIAN; Diano; Dimethyl bis(p-hydroxyphenyl)methane; Dimethylbis(p-hydroxyphenyl)methane; Dimethylmethylene-p,p'-diphenol; Diphenylolpropane; Ipognox 88; Isopropylidenebis(4-hydroxybenzene); Parabis A, Phenol; (1-methylethylidene)bis-, Phenol; 4,4'-(1-methylethylidene)bis-, Phenol; 4,4'-dimethylmethylenedi-, Phenol; 4,4'-isopropylidenedi-, Pluracol 245, Propane; 2,2-bis(p-hydroxyphenyl)-; Rikabanol; Ucar bisphenol A; Ucar bisphenol HP; beta,beta'-Bis(p-hydroxyphenyl)propane; beta-Di-p-hydroxyphenylpropane; p,p'-Bisphenol A; p,p'-Dihydroxydiphenyldimethylmethane; p,p'-Dihydroxydiphenylpropane; p,p'-Isopropylidenebisphenol; and p,p'-Isopropylidenediphenol.

1.1.2 Formulae and molecular mass

Bisphenol A has a molecular mass of 228.29 and a molecular formula of $C_{15}H_{16}O_2$ (2). The structure for bisphenol A is shown in Figure 1.

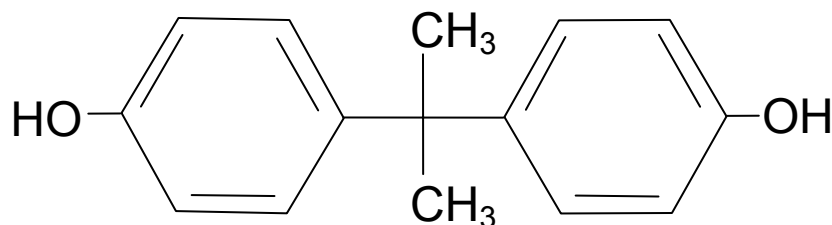


Figure 1. Structure for Bisphenol A.

1.1.3 Chemical and physical properties

Bisphenol A is a colorless white solid with a mild phenolic odor (2). Physicochemical properties are listed in Table 1.

1 **Table 1. Physicochemical Properties of Bisphenol A**

Property	Value
Odor threshold	No data found
Boiling point	220°C at 4 mm Hg; 398 C at 760 mm Hg
Melting point	150–157°C
Specific gravity	1.060–1.195 g/mL at 20–25°C
Solubility in water	120–300 mg/L at 20–25°C
Vapor pressure	8.7×10^{-10} – 3.96×10^{-7} mm Hg at 20–25°C
Stability/reactivity	No data found
Log K_{ow}	2.20–3.82
Henry constant	1.0×10^{-10} atm·m ³ /mol

From Staples et al. (3).

2
3 *1.1.4 Technical products and impurities*
4 Purity of bisphenol A was reported at 99–99.8%, and common impurities observed were phenol and ortho
5 and para isomers of bisphenol A (reviewed in (2)).
6

7 No information on trade names for bisphenol A was located.
8

9 **1.2 Use and Human Exposure**

10 *1.2.1 Production information*

11 Bisphenol A is manufactured by condensation of phenol and acetone catalyzed by an acid or alkaline
12 compound (2).
13

14
15 In 1998, members of the Society of the Plastics Industry Bisphenol A Task Group [**assumed**
16 **manufacturers of bisphenol A**] included Aristech Chemical Corporation, Bayer Corporation, Dow
17 Chemical Company, and Shell Chemical Company (3). Additional current or past manufacturers of
18 bisphenol A in the US include General Chemical, Union Carbide, and ACC Holdings (4). According to
19 the Society of the Plastics Industry, there are currently 6 bisphenol A and 4 polycarbonate plants in the
20 US (5); 3 of the 4 polycarbonate plants are located within bisphenol A plants. In 2000, there were 13
21 epoxy plants in the US, but was not clear if all of the plants manufactured bisphenol A-containing epoxy
22 resins.
23

24 In mid 2004, US bisphenol A production volume was reported at 1024 thousand metric tons [**~2.3 billion**
25 **pounds**] (6). A production volume of 7.26 billion g [**16 million pounds**] was reported for bisphenol A in
26 1991 (reviewed in (4)). US bisphenol A consumption was reported at 856,000 metric tons [**~1.9 billion**
27 **pounds**] in 2003 (6); 2003 consumption patterns included 619,000 metric tons [**~1.4 billion pounds**] used
28 in polycarbonate resins, 184,000 metric tons [**~406 million pounds**] used in epoxy resins, and 53,000
29 metric tons [**~117 million pounds**] used in other applications.
30

31 *1.2.2 Use*

32 In 1999 and 2003, it was reported that most bisphenol A produced in the US was used in the manufacture
33 of polycarbonate and epoxy resins and other products (reviewed in (3, 6)). Polycarbonate plastics may be
34 used in the manufacture of compact discs, “solid and multi wall sheet in glazing applications and film,”
35 food containers (e.g., milk, water, and infant bottles), and medical devices (reviewed in (2)).
36 Polycarbonate blends have been used to manufacture injected molded parts utilized in alarms, mobile
37 phone housings, coil cores, displays, computer parts, household electrical equipment, lamp fittings, and
38 power plugs. Automotive and related uses for polycarbonate blends include light reflectors and coverings,
39 bumpers, radiator and ventilation grills, safety glazing, inside lights, and motorcycle shields and helmets.

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1 Epoxy resins are used in protective coatings, structural composites, electrical laminates, electrical
2 applications, and adhesives. The European Union (2) reported that smaller volumes of bisphenol A are
3 used in production of phenoplast, phenolic, and unsaturated polyester resins, epoxy can coatings,
4 polyvinylchloride (PVC) plastic, alkoxyated bisphenol A, thermal paper, and polyols/polyurethane.
5 Other uses reported for products manufactured from bisphenol A included protective window glazing,
6 building materials, optical lenses, and development of dyes (reviewed in (3)). A search of the National
7 Library of Medicine Household Products Database (7) revealed that bisphenol A-containing coatings,
8 adhesives, and putties are available to the general public for use in automobiles, home maintenance and
9 repair, and hobbies.

10
11 Some polymers manufactured with bisphenol A are Food and Drug Administration (FDA) approved for
12 use in direct and indirect food additives and in dental materials, as reported in the Code of Federal
13 Regulations (CFR) (8). In the CFR, bisphenol A is often referred to as 4,4'-isopropylidenediphenol.
14 Polymers manufactured with bisphenol A are FDA-approved for use as anoxomers (21CFR172.105) and
15 in coatings (21CFR175.300; 21CFR175.320; 21CFR175.380), adhesives (21CFR175.105), single and
16 repeated food contact surfaces (21CFR177.1555; 21CFR177.1595), and tooth shade resin materials
17 (21CFR872.3690).

18
19 The European Union (2) noted that resins, polycarbonate plastics, and other products manufactured from
20 bisphenol A can contain trace amounts of residual monomer and additional monomer may be generated
21 during breakdown of polymer. The Society of the Plastics Industry reports that residual bisphenol A
22 levels in polycarbonate plastics and epoxy resins are generally <50 ppm (5). Polymer hydrolysis can
23 occur at elevated temperature or extreme pH. An example of potential human exposure is migration of
24 bisphenol A from a food container into the food. Exposure to bisphenol A through food is discussed in
25 detail in Section 1.2.3.2.

26 27 *1.2.3 Occurrence*

28 29 *1.2.3.1 Environmental fate and bisphenol A levels in environment*

30 Bisphenol A may be present in the environment as a result of direct releases from manufacturing or
31 processing facilities, fugitive emission during processing and handling, or release of unreacted monomer
32 from products (2). According to the Toxics Release Inventory database, total environmental release of
33 bisphenol A in 2004 was 181,768 pounds, with releases of 132,256 pounds to air, 3533 pounds to water,
34 172 pounds to underground injection, and 45,807 pounds to land (9).

35
36 Bisphenol A released to the atmosphere is likely degraded by hydroxy radicals (2). Half-life for the
37 reaction between bisphenol A and hydroxy radicals was estimated at 0.2 days. It was also noted that
38 photolysis and photodegradation of bisphenol A in the atmosphere is possible and photooxidation half-
39 lives of 0.74–7.4 hours were estimated (reviewed in (2, 3)). The European Union (2) noted that because of
40 its low volatility and relatively short half-life in the atmosphere, bisphenol A is not likely to enter the
41 atmosphere in large amounts. Removal by precipitation and occurrence in rain water were thought likely
42 to be negligible. Because of its short half-life in the atmosphere, bisphenol A is unlikely to be transported
43 far from emission points.

44
45 Based on vapor pressure and Henry constant (Table 1), the European Union (2) and Staples et al. (3)
46 concluded that bisphenol A is of low volatility and not likely to be removed from water through
47 volatilization. Both groups concluded that hydrolysis of bisphenol A in water is unlikely. However, there
48 was disagreement on potential for photooxidation of bisphenol A in water. Based on physical and
49 chemical properties, the European Union concluded that photolysis of bisphenol A in water is unlikely.
50 Staples et al. noted that bisphenol A is able to absorb ultraviolet light, especially in a basic solution.
51 Therefore, it was concluded that photolysis from surface water is possible, depending on conditions such

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1 as pH, turbidity, turbulence, and sunlight. Photooxidation half-life of bisphenol A in water was estimated
2 at 66 hours to 160 days (reviewed in (3)). Rapid biodegradation of bisphenol A from water was reported
3 in the majority of studies reviewed by the European Union (2) and Staples et al. (3). A biodegradation
4 half-life of 2.5–4 days was reported in a study measuring bisphenol A levels in surface waters near the
5 receiving stream of a bisphenol A manufacturer (reviewed in (3)).

6
7 When the Staples et. al. (3) review was published, soil sorption constants had not been measured but were
8 estimated at 314–1524. Based on such data, the European Union (2) and Staples et al. (3) concluded that
9 bisphenol A adsorption to soils or sediments would be “modest” or “moderate.” Based on data for
10 degradation of bisphenol A in water, the European Union (2) predicted that bisphenol A would be
11 degraded in soil and estimated a half-life of 30 days for degradation of bisphenol A in soil. Subsequent to
12 the Staples et al. and European Union reviews, a study examining fate of ¹⁴C-bisphenol A in soils through
13 laboratory soil degradation and batch adsorption tests was released by Fent et al. (10). In that study, ¹⁴C-
14 bisphenol A was rapidly dissipated and not detectable in 4 different soil types within 3 days. Soil
15 distribution coefficients were determined at 636–931, and based on those values, the study authors
16 concluded that bisphenol A has low mobility in soil. The study authors concluded that bisphenol A is not
17 expected to be stable, mobile, or bioavailable from soils.

18
19 In studies reviewed by the European Union (2) and Staples et al. (3), bioconcentration factors for fish
20 were measured at 3.5–68 and were found to be lower than values estimated from the K_{ow} . Both groups
21 concluded that potential for bioconcentration of bisphenol A is low in fish. Higher bioconcentration
22 factors (134–144) were determined for clams (reviewed in (2)).

23
24 Two studies examining aggregate exposures in preschool age children in the US used gas
25 chromatography/mass spectrometry (GC/MS) methods to measure bisphenol A levels in environmental
26 media (11, 12). In the first study (11), bisphenol A levels were measured in air outside 2 day care centers
27 and the homes of 9 children. Bisphenol A was detected in 9 of 13 outdoor air samples at <0.100–4.72
28 ng/m³. In indoor air from day care centers and homes, bisphenol A was detected in 12 of 13 samples at
29 <0.100–29 ng/m³. At those same locations, bisphenol A was detected in all of 13 samples of floor dust at
30 0.567–3.26 ppm (µg/g) and play area soils at 0.004–0.014 ppm (µg/g). In the second study (12), bisphenol
31 A levels were measured inside and outside at least 222 homes and 29 daycare centers. Bisphenol A was
32 detected in 31–44% of outdoor air samples from each location; levels ranged from <LOD (0.9) to 51.5
33 ng/m³. Forty-five to 73% of indoor air samples contained detectable levels of bisphenol A; levels were
34 reported at <LOD (0.9)–193 ng/m³. Bisphenol A was detected in 25–70% of dust samples; levels were
35 reported at <LOD (20)–707 ng/g.

36
37 A second US study used a GC/MS method to measure bisphenol A levels in dust from 1 office building
38 and 3 homes and in air from an office building and 1 home (13). Bisphenol A was detected in 3 of 6 dust
39 samples (reporting limit > 0.01 µg/extract) at concentrations of 0.25–0.48 µg/g dust. In indoor air samples
40 collected from offices and residences, bisphenol A was detected in 3 of 6 samples (detection limit 0.0038
41 µg/extract) at concentrations of 0.002–0.003 µg/m³. In another study using a GC/MS technique, bisphenol
42 A levels in indoor air and dust from 120 US homes were below reporting limits (0.018 µg/m³ for indoor
43 air and 0.2 µg/g for dust) (14).

44
45 Limited information is available for bisphenol A levels in US water. In 1996 and/or 1997, mean bisphenol
46 A levels were reported at 4–8 µg/L in surface water samples near a bisphenol A production site but
47 bisphenol A was not detected (<1 µg/L) in surface water near 6 of 7 bisphenol A production sites in the
48 US (15). Bisphenol A was detected at a median concentration of 0.14 µg/L and a maximum concentration
49 of 12 µg/L in 41.2% of 85 samples collected from US streams in 1999 and 2000 (16). In 2001 and 2002,
50 bisphenol A was not detected (< 0.001 µg/L) in effluent from a wastewater treatment plant in Louisiana,
51 and concentrations were not quantifiable [**quantification limit not defined**] in samples collected from

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1 surface waters near Louisiana and in drinking water at various stages of treatment at plants in Louisiana
2 or Ontario, Canada (17). In water samples collected in Europe and Japan from the 1970s through 1989,
3 bisphenol A levels were ≤ 1.9 $\mu\text{g/L}$ and in most cases were ≤ 0.12 $\mu\text{g/L}$ (reviewed in (2)). Bisphenol A was
4 not detected in drinking water collected from an unspecified location (reviewed in (2)).

6 *1.2.3.2 Potential exposures from food*

7 The European Union (2) noted that the highest potential for human exposure to bisphenol A is through
8 products that directly contact food. Examples of food contact materials that can contain bisphenol A
9 include food and beverage containers with internal epoxy resin coatings and polycarbonate tableware and
10 bottles, such as those used to feed infants.

11
12 Studies have measured migration of bisphenol A from polycarbonate infant bottles or containers into
13 foods or food simulants. Results of those studies are summarized in Table 2. Analyses for bisphenol A
14 were conducted by GC/MS or high performance liquid chromatography (HPLC). The European Union (2)
15 group noted that in many cases bisphenol A levels were below the detection limit in food simulants.
16 When bisphenol A was detected, levels were typically ≤ 50 $\mu\text{g/L}$ in simulants exposed to infant bottles and
17 ≤ 5 $\mu\text{g/kg}$ in simulants exposed to polycarbonate tableware. An exception is 1 study that reported
18 bisphenol A levels at up to ~ 192 $\mu\text{g/L}$ in a 10% ethanol food simulant and 654 $\mu\text{g/L}$ in a corn oil simulant
19 (18). In the study, cut pieces of bottles were incubated, and the study authors acknowledged that
20 bisphenol A could have migrated from the cut edges. One study conducted with actual infant food
21 (formula and fruit juice) reported no detectable bisphenol A (19). Some studies examining the effects of
22 repeated use of polycarbonate items noted increased leaching of bisphenol A with repeated use (20-22). It
23 was suggested that the increase in bisphenol A migration was caused by damage to the polymer during
24 use. Results from other reports suggested that leaching of bisphenol A decreased with repeated use, and it
25 was speculated that available bisphenol A was present at the surface of the product and therefore removed
26 by washing ((23) and Kawamura et al. (1998), reviewed by the European Union (2) and Haighton et al.
27 (24)). One study (Kawamura et al. (1998) demonstrated higher levels of bisphenol A in simulants exposed
28 to products that had been recalled because of unacceptable residual levels of bisphenol A and other
29 compounds. The study by Biles et al. (23) demonstrated that infant bottles exposed to 50 or 95% ethanol
30 at 65°C for 240 hours leached bisphenol A at levels exceeding residual monomer concentrations, and it
31 was suggested that hydrolysis of the polymer had occurred.

1 **Table 2. Examination of Bisphenol A in Polycarbonate Food Contact Surfaces**

Sample (Location)	Procedure	Bisphenol A level	Reference
Commercially available infant bottles containing residual bisphenol A levels of 7–46 ppm (US)	Common use: Bottles were boiled for 5 minutes, filled with water or 10% ethanol, and stored at room temperature for up to 72 hours. Worst case use: Bottles were boiled for 5 minutes, filled with water or 10% ethanol, heated to 100°C for 0.5 hour, cooled to room temperature, and refrigerated for 72 hours.	Not detected (ND; < 5 ppb [$\mu\text{g/L}$]; corresponding to a food level of 1.7 ppb) following either procedure.	FDA (25)
21 new and 12 used (1–2-year-old) infant bottles (UK)	Bottles were pre-washed, steam sterilized, filled with boiling water or 3% glacial acetic acid, refrigerated at 1–5°C for 24 hours, and heated to 40°C prior to sampling.	ND (< 10 $\mu\text{g/L}$) [ppb] in new bottles; ND (< 10) to 50 $\mu\text{g/L}$ in used bottles exposed to either simulant.	Earls et al. (20)
Infant bottles with residual bisphenol A levels of 26 mg/kg [number tested not indicated]. (UK)	Bottles were sterilized with hypochlorite, in dishwasher, or by steam; filled with infant formula, fruit juice, or distilled water; microwaved for 30 seconds; and left to stand for 20 minutes (1 cycle). Samples were analyzed after 3, 10, 20, or 50 cycles. Other bottles were filled with distilled water and left to stand for 10 days at 40°C.	ND (< 0.03 mg/kg) [< 30 $\mu\text{g/kg}$ or ppb] under any condition.	Mountfort et al. (19)
6 infant feeding bottles (country of purchase not known)	Bottles were filled with water at 26°C and left to stand for 5 hours or filled with water at 95°C and left to stand overnight.	ND (< 2 ppb [$\mu\text{g/L}$]) in bottles filled with water at 26°C and 3.1–55 ppb [$\mu\text{g/L}$] in bottles filled with water at 95°C.	Hanai et al. (1997) ^a
14 samples of new infant feeding bottles and tableware including a bowl, mug, cup, and dish recalled because residual bisphenol A and other phenol levels exceeded 500 ppm [mg/kg] (Japan)	Products were exposed to n-heptane, water, 4% acetic acid, or 20% ethanol; in some cases simulant was heated to 60 or 95°C; in other cases, the object was boiled for 5 minutes; analyses were usually conducted after a 30-minute contact period.	Up to 40 ppb [$\mu\text{g/kg}$] in recalled products and ND (< 0.2) to 5 $\mu\text{g/kg}$ in commercially available products.	Kawamura et al. (1998) ^{a,b}
Discs prepared from commercial food-grade polycarbonate resins (residual bisphenol A at 8800 to 11,200 $\mu\text{g/kg}$) from US manufacturers	Materials exposed to water, 10% ethanol, or Miglyol® (fractionated coconut oil) at 100°C for 6 hours or water, 3% acetic acid, 10% ethanol, or Miglyol at 49°C for 6–240 hours.	ND (< 5 ppb [$\mu\text{g/L}$]) under all conditions.	Howe and Borodinsky (26)

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Sample (Location)	Procedure	Bisphenol A level	Reference
2 bottles from Japan	In 3 repeated tests, boiling water was added to bottles; bottles were incubated at 95°C for 30 minutes and cooled to room temperature. Prior to repeating the test a 4 th time, the bottles were scrubbed with a brush.	Below quantification limit (0.57 ppb [$\mu\text{g/L}$]) to mean levels of 0.75 ppb prior to brushing and <0.57 to 0.18 ppb after brushing.	Sun et al. (27)
4 new different brands of infant bottles (Argentina)	Bottles were exposed to distilled water, 3% acetic acid, or 15% ethanol at 80°C for 2 minutes or distilled water at 100°C for 0.5 minutes.	1.1–2.5 ppb [$\mu\text{g/L}$].	D’Antuono et al. (28)
12 infant bottles (Norway)	Bottles were tested prior to washing and following 51 and 169 dishwashings; bottles were occasionally brushed (13 times by 2 nd test and 23 times by 3 rd test) and boiled (12 times by 2 nd testing and 25 times by 3 rd testing). Unwashed bottles were rinsed with boiling water before testing. For testing, bottles were filled with hot water and incubated at 100°C for 1 hour.	0 washes: 0.11–0.43 $\mu\text{g/L}$ [ppb]; 51 washes: 3.7–17 $\mu\text{g/L}$; 169 washes: 2.5–15 $\mu\text{g/L}$.	Brede et al. (21)
18 infant bottles (12 tested) (UK)	Bottles were tested prior to and after 20 and 50 dishwashings; bottles were brushed after every 2 wash cycles. Bottles were sterilized with boiling water, filled with 3% acetic acid, or 10% ethanol, and incubated at 70°C for 1 hour.	Prior to washing: ND (< 1.1 ppb or $\mu\text{g/L}$) in 10% ethanol and ND (< 0.34 ppb or $\mu\text{g/L}$) in 3% acetic acid; 20 washes: ND to 4.5 ppb in 10% ethanol and ND to 0.64 ppb in 3% acetic acid; 50 washes: ND to 3.1 ppb in 10% ethanol and ND to 2.6 ppb in 3% acetic acid.	CSL (22)
28 brands of new infant bottles (residual bisphenol A levels of <3 to 141 mg/kg) manufactured in Europe or Asia (Singapore)	Bottles were cut, and the pieces were exposed to 10% ethanol at 70°C or corn oil at 100°C for 8–240 hours.	ND (< 0.05) to 1.92 $\mu\text{g/in}^2$ [5–192 $\mu\text{g/L}$ or ppb] in 10% ethanol and ND (< 0.05) to 6.54 $\mu\text{g/in}^2$ [<5–654 $\mu\text{g/L}$] in corn oil over the 240-hour exposure period.	Onn Wong et al. (18)
22 new infant bottles and 20 used (3–36 months) bottles (Netherlands)	Bottles were immersed in boiling water for 10 minutes prior to testing and filled with distilled water or 3% acetic acid and incubated at 40 °C for 24 hours.	ND in new bottles (< 2.5 $\mu\text{g/L}$ [ppb] in distilled water and < 3.9 $\mu\text{g/L}$ in 3% acetic acid) or in used bottles exposed to 3% acetic acid; not detected to non-quantifiable (<5 $\mu\text{g/L}$) in distilled water from used bottles.	FCPSA (29)

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Sample (Location)	Procedure	Bisphenol A level	Reference
New unwashed bottles (number not indicated) (Japan)	Bottles were exposed to water at 95°C for 30 minutes.	ND (< 0.05 µg/L [ppb]) to 3.9 µg/L.	Japanese studies reviewed in Miyamoto and Kotake (30)
5-gallon water carboys	Water was stored in the carboys for 3, 12, or 39 weeks, temperature not indicated.	0.0001–0.0005 µg/L [ppb] at 3 and 12 weeks and 0.0046–0.0047 µg/L at 39 weeks.	Biles et al. (23)

^aReviewed by European Union (2).

^bReviewed by Haighton et al. (24).

High molecular weight, heat-cured bisphenol A-based epoxy resins are used as protective linings in cans for food and beverages and occasionally in wine storage vats (2). Residual bisphenol A monomer can migrate from the coatings to foods or beverages contained within cans. Studies were conducted to measure actual levels of bisphenol A in commercially available foods or to measure levels of bisphenol A leaching from can linings into food simulants. Because the actual measurement of bisphenol A concentrations in canned foods represents the most realistic situation, the CERHR review will focus on those data. Studies conducted with simulant will not be reviewed, with the exception of one study by Howe et al. (31) that was considered by the FDA (25) in their estimates of bisphenol A intake.

Bisphenol A levels detected in infant foods are summarized in Table 3, and bisphenol A concentrations detected in non-infant foods are summarized in Table 4. With the exception of isolated cases in which bisphenol A levels were measured at up to ~0.6 mg/kg food, most measurements were below 0.1 mg/kg. The European Union also noted an extraction study conducted with an epoxy resin that is occasionally used to line wine vats. Based on that study, a worst-case scenario of 0.65 mg/L bisphenol A in wine was used. The European Union noted that the value represents a very worst-case exposure scenario but decided to use that number in risk estimates because no other value was available.

In one study, empty cans were filled with soup, beef, evaporated milk, carrots, or 10% ethanol (32). The cans were then sealed, processed at 5, 20, or 40°C, and sampled at 1 or 10 days or 1, 3, or 9 months. Half the cans processed according to each condition were dented. It was determined that 80–100% of the bisphenol A migrated to food immediately after processing, and that bisphenol A concentrations did not change during storage or as a result of denting. The study authors concluded that most migration occurred during can processing. Boiling the cans or heating to 230°C did not increase migration of bisphenol A, but that finding appears to contrast with findings of others. Kang et al. (33) examined the effects of temperature, duration of heating, glucose, sodium, and oil on migration of bisphenol A from cans. In cans filled with water, heating to 121°C compared to 105°C increased migration of bisphenol A but the duration of heating had no significant effect. Compared to cans filled with water, increased amounts of bisphenol A migrated from cans filled with 1–10% sodium chloride, 5–20% glucose, or vegetable oils and heated to 121°C. Takao et al. (34) reported increased leaching of bisphenol A from cans into water when the cans were heated to $\geq 80^\circ\text{C}$.

Table 3. Surveys of Bisphenol A Levels in Canned Infant Formulas or Foods

Food (no. sampled)	Bisphenol A level, $\mu\text{g}/\text{kg}$ or $\mu\text{g}/\text{L}$	Country	Reference
Infant formula (14)	Mean 5 (0.1–13.2); when diluted with water to make prepared formula, mean levels would be 0.0025 (0.00005–0.0066).	US	Biles et al. (35) and FDA (25)
Infant formula (4)	Not detected (<0.002)	UK	Goodson et al. (36) and UKFSA (37)
Infant formula (5)	44–113	Taiwan	Kuo and Ding (38)
Infant dessert ^a (1)	49.2–77.3	UK	Goodson et al. (32)
Infant food ^b (2)	18.9–46.7		
Infant vegetable food (4)	< LOQ (10)	New Zealand	Thomson and Grounds (39)
Infant dessert (3)	< LOQ (10)		

^aValues from heated and non-heated cans presented together; it could not be determined if heating resulted in different extraction rates.

^bValues prior to and following heating in can and from non-dented and dented cans; values did not differ under the various conditions and were presented together.

1 **Table 4. Surveys of Bisphenol A Levels in Canned or Bottled Foods or Food Simulants**

Food (no. sampled)	Bisphenol A level, µg/kg unless specified	Country of purchase ^a	Reference
Vegetables with liquid (6)	Mean 16 (4–39)	US	FDA (25)
Liquids from canned vegetables or mushrooms (10)	4.2 ± 4.1 (SD) to 22.9 ± 8.8 µg/can [12 ± 12 to 76 ± 29 µg/kg]	Spain and US	Brotons et al. (40)
Coffee (13)	3.3–213	Japan	Kawamura et al. (41)
Black tea (9)	8.5–90		(reviewed in (2);
Other tea (8)	3.7–22		English abstract available)
Alcoholic beverages (10)	Not detected (<2) to 12		
Soft drinks (7)	Not detected (<2)		
Vegetables (10)	9–48	UK	Goodson et al. (36)
Desserts (5)	Not detected (<2) to 14		and UKFSA (37)
Fruits (2)	19–38		
Pastas (5)	<7 to 41		
Meats (5)	16–70 (up to 422 in 1 sample) ^b		
Fish (10)	Not detected (<2) to 44		
Non-alcoholic or alcoholic beverages (11)	Not detected (<2) to <7		
Soups (10)	Not detected (<2) to 21		
Vegetables, fruits, or mushrooms (14)	Not detected (< 10) to 95.3 in solid portion; not detected (< 0.005) to 0.004 µg/mL in liquid portion; not detected to 11.1 µg/can [85 µg/kg] total		Yoshida et al. (42)
Meat products ^d (2)	8.6–25.7	UK	Goodson et al. (32)
Pasta ^d (1)	67.3–129.5		
Vegetables or beans ^c (2)	11.3–14.4		
Soup ^c (1)	18.5–39.1		
Pudding ^c (3)	3.8–53.2		
Pudding ^d (1)	18.5–28.1		
Grains and potatoes ^e	0 ^f –75	Japan	Reviewed in
Sugar, sweets, snacks	0 ^f –4		Miyamoto and Kotake (30)
Fats	0 ^f		
Fruits (including canned drinks), vegetables, mushrooms, seaweeds	0 ^f –450		
Seasoning and beverages	0 ^f –213		
Fish	9–480		
Meat and eggs	12.5–602		
Milk and dairy products	0 ^c –6		
“Other” [not specified further]	36–310		
Fruits and vegetables (38)	< LOQ (10) to 24	New Zealand	Thomson and Grounds (39)
Fish (8)	< LOQ (20) to 109		
Soup (4)	< LOQ (20) to 16		
Sauces (4)	< LOQ (10) to 21		
Meat (6)	< LOQ (20) to 98		
Pasta (4)	< LOQ (10)		
Dessert (2)	< LOQ (20)		
Coconut cream (3)	< LOQ (20) to 192		

1.0 Chemistry, Use, and Human Exposure

Food (no. sampled)	Bisphenol A level, µg/kg unless specified	Country of purchase ^a	Reference
Soft drinks (4) Beverages (7) Vegetables (6) (only solid portion was analyzed, with the exception of tomatoes) Fruits (4) Canned fat products such as soups, meats, and cream (9) (only solid portion was analyzed for fish) Tuna (9)	< LOQ (10) Not detected (< 0.9) to 3.4 8.5–35 5–24 2.1–37.6 < LOQ (7.1) to 102.7	Austria Mexico	Braunrath et al. (43) Munguía-López et al. (44)
Beverage/beer cans exposed to 10% ethanol at 150°F [65.6°C] for 30 minutes and then 120°F [48.9 °C] for 10 days. Food cans exposed to 10 or 95% ethanol at 250°F [121°C] for 2 hours and then 120°F [48.9 °C] for 10 days or at 212°F [100 °C] for 30 minutes and then 120°F [48.9 °C] for 10 days.	Not detected (< 5) Not detected (< 5) to 95 (mean 37) ^g	US	Howe et al. (31) and FDA (25)
Honey (107 samples; ~90% imported in epoxy-lined drums) Wine stored in steel, wood, or plastic vats; filled into glass bottles; or purchased in local markets (59)	Not detected (<2) to 33.3 < LOQ (0.2 ng/mL) to 2.1 ng/mL	Japan Austria	Inoue et al. (45) Brenn-Struckhofova and Chichna-Markl (46)

^aAlthough cans were purchased in 1 or 2 countries for each study, most studies reported that cans were packaged in various locations throughout North America, Europe, and/or Asia.

^bThe UKFSA noted that the higher levels of bisphenol A detected in 1 meat product likely resulted from the use of bisphenol A as a cross-linking agent in the resin at that time.

^cValues were obtained from heated and non-heated cans but presented together because it could not be determined if heating resulted in differing extraction rates.

^dValues were determined before and after heating in can and from non-dented and dented cans; because the values did not differ under the various conditions, they were presented together.

^eTotal number of samples analyzed was not reported.

^fAs reported by study authors; detection limits not specified.

^gA maximum level of 121 ppb reported in the first phase of the study was determined to have resulted from analytical interference.

1
2 A study examining aggregate exposures of US preschool age children measured bisphenol A levels in
3 liquid food and solid food served to the children at home and at child care centers (11). Duplicate plates
4 of food served to 9 children were collected over a 48-hour period. GC/MS analyses were conducted on 4
5 liquid food samples and 4 solid food samples from the child care center and 9 liquid food samples and 9
6 solid food samples from home. Bisphenol A was detected in all solid food samples, 3 liquid food samples
7 from the child care center, and 2 liquid food samples from the home. Concentrations of bisphenol A

1 ranged from <0.100 to 1.16 ng/g [**µg/kg**] in liquid foods and from 0.172 to 4.19 ng/g [**µg/kg**] in solid
2 food.

3
4 The study examining aggregate exposures of US preschool age children was repeated with a larger
5 sample and again measured bisphenol A levels in liquid food and solid food served to the children at
6 home and at child care centers (12). Bisphenol A levels were measured by GC/MS in food served over a
7 48 hour period to at least 238 children at home and 49 children at daycare centers. Bisphenol A was
8 detected in 83–100% of solid food samples; levels were reported at <LOD (0.8) to 192 ng/g [**µg/kg**].
9 Sixty-nine to 80% of liquid food contained detectable levels of bisphenol A; levels were reported at
10 <LOD (0.3)–17.0 ng/mL in liquid food. Data were also collected for hand wipes of 193 children at
11 daycare centers and 60 children at home. Bisphenol A was detected in 94–100% of handwipe samples;
12 levels ranged from <LOD [**not defined**] to 46.6 ng/cm². and food preparation surface wipes. Bisphenol A
13 was detected in 85–89% of food preparation surface wipes from homes; levels were reported at < LOD
14 [**not defined**] to 0.357 ng/cm².

15
16 A review by Miyamoto and Kotake (30) reported bisphenol A levels of 0.011–0.086 mg/kg in non-canned
17 foods such as fats, fruits, fish, meat, and eggs. One study was identified that measured bisphenol A levels
18 in fresh produce purchased in southern Italy (47). Fourteen types of produce were homogenized, and
19 bisphenol A was measured by GC/MS. Bisphenol A levels were below the detection limit [**not reported**]
20 in 5 produce samples. In the remaining samples, bisphenol A was detected at concentrations of $0.25 \pm$
21 0.02 (SD) to 1.11 ± 0.09 mg/kg. [**It is unexpected that many bisphenol A concentrations exceeded**
22 **levels detected in canned foods (Table 4). Study authors did not compare their findings with those**
23 **of other studies examining bisphenol A levels in foods.**]

24 25 1.2.3.3 Potential migration from dental sealants

26 Bisphenol A is used in the manufacture of materials found in dental sealants or composites (i.e., fillings)
27 (2). Examples of bisphenol-A derived materials used in dental sealants include bis-glycidylmethacrylate
28 and bisphenol A-dimethyl acrylate. Bisphenol A could potentially be present as an impurity or be released
29 during degradation of the dental materials. Sealants are comprised of an organic matrix, while composites
30 contain inorganic filler in addition to the organic matrix. The British Dental Association therefore
31 concluded that because composites contain less resin than sealers, there is likely to be lower exposure to
32 bisphenol A from composites than sealants (reviewed in (2)). During dental procedures, resin mixtures are
33 applied as fluid monomers and polymerized in situ by ultraviolet or visible light. According to the
34 European Union (2), patients can be exposed to bisphenol A during the polymerization stage.

35
36 In a review of in vitro studies examining bisphenol A migration from dental sealants, the European Union
37 (2) concluded that release of bisphenol A is likely to occur only with degradation of the parent monomer.
38 The data suggested that bis-glycidylmethacrylate does not degrade; therefore, release of bisphenol A is
39 only likely to occur with bisphenol A-dimethyl acrylate use. In vivo studies measuring bisphenol A in
40 saliva following sealant application were reviewed in detail by CERHR because they provide the most
41 relevant human exposure information.

42
43 Olea et al. (48) measured saliva levels of bisphenol A for 1 hour before and 1 hour after application of 50
44 mg bis-glycidylmethacrylate- and bisphenol A-dimethyl acrylate-based sealant across 12 molars of 18
45 patients. Levels of bisphenol A in saliva were measured by GC/MS and HPLC. Following treatment,
46 saliva contained ~90–931 µg bisphenol A. Based on an assumed saliva production rate of 0.5 mL/minute,
47 a saliva concentration of 3–30 µg/mL was estimated by the study authors. With the exception of 1 patient
48 who was excluded from the study, bisphenol A was not detected in saliva prior to sealant application.

49
50 Arenholt-Binslev (49) measured bisphenol A in saliva of 8 adult patients who each had 4 molars treated
51 with 38 mg of 1 of 2 sealants, Delton LC or Visio-seal. Saliva was collected prior to, immediately after,

1 and at 1 or 24 hours following treatment for measurement of bisphenol A levels by HPLC. Bisphenol A
2 was detected at 0.3–2.8 ppm immediately after application of Delton SC sealant (bisphenol A-dimethyl
3 acrylate sealant according to the European Union (2)) but was not detected 24 hours later (detection limit
4 = 0.1 ppm [**mg/L**]). Bisphenol A was not detected in saliva of patients who received the Visio-seal sealant
5 (bis-glycidyl dimethacrylate sealant, according to the European Union). It was noted that saliva bisphenol
6 A levels were much lower than those reported by Olea et al. (48). Possible reasons for the inconsistencies
7 in results between the 2 studies were stated to be differences in the amount of sealant used, co-elution of
8 compounds that could have confounded bisphenol A analysis, or possible hydrolysis of resin by saliva
9 esterases in the Olea et al. study.

10
11 Fung et al. (50), measured saliva bisphenol A levels in 40 patients treated with a dental sealant (Delton
12 Opaque Light-cure Pit and Fissure Sealant) that was understood to contain bisphenol A-dimethyl acrylate,
13 according to the European Union (2). Eighteen patients in the low-dose group received 8 mg dental
14 sealant on 1 tooth, and 22 patients in the high-dose group received 32 mg sealant on 4 teeth. Saliva and
15 blood were collected for HPLC analysis before the procedure and at 1 and 3 hours and 1, 3, and 5 days
16 after the procedure. More details of this study are included in Section 2.1.1.1. Analysis of the dental
17 sealant revealed that bisphenol A levels were below the detection limit of 5 ppb. At 1 hour following
18 treatment, Bisphenol A was detected only in saliva samples from 3 of the 18 volunteers in the low-dose
19 group and 13 of 22 samples from volunteers in the high-dose group. At 3 hours post-treatment, bisphenol
20 A was detected in samples from 1 of 18 volunteers in the low-dose group and 7 of 22 volunteers from the
21 high-dose group. Levels of bisphenol A in saliva at 1 and 3 hours following exposure were reported at
22 5.8–105.6 ppb [**µg/L**]. No bisphenol A was detected in saliva samples at 24 hours after treatment or in
23 serum samples at any time point. Differences in bisphenol A levels and the presence of bisphenol A in
24 saliva of the low-dose compared to the high-dose group at 1 and 3 hours achieved statistical significance.
25 The European Union (2) noted that the concentrations of saliva bisphenol A reported by Fung et al. (50)
26 were more than 250 times lower than those reported by Olea et al. (48).

27
28 Sasaki et al. (51) used an enzyme-linked immunosorbent assay (ELISA) technique to examine saliva
29 bisphenol A levels in 21 patients before and after 1 cavity was filled with 0.1 g of composite resin. The
30 resins consisted of bisphenol A diglycidylether methacrylate (i.e., bis-glycidyl dimethacrylate), triethylene
31 glycol dimethacrylate, and/or urethane dimethacrylate. Saliva was collected prior to treatment, during the
32 5 minutes following treatment, and then immediately after gargling with water. Following treatment,
33 saliva bisphenol A increased [**from ≤ 2 to ~ 15 – 100 µg/L**]. Gargling reduced bisphenol A to near
34 pretreatment levels [**≤ 5 µg/L**] in most patients, with the exception of 1 patient with the highest bisphenol
35 A level [**reduced from ~ 100 to 18 µg/L**]. [**An increase in saliva bisphenol A levels was noted in 1 of 2**
36 **patients receiving a composite consisting solely of urethane dimethacrylate.**] The study authors noted
37 that cross-reactivity is possible with the ELISA technique, but that cross reactivity between bisphenol A
38 diglycidylether methacrylate and triethylene glycol dimethacrylate is low. Therefore, the study authors
39 thought it possible that they were measuring only bisphenol A.

40
41 Joskow et al. (52) examined bisphenol A in urine and saliva of 14 adults treated with dental sealants. The
42 volunteers received either HeliOSEAL F (n = 5) or Delton LC (n = 9) sealant. Only the HeliOSEAL F sealant
43 was noted to carry the American Dental Association (ADA) Seal of Acceptance. Sealant was weighed
44 before and after application to determine the amount applied, and the numbers of treated teeth were
45 recorded. The mean number of teeth treated was 6/person and the mean total weight of sealant applied
46 was 40.35 mg/person. In a comparison of the 2 different sealants, no differences were reported for the
47 number of teeth treated or amount of sealant applied. Saliva samples were collected before, immediately
48 after, and 1 hour after sealant application. Urine samples were collected before and at 1 and 24 hours after
49 sealant placement. A total of 14–15 saliva samples and 12–14 urine samples were collected at each time
50 point. Samples were treated with β -glucuronidase and analyzed for bisphenol A levels using selective and
51 sensitive isotope-dilution-MS-based methods. Saliva levels were highest immediately following

1 treatment; mean levels were reported at 42.8 ng/mL in patients treated with Delton LC and 0.54 ng/mL in
2 patients treated with Heliaseal F. The highest mean urinary levels of bisphenol A were measured at 1 hour
3 following exposure and were reported at 27.3 ng/mL in patients treated with Delton LC and 7.26 ng/mL
4 in patients receiving the Heliaseal F sealant. The study authors noted that saliva and urine bisphenol A
5 levels following application of Heliaseal F were comparable to baseline levels. More information on
6 bisphenol A levels in saliva and urine is included in Section 2, and exposure estimates are provided in
7 Section 1.2.4.1.2. The study authors noted that saliva levels detected in their study were ~1000 times
8 lower than those reported by Olea et al. (48) but were within the ranges reported by Fung et al. (50) and
9 Sasaki et al. (51). Analytical procedures and use of a large amount of sealant were noted as possible
10 reasons for the higher values reported by Olea et al. (48).

11
12 The European Union noted a study by Lewis et al. (53) that characterized materials in 28 commercial
13 resin-based composites and sealants, including those examined by Olea et al. (48). HPLC and infrared
14 analysis could not verify the presence of bisphenol A in any sealant product. Lewis et al. noted that in the
15 study by Olea et al. another component in the resin may have been misidentified as bisphenol A because
16 of difficulties with resolution.

17
18 In their review of studies examining bisphenol A levels in saliva of patients treated with dental sealants,
19 the European Union (2) noted that the higher concentrations reported may have resulted from interference
20 during analysis and thus may overestimate bisphenol A exposures from dental treatments. It was
21 concluded that dental treatment would likely result in saliva bisphenol A levels of 0.3–3 ppm. Because
22 bisphenol A was generally not detected in saliva at time points beyond 1 hour after treatment, it was
23 concluded that bisphenol A exposure resulting from dental treatments is likely to be an acute event. In
24 their 2002 position statement, the ADA stated that none of the 12 dental sealants that carry the ADA Seal
25 release bisphenol A (54). Upon initial analysis, one of the sealants was found to leach trace levels of
26 bisphenol A, but following implementation of quality controls by the manufacturer, bisphenol A could no
27 longer be detected in the final product.

28 *1.2.3.4 Bisphenol A levels measured in biological samples*

29 Bisphenol A levels detected in human blood are summarized in Table 5. Additional studies reporting
30 bisphenol A levels in blood of pregnant women and fetal tissues and fluids are summarized in Section 2.1.
31 Goodman et al. (55) noted that although blood level may provide information on internal dose, it does not
32 allow for estimates of daily intake. It was also noted that in many studies in which blood level of
33 bisphenol A was measured, sample preparation and analysis methods were poorly reported. Therefore, it
34 was difficult to determine if parent or total bisphenol A (the sum of parent and metabolized bisphenol A)
35 was reported. As noted in greater detail in Section 2, the majority of bisphenol A is systemically absorbed
36 and circulated as a glucuronide compound. Many study groups used an ELISA method to measure blood
37 bisphenol A level. Goodman et al. (55) and Fukata et al. (56) stated that the ELISA technique is likely to
38 overestimate bisphenol A levels as a result of cross-reactivity with substances such as bisphenol A
39 glucuronide.

40
41
42 Several studies reported levels of bisphenol A in human urine; those studies are summarized in Table 6.
43 As discussed in greater detail in Section 2, the majority of ingested bisphenol A is excreted in urine as
44 bisphenol A glucuronide. Smaller amounts of bisphenol A are metabolized to and excreted as bisphenol A
45 sulfate. Some of the studies determined levels of parent bisphenol A before and after digestion with
46 glucuronidases. With the exception of Fujimaki et al. (57) who used an ELISA technique to measure
47 urinary bisphenol A, other study authors used techniques such as HPLC, GC/MS, and liquid
48 chromatography (LC)/MS. Results from 394 participants of the National Health and Nutrition
49 Examination Survey (NHANES) III survey are included in Table 6 (58). Bisphenol A was detected in
50 95% of the participants, which indicated widespread exposure to bisphenol A in the US. Consistent with
51 those findings, bisphenol A was detected in urine from 85 of 90 (94.4%) 6–8-year-old girls from the US

1 (59) In a review of urinary bisphenol A data, Goodman et al. (55) noted that in most cases, median total
 2 urinary bisphenol A concentration (the sum of parent and conjugated bisphenol A) were ~1–2 µg/L. Two
 3 studies (60, 61) reported urinary bisphenol A levels that were orders of magnitude higher than commonly
 4 observed levels, despite the use of apparently reliable analytical techniques. Goodman et al. (55) stated
 5 that reported hormone levels for the study volunteers were also higher than expected, indicating the
 6 possibility of laboratory or reporting error.

7
 8

Table 5. Blood Levels of Bisphenol A in Adults

Population	Bisphenol A, µg/L ^a	Method	Reference
7 males and 12 females in Germany	<0.5	HPLC	Völkel et al. (62)
<i>Japanese women</i>			
Healthy premenopausal (n = 30)	2.0 ± 0.8	ELISA	Ikezuki et al. (63)
Non-obese, average age 27.5 years (n = 19)	0.71 ± 0.09 (SEM)	ELISA	Takeuchi et al. (64)
Obese, average age 28.8 years (n = 7)	1.04 ± 0.09 (SEM)		
Hyperprolactinemic, average age 27.7 years (n = 7)	0.83 ± 0.12 (SEM)		
Amenorrhic, average age 25.1 years (n = 7)	0.84 ± 0.10 (SEM)		
Non-obese with polycystic ovarian syndrome, average age 26.5 years (n = 13)	1.05 ± 0.10 (SEM)		
Obese with polycystic ovarian syndrome, average age 24.7 years (n = 6)	1.17 ± 0.16 (SEM)		
Normal (n = 14)	0.64 ± 0.10 (SEM)	ELISA	Takeuchi et al. (65)
Polycystic ovary syndrome (n = 16)	1.04 ± 0.10 (SEM)		
Unspecified age (n = 12)	0.33 ± 0.54	HPLC	Sajiki et al. (66)
45 patients (mean age 31.6 years) who had experienced multiple miscarriages	2.59 ± 5.23	ELISA	Sugiura-Ogasawara (67)
32 healthy woman (mean age 32 years)	0.77 ± 0.38	ELISA	Sugiura-Ogasawara (67)
<i>Japanese men and unspecified sex</i>			
Normal (n = 11)	1.49 ± 0.11 (SEM)	ELISA	Takeuchi et al. (65)
Unspecified age (n = 9)	0.59 ± 0.21	HPLC	Sajiki et al. (66)
21 sets of serum samples obtained from sterile patients of unspecified sex and age, median (range)	0.44 (0.22–0.87)	HPLC	Kuroda et al. (68)

^aMean ± SD unless otherwise specified.

1 **Table 6. Urinary Levels of Bisphenol A and Metabolites in Adults or Children**

Country	Study population	Urinary bisphenol A or metabolite concentrations in median (range) or mean \pm SEM, $\mu\text{g/L}^a$				Reference
		Free	Total	Glucuronide	Sulfate	
US	30 urine samples from demographically diverse, anonymous adult volunteers	< 0.3 (<0.3–0.6)	2.12 (<LOD ^b –19.8)	1.4 (<LOD ^b –19.0)	0.3 (<LOD ^b –1.8)	Ye et al. (69)
US	394 adult volunteers (males and females; 20–59 years old) from the NHANES III survey		1.28 (10 th to 95 th percentile: 0.22–5.18) ^c			Calafat et al. (58)
US	90 girls (6–8-years-old; White, Black, Asian, or Hispanic ethnicity)		1.8 (<0.3–54.3)			Wolff et al. (59)
Germany	7 males and 12 females	<1.14		<65 nM [15 $\mu\text{g/L}$]		Völkel et al. (62)
Korea	15 men (age 42.6 \pm 2.4 ^d years)	0.28–2.36; 0.58 \pm 0.14	0.85–9.83; 2.82 \pm 0.73	0.16–11.67; 2.34 \pm 0.85	<MDL ^e –1.03; 0.49 \pm 0.27	Kim et al. (70)
Korea	15 women (age 43.0 \pm 2.7 ^d years)	0.068–1.65; 0.56 \pm 0.10	1.00–7.64; 2.76 \pm 0.54	<MDL ^e –4.34; 1.00 \pm 0.34	<MDL ^e –3.40; 1.20 \pm 0.32	Kim et al. (70)
Japan	48 female college students	\leq 0.2		1.2 (0.2–19.1)		Ouchi and Watanabe (71)
Japan	23 females and 46 males; in each volunteer, 2 samples per volunteer were combined	0.01–0.27	Mean: 0.81 (range: 0.14–5.47)			Tsukioka (72)
Japan	Whole-day urine samples collected from 11 males and 11 females		Mean: 0.81 (range 0.24–2.03)			Tsukioka (72)
Japan	Spot urine samples collected from 56 women who were 1–9 months pregnant; 21–43 years of age		<1.1 (<1.1–5.4) ^c (ELISA)			Fujimaki et al. (57)
China	10 healthy male volunteers age 21–29 years		<2.7 to 3950; 1220 \pm 1380 ^d			Mao et al. (60)
China	10 healthy female volunteers age 21–29 years		30–3740; 1290 \pm 1220 ^d			Mao et al. (60)
Korea	34 males and 39 females (mean age 48.5 years)		Geometric mean: 9.54 (<0.012–586.14) ^b			Yang et al. (61)

^aWith the exception of the study by Fujimaki et al. (57), the studies used analytical techniques based on HPLC, GC/MS, and LC/MS. ^bLimit of detection (LOD) for bisphenol A following digestion of conjugate was 0.3 $\mu\text{g/L}$. ^cSamples were only digested with β -glucuronidase and do not account for bisphenol A conjugated to sulfate.

^dVariance not indicated. ^eMinimum detection limit based upon free bisphenol A.

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1.2.4 Human exposure

1.2.4.1 General population exposure

1.2.4.1.1 Estimates based on bisphenol A levels in food or environment

Wilson et al. (11) estimated aggregate exposures to bisphenol A in preschool aged children (2–5 years) from the US. In 1997, numerous chemicals were surveyed, but only bisphenol A results are reported here. Ten child care centers were surveyed and the 2 centers with the highest and lowest overall concentrations of target pollutants were selected for the study. Both centers were located in North Carolina. Nine children who attended one of the child care centers participated in the study. Over a 48-hour period, bisphenol A levels were measured in indoor and outdoor air, dust, soil, and food; the ranges detected are summarized in Sections 1.2.3.1 and 1.2.3.2. In estimating exposures, absorption was considered to be 100%. Calculations considered ventilation rates, time spent indoors and outdoors, time spent at home and in day care, the measured weight of each child, assumed ingestion of dust and soil, and total weight of foods consumed. Mean (range) bisphenol A intake was estimated at 0.042981 (0.018466–0.071124) µg/kg bw/day.

Wilson et al. (12) conducted a second study to estimate aggregate exposures in 257 US children aged 1.5–5 years. Bisphenol A was one of the compounds assessed in homes and daycare centers in 6 North Carolina and 6 Ohio counties in 2000–2001. Over a 48-hour period, bisphenol A levels were measured in indoor and outdoor air, dust, soil, food, and surface and handwipes; the ranges detected are summarized in Sections 1.2.3.1 and 1.2.3.2. In estimating exposures, absorption was considered to be 50%. Calculations considered ventilation rates, time spent indoors and outdoors, time spent at home and in day care, the measured weight of each child, assumed ingestion of dust and soil, and total weight of foods consumed. Median (25th percentile to maximum) bisphenol A aggregate exposures were estimated at 2.56 (1.5–57.2) µg/day for children from North Carolina and 1.88 (1.27–48.6) µg/day in children from Ohio. Median (25th percentile to maximum) potential aggregate dose, assuming 50% absorption was estimated at 0.0714 (0.0424–1.57) µg/kg bw/day in children from North Carolina and 0.0608 (0.0341–0.775) µg/kg bw/day in children from Ohio. The study authors noted that 99% of exposure occurred through dietary ingestion.

The European Union (2) conducted a comprehensive exposure estimate that considered exposures resulting from food and environmental sources. Oral exposure estimates for children and adults were reported and are summarized in Table 7. Estimates were based on migration studies conducted with polycarbonate and levels of bisphenol A measured in foods packaged in epoxy-lined cans. Assumptions used in exposure estimates included 100% oral absorption and body weights of 70 kg for adults, 14.5 kg for 1.5–4.5-year-old children, 4.5 kg for 1–2-month-old infants, 7 kg for 4–6-month-old infants, and 8.7 kg for 6–12-month-old infants. Estimated exposures for children were said to represent realistic worst-case scenarios for food and drink intake relative to body weight.

1 **Table 7. Bisphenol A Oral Exposure Estimates by the European Union**

Exposure source (exposed population)	Daily food intake	Bisphenol A level in food	Bisphenol A intake	
			µg/day	µg/kg bw/day
Infant bottles (1–2 month-old infant)	0.699 L/day milk	50 µg/L	35	8
Infant bottles (4–6-month-old infant)	0.983 L/day milk	50 µg/L	50	7
Polycarbonate tableware (1.5–4.5- year old child)	2 kg food/day	5 µg/kg	10	0.7
Canned food (6–12- month-old infant)	0.375 kg canned food/day	100 µg/kg	40	5
Canned food (1.5– 4.5-year-old child)	2 kg canned food/day	100 µg/kg	200	14
Canned food (adult)	1.0 kg canned food/day	100 µg/kg	100	1.4
Wine (adult)	0.75 L/day	650 µg/L	500	7 ^a
Canned food and wine (adult)	0.75 L/day wine and 1.0 kg canned food/day	650 µg/L in wine and 100 µg/kg food	600	9 ^a

^aThe European Union acknowledged that exposure through wine represents a very worst-case scenario. From the European Union (2).

2
3 The European Union (2) also estimated human environmental exposure to bisphenol A from sources such
4 as drinking water, fish, plants, milk, meat, and air. The values were apparently obtained using the
5 “EUSES” model. Total regional exposure to bisphenol A was estimated at 0.0178 µg/kg bw/day. The
6 highest local exposure was thought to occur in the vicinity of PVC-producing plants and was estimated at
7 59 µg/kg bw/day. Aggregate exposures in adults involving food, wine, and environmental sources were
8 estimated at 9 µg/kg bw/day for regional scenarios and 69 µg/kg bw/day for worst-case local scenarios
9 occurring near a PVC-manufacturing plant. However, it was noted in the European Union report that use
10 of bisphenol A in PVC manufacture was being phased out.

11
12 The European Union (2) noted that exposures to bisphenol A through dental sealant are single and rare
13 events and do not lead to repeated exposure. Therefore, the issue was not considered further.

14
15 Exposures to bisphenol A from some consumer products were identified and characterized by the
16 European Union (2). Products included: marine antifouling agents used on boats, wood varnish, wood
17 fillers, and adhesives. With the exception of adhesives for which frequent use was thought possible,
18 exposure to the other products was considered to be relatively rare. Exposures were estimated based on
19 factors such as epoxy and residual bisphenol A levels, exposure time, area of skin exposed, and possible
20 generation of mists during processes such as brushing. Inhalation exposures per product use were
21 estimated at 3×10^{-4} µg for antifouling agents and 0.02 µg for wood varnish. Dermal exposure per
22 product use without protective clothing was estimated at 29 µg for antifouling agents, 3.6 µg for wood
23 varnish, 9 µg for wood filler, and 14 µg for adhesives. **[Dermal exposure to adhesives appears to be**
24 **incorrectly reported as 1 µg in Table 4.20 of the European Union review.]** Exposure was estimated to
25 be 1–2 orders of magnitude lower when protective clothing such as gloves was used. Assuming an
26 absorption rate of 10%, dermal exposure to bisphenol A through adhesives was estimated at 0.02 µg/kg
27 bw/day.
28

1 The European Commission (73) reviewed the report by the European Union (2) in draft and suggested
 2 alternate exposure estimates. Those estimates and the assumptions used to support those estimates are
 3 summarized in Table 8.

4
 5 **Table 8. Bisphenol A Exposure Estimates by the European Commission**

Age and body weight	Type of food and amount consumed	Concentration of bisphenol A in food, $\mu\text{g}/\text{kg}$	Estimated body burden, $\mu\text{g}/\text{kg bw}/\text{day}$
0–4-month old infant, 4.5 kg	0.7 L of formula/day	10	1.6
6–12-month old infant, 8.8 kg	0.7 L of formula/day	10	0.8
6–12-month old infant, 8.8 kg	0.38 kg canned food/day	20	0.85
4–6-year-old child, 18 kg	1.05 kg canned food/day	20	1.2
Adult, 60 kg	1.05 kg canned food/day	20	0.37
Adult, 60 kg	0.75 L wine/day	9	0.11

From the European Commission (73).

6
 7 Miyamoto and Kotake (30) estimated aggregate oral and inhalation exposure to bisphenol A in Japanese
 8 male children and adults. The estimates were based on unpublished Japanese data. This report is the only
 9 known study investigating potential exposure to children through mouthing of toys. Mouthing times were
 10 estimated by surveying the mothers of 50 infants and recording 25 infants on video camera. Mean \pm SD
 11 mouthing times were reported at 41.7 ± 13.7 minutes for infants 0–5 months of age and 73.9 ± 32.9
 12 minutes for infants 6–11 months of age. Migration rates were estimated from $0 \mu\text{g}/\text{cm}^2/\text{minute}$ for toys
 13 that do not contain bisphenol A to $0.0162 \mu\text{g}/\text{cm}^2/\text{minute}$, the highest value reported in the Japanese
 14 literature. It was assumed that most toys were not manufactured with polycarbonate, epoxy resins, or
 15 grades of PVC that contain bisphenol A. Surface area of toys was assumed to be 10 cm^2 . In estimating
 16 oral exposures to bisphenol A, intake from food was also considered. Bisphenol A levels measured in
 17 migration testing of polycarbonate bottles and food surveys are summarized in Section 1.2.3.2. Volume of
 18 food consumption and frequency of article use were considered in estimates of bisphenol intake through
 19 food. Bisphenol A concentrations in drinking water were considered to be 0– $0.17 \mu\text{g}/\text{L}$, and water intake
 20 was assumed to be 2 L/day. In estimating inhalation exposures, concentrations of bisphenol A were
 21 considered to range from 0 to $8.1 \text{ ng}/\text{m}^3$ in indoor air and 0 to $28 \text{ ng}/\text{m}^3$ in outdoor air. Time spent indoors
 22 and outdoors and breathing rates were considered. Absorption from lungs was assumed at 100%.
 23 Estimated exposures from mouthing of toys, food and water intake, and inhaled air are summarized in
 24 Table 9.

25

1 **Table 9. Average Estimated Exposure to Bisphenol A in Japanese Male Adults and Children**

Exposure source	Bisphenol A level (other assumptions)	Average estimated exposures ($\mu\text{g}/\text{kg}$ bw/day) in each age group ^a					
		0–5 months	6–11 months	1–6 years	7–14 years	15–19 years	19 years
Human milk	Negligible	0	0				
Formula (water)	0–0.17 $\mu\text{g}/\text{L}$	0.012	0.0096				
Feeding bottle	0–3.9 $\mu\text{g}/\text{L}$	0.015	0.014				
Infant food	0–5.0 $\mu\text{g}/\text{kg}$		0.085				
Toys	0–0.0162 $\mu\text{g}/\text{cm}^2/\text{minute}$ (mean mouthing times of 41.7 minutes in 0–5 month olds and 73.9 minutes in 6–11 month olds)	0.026	0.069				
Air	0–8.1 ng/m^3 in indoor air and 0–28 ng/m^3 in outdoor air (90% indoors and 10% outdoors)	0.0026	0.0024	0.0021	0.0017	0.0015	0.0015
Water	0–0.17 $\mu\text{g}/\text{L}$ (intake of 2 L/day)			0.012	0.0053	0.0029	0.0027
Food and drink							
Canned	0–602 $\mu\text{g}/\text{kg}$			0.38	0.21	0.20	0.29
Non-canned	0–3 $\mu\text{g}/\text{kg}$			0.38	0.21	0.13	0.12
Tableware	0–39.4 $\mu\text{g}/\text{meal}/\text{utensile}$ (3 meals/day; 1–5 types of utensiles used/meal)			0.40	0.12	0.024	0.022
Total		0.028 (breast- fed) 0.055 (formula- fed)	0.16 (breast- fed) 0.18 (formula- fed)	1.2	0.55	0.36	0.43

^aAssumptions for bodyweights and most media intake levels were not provided.

Source: Miyamoto and Kotake (30).

2
3 Additional estimates of bisphenol A exposure through food are summarized in Table 10. Details of
4 studies conducted by Earls et al. (20) and Onn Wong et al. (18) are presented in Section 1.2.3.2. Exposure
5 estimates conducted by the FDA are described below. Limited details were available from the other
6 studies that were presented in reviews.

7
8 The FDA (25) estimated bisphenol A intake in infants and adults resulting from exposures to epoxy food-
9 can linings and polycarbonate plastics. Exposure estimates occurring through contact of formula with
10 polycarbonate bottles were based on results of a study conducted by the Chemistry Methods Branch of the
11 FDA. The Chemistry Methods Branch also measured levels of bisphenol A in 5 brands of infant formula
12 (14 samples total); the study is also published as Biles et al. (35). In estimating adult bisphenol A
13 exposure through the consumption of canned foods, the FDA considered surveys conducted by the
14 Chemistry Methods Branch, Brotons et al. (40), and the Society of Plastics Industry Group. It appears that
15 the study by the Society of Plastics Industry Group was later published by Howe et al. (31) and included a

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1 re-analysis to correct some interferences observed in analytical methods. Exposure estimates and
 2 assumptions used to make the estimates are summarized in Table 10.

3
 4

Table 10. Summaries of Studies Estimating Bisphenol A Exposures Solely from Foods

Population	Exposure source	Basis and assumptions for estimates	Exposure estimate, $\mu\text{g}/\text{kg}$ bw/day	Reference
Infants	Polycarbonate bottles	Bisphenol A migration level of 15–20 $\mu\text{g}/\text{L}$; milk consumption of up to 550 mL/day; mean body weight of 11 kg.	0.75–1	Earls et al. (20)
Infants (0–3 months old)	Polycarbonate bottles	Mean upper-bound level of bisphenol A migration in 10% ethanol (0.64 $\mu\text{g}/\text{in}^2$) and in corn oil (0.43 $\mu\text{g}/\text{in}^2$); body weights reported by National Center for Health Statistics, and FDA Dietary Exposure Guidelines with modifications for properties of infant formula.	15–24 ^a	Onn Wong et al. (18)
Not reported	Food from epoxy-lined cans	Bisphenol A levels of 5 ppb [$\mu\text{g}/\text{L}$] in beverages and 37 ppb [$\mu\text{g}/\text{kg}$] in other foods; FDA Dietary Exposure Guidelines: dietary intake of 3 kg/day, body weight of 60 kg.	0.105	Howe et al. (31), Haighton et al. (24), and NAS (74)
Adults	Cumulative exposures from food contacting cans and polycarbonate plastics	22 ppb [$\mu\text{g}/\text{kg}$] bisphenol A in vegetables, consumption factor of 0.17 for food contacting polymer-coated metal, intake of 3 kg food/bw/day, 60 kg bw, and insignificant contribution from polycarbonate	0.183	FDA (25)
Infants	Cumulative exposures from food contacting cans and polycarbonate plastics	Bisphenol A concentration of 6.6 $\mu\text{g}/\text{kg}$ in prepared infant formula, < 1.7 ppb [$\mu\text{g}/\text{L}$] in infant formula from polycarbonate bottles, consumption of 820 g food/day, and 4 kg infant weight	1.75	
Adults	Canned foods	Data from survey of canned foods and food intake patterns determined from surveys	Mean 0.0083 (0–0.29)	Thomson and Grounds (39)
Adults	Canned foods and canned fish	Data from survey of canned foods and food intake patterns determined from surveys	0.0044 for males ≥ 25 , 0.0041 for females ≥ 25 , and 0.0048 for males age 19–24	Thomson et al. (75)
Adults	Wine	Maximum bisphenol A concentration of 2.1 ng/mL in wine, consumption of 0.75 L/day, and 60 kg body weight.	<0.026	Brenn-Struckhoffova and Cichna-Markel (46 2393)

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Population	Exposure source	Basis and assumptions for estimates	Exposure estimate, $\mu\text{g}/\text{kg bw}/\text{day}$	Reference
Hospital patients	Meals served at 2 hospitals	Mean intake from hospital diets was estimated at 1.3 (0.19–3.7 $\mu\text{g}/\text{day}$); [60 kg body weight was assumed]	[0.02 (0.003–0.06)]	Imanaka (2001) as cited in Miyamoto and Kotake (30) and Fujimaki et al. (57)
Japanese adults and children	~200 food items were collected in a total diet study	No details	0.00475 for children age 2–6 years and 0.00195 for adults	Tokyo Metropolitan Government (2003) as cited in Miyamoto and Kotake (30)

^aThe study authors acknowledged the use of aggressive migration testing conditions and conservative assumptions in calculations, thus leading to overestimated infant exposures.

1
2 Table 11 summarizes exposure estimates for aggregate or food exposures. Studies suggest that the
3 majority of bisphenol A exposure occurs through food and that environmental exposures do not appear to
4 substantially affect exposure, with the possible exception of exposure near point sources. Table 11
5 includes estimates that CERHR believes to represent potentially realistic exposure scenarios and does not
6 include data from extreme worst-case scenarios such as possible point-source exposures.

7

8 **Table 11. Summary of Aggregate or Food Exposures to Bisphenol A**

Population	Basis of Estimates	Exposure estimate, $\mu\text{g}/\text{kg bw}/\text{day}^a$	Reference
1–2-month old infant	Food exposure (data from migration studies of polycarbonate bottles)	8	European Union (2)
0–4-month old infant	Food exposure (data from migration studies of polycarbonate bottles)	1.6	European Commission (73)
0–5-month old infant (formula-fed)	Aggregate exposure (based on formula, environmental, and toy exposures)	0.055	Miyamoto and Kotake (30)
0–5-month old infant (breast fed)	Aggregate exposure (based on human milk, environmental, and toy exposures)	0.028	Miyamoto and Kotake (30)
4–6-month old infant	Food exposure (data from migration studies of polycarbonate bottles)	7	European Union (2)
6–11-month-old infant (formula-fed)	Aggregate exposure (based on formula, food, environmental, and toy exposures)	0.18	Miyamoto and Kotake (30)
6–11-month-old infant (breast-fed)	Aggregate exposure (based on human milk, food, environmental, and toy exposures)	0.16	Miyamoto and Kotake (30)
6–12-month-old infant	Food exposure (data from survey of canned foods)	5	European Union (2)
6–12-month-old infant	Food exposure (data from migration studies with infant bottles and canned foods)	1.65	European Commission (73)
Infant	Food exposure (data from polycarbonate bottle leaching studies)	0.75–1	Earls et al. (20)
Infant	Food exposures (contact with cans and polycarbonate plastics)	1.75	FDA (25)

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Population	Basis of Estimates	Exposure estimate, $\mu\text{g}/\text{kg bw}/\text{day}^{\text{a}}$	Reference
1.5–4.5-year-old child	Food exposure (data from survey of canned foods and migration studies with polycarbonate tableware)	14.7	European Union (2)
1–6-year-old child	Aggregate exposure (based on food, environmental, and tableware exposures)	1.2	Miyamoto and Kotake (30)
1.5–5 year old child	Aggregate exposure (surveys of bisphenol in food, air, dust, soil and hand and surface wipes)	0.0714–0.0608 (0.0341–1.57)	Wilson et al. (12)
3–5-year-old child	Aggregate exposure (surveys of bisphenol in food, air, dust, and soil)	0.042981 (0.018466– 0.071124)	Wilson et al. (11)
2–6 year-old child	Food exposure (collection of 200 food items)	0.00475	Tokyo Metropolitan Government (2003) as cited in Miyamoto and Kotake (30)
4–6 year-old child	Food exposure (data from survey of canned foods)	1.2	European Commission (73)
7–14 year-old child	Aggregate exposure (based on food, environmental, and tableware exposures)	0.55	Miyamoto and Kotake (30)
15–19 year-old individual	Aggregate exposure (based on food, environmental, and tableware exposures)	0.36	Miyamoto and Kotake (30)
Adult, ≥ 19 years	Aggregate exposure (based on food, environmental, and tableware exposures)	0.43	Miyamoto and Kotake (30)
Adult	Food exposure (data from survey of canned foods not including wine)	1.4	European Union (2)
Adult	Food exposure (data from surveys of canned food)	0.37	European Commission (73)
Adult	Wine exposure (data from study of epoxy-lined wine drums)	0.11	European Commission (73)
Adult	Wine exposure (data from wine samples)	<0.026	Brenn-Struckhoffova and Cichna-Markel (46)
Adult	Food exposure (from contact with epoxy-lined cans and polycarbonate)	0.183	FDA (25)
Adults	Food exposure (survey of canned foods)	0.0083	Thomson and Grounds (39)
Adult	Food exposure (collection of 200 food items)	0.00195	Tokyo Metropolitan Government (2003) as cited in Miyamoto and Kotake (30)

^aEstimates involving extreme worst case scenarios and Japanese data with very limited information were not included in this table.

- 1
- 2 *1.2.4.1.2 Estimates based on biological monitoring*
- 3 Goodman et al. (55) noted that total urinary bisphenol A concentrations were useful for estimating
- 4 bisphenol A intake. Because nearly 100% of bisphenol A is excreted in urine within 24 hours, bisphenol

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1 A intake can be estimated by measuring bisphenol A in urine over a specified time interval. Two studies
2 were identified that measured urinary bisphenol A over a 24-hour period; those studies are summarized in
3 Table 12. An estimate based on a Monte Carlo analysis of data from the 2 studies is also summarized in
4 Table 12. One of the studies (76) measured bisphenol A excretion over a 5-day period and reported intra-
5 and inter-individual variability. As a result, caution was urged in using single time-point values to
6 estimate long-term exposure. However, it was noted that single values can be useful in estimating mean
7 population values in a cross-sectional study because the variations average out over time. Goodman et al.
8 (55) noted a study by Fugimaki et al. (57) that estimated bisphenol A intake in pregnant woman based on
9 spot urine samples. Details and results of the study are summarized in Table 12. Goodman et al. (55)
10 concluded typical daily intakes of bisphenol A are ~0.02–0.06 µg/kg bw/day in adults, based on urinary
11 levels. **[Consistent with adult findings, estimated mean exposure based on urinary bisphenol A**
12 **levels in 6–8-year-old girls was 0.059 µg/kg bw/day; see Wolff et al. (59) in Table 12.]** Because of
13 extensive first-pass metabolism, a small percentage of the parent compound is systemically circulated, as
14 discussed in more detail in Section 2.

15
16 Joskow et al. (52) used values for total bisphenol A in urine to estimate exposure to bisphenol A
17 following dental sealant application. Urinary levels of bisphenol A were reported in Section 2. Factors or
18 assumptions used in the exposure estimates were recovery of bisphenol A in urine as its glucuronide
19 conjugate within 24–34 hours following exposure, a 5.4 hour half-life of elimination for bisphenol A
20 glucuronide, and a 1.5 L/day urinary excretion volume. Estimated doses of bisphenol A **[based on a 60-**
21 **kg bw]** were 49–239 µg **[0.82–4.0 µg/kg bw]** following application of Delton LC and 0–9.5 µg **[0–0.16**
22 **µg/kg bw]** following application of Helioseal F. The study authors stated that the estimates were likely
23 low because a substantial amount of bisphenol A was potentially eliminated by collection of saliva
24 samples immediately following treatment.

25

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1 **Table 12. Estimates of Bisphenol A Intakes Based on Urinary Excretion**

Population	Basis for estimates	Mean or median (range) of estimated intake, $\mu\text{g}/\text{kg bw}/\text{day}^{\text{a}}$	Reference
22 Japanese adults	Mean excretion of 1.68 $\mu\text{g}/\text{day}$ (0.48–4.5 $\mu\text{g}/\text{day}$)	0.028 (0.008–0.075)	Tsukioka et al. (72)
36 Japanese male students	Median excretion of 1.2 $\mu\text{g}/\text{day}$ (<0.21–14 $\mu\text{g}/\text{day}$)	0.02 (<0.0035–0.23)	Arakawa et al. (76)
5 Japanese males	Median excretion of 1.3 $\mu\text{g}/\text{day}$ (<0.58–13 $\mu\text{g}/\text{day}$) over a 5-day period	0.022 (<0.01–0.22)	Arakawa et al. (76)
Data from Tsukioka et al. (72) and Arakawa et al. (76)	Monte Carlo simulations	Mean exposure: 0.028–0.049 in males and 0.034–0.059 in females; low exposures (5 th percentile) 0.021–0.037 in males and 0.025–0.044 in females; high exposures (95 th percentile): 0.037–0.064 in males and 0.043–0.075 in females	Miyamoto and Kotake (30)
56 pregnant Japanese women	Bisphenol A level in 1 spot sample was normalized to creatinine and exposure was estimated using average creatine and urine volume excretion rates, which resulted in a median intake of <2 $\mu\text{g}/\text{day}$ (<0.3–7.9 $\mu\text{g}/\text{day}$).	<0.04 (<0.006–0.16) ^b	Fugimaki et al. (57)
48 Japanese female college students	Authors estimated bisphenol A intake of 0.6–71.4 $\mu\text{g}/\text{day}$, based on a median bisphenol A level of 0.77 ng/mg (0.1–11.9 ng/mg) creatine in a spot urine sample, assumed creatine excretion of 1200 mg/day and that 20% of the dose is excreted in urine. [CERHR recalculated values using a 100% urinary excretion rate which is consistent with human data]	0.01–1.2 based on study author assumptions [0.015 (0.002–0.24) based on a 100% urinary excretion rate]	Ouchi and Watanabe (71)
394 participants in the NHANES III survey (US)	Median (10 th –95 th percentile) 1.32 (0.23–7.95) μg bisphenol A/g creatine in a spot urine sample; [assumed 100% urinary excretion of bisphenol A in 24 hours and creatinine excretion of 1200 mg/day]	[median: 0.026; 10th–95th percentile: 0.0046–0.16]	Calafat et al. (58)
90 girls, 6–8 years-old (US)	Geometric mean \pm SD of 3.0 \pm 3.0 μg bisphenol A /g creatine in sport urine or early morning void samples; [assumed 100% urinary excretion of bisphenol A in 24 hours and creatinine excretion of 1200 mg/day]	[mean 0.059]	Wolff et al. (59)

^aConsistent with estimates conducted by Goodman et al. (55), body weights of 60 kg were assumed, unless otherwise indicated.

^bA 50 kg body weight was assumed.

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1.2.4.2 Occupational exposure

Occupational exposure to bisphenol A could potentially occur during its manufacture, in the production of polycarbonate plastics, and during the manufacture or use of epoxy resins, powder coatings paints, or lacquers (2). Possible exposure to bisphenol A during PVC manufacture has been considered, but the European Union (2) stated that the application was being phased out. According to the European Union, bisphenol A is generally available as granules, flakes, or pellets, thus reducing exposure potential. Bisphenol A is manufactured in closed systems, but exposure is possible during sampling, container filling, and plant maintenance. In the manufacture of polycarbonate, bisphenol A enters the plant and remains in a closed system prior to extrusion. Sampling is conducted by a closed loop system. Following extrusion, the polycarbonate is chopped into granules and bagged, and it is during that stage that exposure to residual bisphenol A (reported at ≤ 100 ppm) through dust is possible. However, it is noted that polycarbonate is stable and that residual bisphenol A is contained within the polymer matrix. The European Union stated that exposure to bisphenol A during the manufacture of polycarbonate items is not likely to exceed values observed during the manufacture of polycarbonate. In the production of epoxy resin, bisphenol A exposure is most likely during reactor charging, but exposure during maintenance is also possible. A residual bisphenol A level of 300 ppm was reported for epoxy resins, but it was noted that most bisphenol A was trapped within the resin matrix. Exposure to bisphenol A during production of epoxy paints is reported to be negligible. In the manufacture of powder epoxy coatings, exposure is thought possible during weighing and milling. Exposure to bisphenol A during the use of powder paints has been documented.

There are no known regulatory limits for occupational exposure to bisphenol A in the US. In 2004, the American Industrial Hygiene Association proposed a workplace environmental exposure level (WEEL) of 5 mg/m^3 for bisphenol A. The draft WEEL was based upon irritation observed in an inhalation toxicity study (77). The value is consistent with the time weighted average (TWA) exposure limits established in Germany and Holland (2).

The European Union (2) summarized occupational exposure data for bisphenol A in Europe and the US. In some cases, industry sources provided data on total inhalable or respirable particulates that were not specifically analyzed for bisphenol A. It was sometimes difficult to discern if values reported in the European Union report were specifically for bisphenol A or for particulates in general. Unless otherwise stated, this report contains only values the Expert Panel was reasonable certain were for bisphenol A. The European Union estimated some exposures through modeling or assumptions of residual monomer levels. Only measured data are summarized in this report. The European Union stated that the values reported did not account for the effects of personal protective equipment in order to avoid difficulties in attempting to quantify protection provided. TWA bisphenol A levels measured in occupational settings are summarized in Table 13. The limited number of values reported indicated that bisphenol A levels were below 5 mg/m^3 . High bisphenol A exposures ($>1 \text{ mg/m}^3$) were observed during spraying of powdered bisphenol A-containing coatings. The highest exposures were observed in the manufacture of bisphenol A and occurred during tanker filling, plant operation activities, and maintenance work. Limited information on short-term exposure to bisphenol is summarized in Table 14. In most cases, bisphenol A values were below 0.1 mg/m^3 ; the highest value of 9.5 mg/m^3 was measured during charging of bags during the manufacture of bisphenol A.

Data for powder paint use summarized in Table 13 were obtained from a NIOSH Health Hazard Evaluation conducted at a company that manufactured fan and ventilation equipment (78). In plant 1 of the company, parts were coated with an epoxy-based powder paint by dipping. At plant 2, an epoxy-based powder was applied to parts via electrostatic spraying. As evident in the data in Table 13, exposures were higher at the plant utilizing electrostatic spraying. Monitoring for bisphenol A was discussed in 2 other NIOSH Health Hazard Evaluation reports. In those reports, bisphenol A was not detected in a plant where an epoxy resin coating was used in the manufacture of electronic resistors (79) or in a plant where an

1 epoxy resin coating was applied to steam turbine generators (80). Rudel et al. (13) used a GC/MS
 2 technique to measure bisphenol A levels at 1 US workplace where plastics were melted and glued; a
 3 concentration of 0.208 $\mu\text{g}/\text{m}^3$ was reported.

4
 5 **Table 13. TWA Measurements of Bisphenol A in the Workplace**

Industry or activity	Location/year	Number of samples	Sample type ^b	8-hour TWA (mg/m^3)
Bisphenol A manufacture	US/not specified	Not specified	Bisphenol A	Not detected (not specified) to 2.6
	Europe/1998	14	Inhalable bisphenol A	<0.5–1.79
	Europe/not specified	10	Inhalable bisphenol A	0.21–1.35
	Europe/1998–2000	8	Bisphenol A	<0.05–0.62
	Europe/1996–1997	5	Inhalable bisphenol A	0.02–0.93
	Europe/not specified	15	Bisphenol A	0.02–2.13
Polycarbonate production	Europe/2000	1	Bisphenol A	<0.001
Powder paint use ^a	US/~1979	7 (3 personal and 4 area samples)	Bisphenol A (plant 1)	0.004–0.006
		21 (15 personal and 6 area samples)	Bisphenol A (plant 2)	0.001–1.063

^aFrom NIOSH (78).

Other data are from the European Union (2).

6
 7
 8 **Table 14. Short-term Bisphenol A Exposures in the Workplace**

Industry	Location/year	Number of samples	Sample type	Exposure level (mg/m^3)
Bisphenol A manufacture	Europe/1990s	18	Inhalable bisphenol A	0.13–9.5
Polycarbonate manufacture	US/not specified	6	Not clear ^a	<0.64 ^a
Epoxy resin manufacture	Europe/not specified	Not known	Bisphenol A	<0.5

^aIt was not clear that the samples were analyzed specifically for bisphenol A or for particulates in general, but the values indicate that if exposure to bisphenol A did occur, it was low. This was the only data available for the US. Source: European Union (2).

9
 10 **[Bisphenol A exposures in US powder paint workers were estimated at ~0.1–100 $\mu\text{g}/\text{kg}$ bw/day based on TWA exposures of 0.001–1.063 mg/m^3 , an inhalation factor of 0.29 m^3/kg day (81), 100% absorption from the respiratory system, and 8 hours worked per day.]**

11
 12
 13
 14 No information was located for dermal exposure to bisphenol A in occupational settings. Using their
 15 Estimation and Assessment of Substance Exposure model, the European Union (2) estimated that dermal
 16 exposure of workers to bisphenol A was unlikely to exceed 5 $\text{mg}/\text{cm}^2/\text{day}$. It was noted that the highest
 17 potential exposure to bisphenol A would occur during bag filling and maintenance work.
 18

1 One study provided information on biological monitoring of bisphenol A in workers exposed to an epoxy
2 compound. In 3 Japanese plants, exposed workers included 42 men who sprayed an epoxy hardening
3 agent consisting of a mixture of bisphenol A diglycidyl ether (10–30%), toluene (0–30%), xylene (0–
4 20%), 2-ethoxyethanol (0–20%), 2-butoxyethanol (0–20%), and methyl isobutyl ketone (0–30%) (82).
5 The workers wore “protection devices” during spraying. Controls consisted of 42 male assembly workers
6 from the same plants who did not use bisphenol A diglycidyl ether. In 1999, urine samples were
7 periodically collected, treated with β -glucuronidase, and examined for bisphenol A by HPLC. Urinary
8 bisphenol A concentrations were significantly higher in exposed workers (median: 1.06 $\mu\text{mol/mol}$
9 creatinine [**2.14 $\mu\text{g/g creatinine}$**]; range: <0.05 pmol to 11.2 $\mu\text{mol/mol creatinine}$ [**<0.1 pg to 22.6 $\mu\text{g/g}$**
10 **creatinine**]) compared to controls (median: 0.52 $\mu\text{mol/mol creatinine}$ [**1.05 $\mu\text{g/g creatinine}$**]; range:
11 <0.05 pmol to 11.0 $\mu\text{mol/mol creatinine}$ [**<0.1 pg to 22.2 $\mu\text{g/g creatinine}$**]). Average difference was
12 reported as 2.5 (95% CI 1.4–4.7; $P = 0.002$). Bisphenol A was not detected in 3 exposed workers and 1
13 control. **[Assuming excretion of 1200 mg/day creatinine (71), mean (ranges) of bisphenol excretion**
14 **in urine were 2.57 $\mu\text{g/day}$ (<0.12 pg to 27.1 $\mu\text{g/day}$) in exposed workers and 1.26 $\mu\text{g/day}$ (<0.12 pg to**
15 **26.6 $\mu\text{g/day}$) in unexposed workers. With an assumed body weight of 60 kg, bisphenol A intake was**
16 **estimated at 0.043 $\mu\text{g/kg bw/day}$ (<0.002 pg to 0.45 $\mu\text{g/kg bw/day}$) in exposed workers and 0.021**
17 **$\mu\text{g/kg bw/day}$ (<0.002 pg to 0.44 $\mu\text{g/kg bw/day}$) in unexposed workers.]**
18

19 1.3 Utility of Data

20 There is a small amount of data available for bisphenol A levels in the environment. Numerous studies
21 reported bisphenol A levels in canned foods and infant formula. Potential levels of bisphenol A in infant
22 formula resulting from leaching of bisphenol A from polycarbonate were estimated. Several estimates of
23 human bisphenol A exposure were developed using bisphenol A levels measured in food and the
24 environment. Although very limited for US populations, there are data reporting bisphenol A levels in
25 urine. There are a limited number of human exposure estimates based on urinary bisphenol A levels. Data
26 for occupational exposure to bisphenol A in the US are very limited. Only 2 studies reported TWA
27 exposures to bisphenol A in US workers. The time period for one study was not reported and the other
28 study was conducted around 1979.

30 1.4 Summary of Human Exposure

31 In 1999 and 2003, it was reported that most bisphenol A produced in the US was used in the manufacture
32 of polycarbonate and epoxy resins and other products (reviewed in (3, 6)). Polycarbonate plastics are used
33 in various consumer products and the products most likely to contribute to human exposure are
34 polycarbonate food containers (e.g., milk, water, and infant bottles). Epoxy resins are used in protective
35 coatings. Food cans lined with epoxy resin are a potential sources of human exposure. Some polymers
36 manufactured with bisphenol A are FDA-approved for use in direct and indirect food additives and in
37 dental materials (8). Resins, polycarbonate plastics, and other products manufactured from bisphenol A
38 can contain trace amounts of residual monomer and additional monomer may be generated during
39 breakdown of polymer (2).

41 Bisphenol A may be present in the environment as a result of direct releases from manufacturing or
42 processing facilities, fugitive emission during processing and handling, or release of unreacted monomer
43 from products (2). Because of its low volatility and relatively short half-life in the atmosphere, bisphenol
44 A is unlikely to enter the atmosphere in large amounts (2). Rapid biodegradation of bisphenol A in water
45 was reported in the majority of studies reviewed by the European Union (2) and Staples et al. (3).
46 Bisphenol A is not expected to be stable, mobile, or bioavailable from soils (10). THE potential for
47 bioconcentration of bisphenol A in fish is low (2, 3). Table 15 summarizes levels of bisphenol A detected
48 in environmental samples and drinking water.
49

1 **Table 15. Summary of Bisphenol A Levels in US Environmental Samples and Drinking Water**

Sample	Bisphenol A Level	Reference
Outdoor air	<52 ng/m ³	Wilson et al. (11, 12)
Indoor air	≤29 ng/m ³	Wilson et al. (11, 12); Rudel et al. (13, 14)
Indoor dust	≤3.24 µg/g	Wilson et al. (11, 12); Rudel et al. (13, 14)
Water bodies	≤12 µg/L	Staples et al. (15); Kolpin et al. (16)
Drinking water from a Louisiana treatment plant	Below quantification limit [not defined]	Boyd et al. (17)

2
3 The highest potential for human exposure to bisphenol A is through products that directly contact food
4 such as food and beverage containers with internal epoxy resin coatings and polycarbonate tableware and
5 bottles, such as those used to feed infants (2). Studies examining the extraction of bisphenol A from
6 polycarbonate bottles or tableware into food simulants are summarized in Table 2. Studies measuring
7 bisphenol A levels in canned infant foods are summarized in Table 3 and studies measuring bisphenol A
8 levels in canned food are summarized in Table 4. Table 16 summarizes the general findings from all the
9 food contact-material studies. Bisphenol A levels were measured in canned foods purchased from various
10 countries. Most of the studies reported that cans were packaged in various locations throughout North
11 America, Europe, or Asia.

13 **Table 16. Summary of Bisphenol A Levels Measured in Foods or Food Simulants**

Exposure Source	Bisphenol A level	Table Reference
Polycarbonate infant bottles	≤50 µg/L	Table 2
Polycarbonate tableware	≤5 µg/kg	Table 2
Canned infant formulas or foods	Generally ≤ 5 µg/kg (up to 113 µg/kg measured in 1 sample from Taiwan)	Table 3
Canned foods	Generally ≤ 100 mg/kg (levels up to ~ 600 mg/kg occasionally detected)	Table 4

14
15 Table 11 summarizes aggregate or food exposure estimates. Estimates of exposure from food range from
16 0.02–8 µg/kg bw/day for infants, 0.00475–14.7 µg/kg bw/day for children, and 0.00195–1.4 µg/kg
17 bw/day for adults. In an aggregate exposure study of US children, it was noted that dietary sources
18 account for 99% of exposure (12).

19
20 Urinary levels of total bisphenol A (free bisphenol A in addition to conjugated bisphenol A metabolites)
21 have been used to estimate human exposures to bisphenol A. Table 6 summarizes urinary levels of total
22 bisphenol A measured in the US, Europe, and Asia. Table 12 summarizes bisphenol A intake estimates
23 based on bisphenol A and metabolite levels in urine. The data suggests that typical daily intakes of
24 bisphenol A are ~0.02–0.06 µg/kg bw/day in adults and children (55, 59).

25
26 Exposure to bisphenol A through dental sealants is considered an acute and rare event (2). Following
27 application of dental sealant, saliva bisphenol A level is typically reported at 0.3–3 ppm [µg/L]. Based on
28 urinary levels of total bisphenol detected following application of dental sealant, exposures to bisphenol
29 A were estimated at 0–239 µg [0–4.0 µg/kg bw based on a 60-kg bw] and were dependent on the type of
30 sealant used (52).

31
32 Very limited information is available for bisphenol A exposure in the US workplace. Data obtained from
33 the US and Europe indicate highest potential exposures during spraying of powdered bisphenol A-

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1 containing coatings and during tanker filling, plant operation activities, and maintenance work in plants
2 where bisphenol A is manufactured. (2). According to limited data summarized in **Table 13**, TWA
3 exposures to bisphenol A were below the draft workplace environmental exposure level (WEEL) of 5
4 mg/m³ proposed by the American Industrial Hygiene Association in 2004 (77). **[Bisphenol A exposures**
5 **in US workers who spray powder paint were estimated at ~0.1–100 µg/kg bw/day based on TWA**
6 **exposures of 0.001–1.063 mg/m³, an inhalation factor of 0.29 m³/kg day (81), 100% absorption from**
7 **the respiratory system, and 8 hours worked per day.]** One study measured total urinary bisphenol A in
8 Japanese workers who sprayed an epoxy compound (82). **[Bisphenol A exposures were estimated at**
9 **0.043 µg/kg bw/day (<0.002 pg to 0.45 µg/kg bw/day) in exposed workers and 0.021 µg/kg bw/day**
10 **(<0.002 pg to 0.44 µg/kg bw/day) in unexposed workers.]**

2.0 GENERAL TOXICOLOGY AND BIOLOGICAL EFFECTS

2.1 Toxicokinetics and metabolism

The studies presented in this section demonstrate that bisphenol A is absorbed in humans and experimental animals following oral exposure. In humans and experimental animals, most of the dose is present in blood as the main metabolite, bisphenol A glucuronide, and smaller percentages are present as the parent compound. Bisphenol A and its metabolites are widely distributed in humans and animals. More than 90% of unmetabolized bisphenol A is reportedly bound to plasma protein. Bisphenol A is distributed to fetal fluids in humans and experimental animals, and a limited number of studies in humans demonstrate fetal levels of bisphenol A within an order of magnitude of levels in maternal blood. None of the studies detected bisphenol A glucuronide in fetal fluids. Transfer of bisphenol A to milk was demonstrated in humans and experimental animals. One study in humans reported bisphenol A in milk at levels exceeding maternal blood levels. In humans and experimental animals, most of a bisphenol A dose is metabolized to bisphenol A glucuronide prior to absorption. In rats, there is evidence that glucuronidation does not occur in fetal liver and is lower in livers of neonates than in older animals. Studies in humans and experimental animals demonstrated that glucuronidation of bisphenol A can occur in the liver, and 1 study in rats demonstrated that bisphenol A is glucuronidated upon passage through the intestine. Bisphenol A glucuronide is excreted in the bile of rats, and enterohepatic cycling is thought to occur in rats but not humans. In humans, most of a bisphenol A dose is eliminated through urine as bisphenol A glucuronide. In rats, bisphenol A is eliminated through feces as bisphenol A and in urine as bisphenol A glucuronide.

2.1.1 Humans

Human toxicokinetics studies that were judged potentially important to interpret developmental and reproductive toxicity were reviewed in full. These studies include reports of potential exposure of fetuses during pregnancy or of infants through human milk and reports of toxicokinetics or metabolism following low-dose exposure of humans. Information from secondary sources was included if the information was not considered to be critical to the interpretation of developmental and reproductive toxicity data.

2.1.1.1 Absorption

Two studies described here examined oral absorption of bisphenol A from dental sealants, and 1 study examined in vitro dermal absorption. Bisphenol A is absorbed in humans as indicated by the detection of bisphenol A in blood from the general population (Section 1) and in maternal and fetal fluids (Table 18).

Fung et al. (50) examined the toxicokinetics of bisphenol A leaching from dental sealant. Volunteers included 18 men and 22 non-pregnant women (ages 20–55 years) who did not have dental disease, existing composite resin restorations or pit and fissure sealants, or a history of resin exposure. Volunteers were treated with a widely used commercial dental sealant (Delton Opaque Light-cure Pit and Fissure Sealant). Components of the sealant were analyzed by HPLC. The low-dose group (n = 7 men, 11 women) received 8 mg dental sealant on 1 tooth, and the high-dose group (11 men, 11 women) received 32 mg sealant on 4 teeth. Saliva and blood samples were collected before the procedure and at 1 and 3 hours and 1, 3, and 5 days after the procedure. Blood and saliva were analyzed by HPLC. Statistical analyses of data were conducted by nonparametric test, Wilcoxon signed rank test, and chi-squared test. Analysis of the dental sealant revealed that bisphenol A levels were below the detection limit of 5 ppb. At 1 hour following treatment, bisphenol A was detected in samples from 3 of the 18 volunteers in the low-dose group and 13 of 22 samples from volunteers in the high-dose group. At 3 hours post-treatment, bisphenol A was detected in samples from 1 of 18 volunteers in the low-dose group and 7 of 22 volunteers in the high-dose group. Levels of bisphenol A in saliva at 1 and 3 hours following exposure were reported at 5.8–105.6 ppb [$\mu\text{g/L}$]. No bisphenol A was detected in saliva samples at 24 hours or in serum samples at any time point. Differences between the low-dose and high-dose groups in bisphenol A

2.0 General Toxicology and Biological Effects

1 saliva levels and in the proportion of bisphenol A-positive saliva samples at 1 and 3 hours achieved
 2 statistical significance. In the high-dose group, a significant difference in “readings” was observed
 3 between 1 and 3 hours. **[The data as presented did not illustrate possible quantitative differences in**
 4 **saliva bisphenol A levels from the 2 dose groups or at different sampling times.]**

5
 6 Joskow et al. (52) examined bisphenol A in urine and saliva of 14 adults (19–42 years old) treated with
 7 dental sealants. Excluded from the study were individuals with resin-based materials on their teeth,
 8 smokers, users of antihistamines, and patients with Gilbert syndrome. The volunteers received either
 9 Heliaseal F (n = 5) or Delton LC (n = 9) sealant. Sealant was weighed before and after application to
 10 determine the amount applied, and the number of treated teeth was recorded. The mean number of teeth
 11 treated was 6/person and the mean total weight of sealant applied was 40.35 mg/person. In a comparison
 12 of the 2 sealants, no differences were reported for number of teeth treated or amount of sealant applied.
 13 Saliva samples were collected prior to treatment, immediately after, and at 1 hour following sealant
 14 application. Urine samples were collected prior to treatment and at 1 and 24 hours following sealant
 15 placement. A total of 14–15 saliva samples and 12–14 urine samples were collected at each time point.
 16 Samples were treated with β -glucuronidase and analyzed for bisphenol A levels using selective and
 17 sensitive isotope-dilution-MS-based methods. Table 17 summarizes changes in saliva and bisphenol A
 18 levels. Immediately and at 1 hour after sealant application, salivary levels of bisphenol A compared to
 19 baseline were significantly higher in the patients who received the Delton LC sealant. Bisphenol A levels
 20 in saliva increased more than 84-fold following application of the Delton LC sealant. Urinary levels of
 21 bisphenol A were increased 1 hour following application of the Delton LC sealant. Levels of bisphenol A
 22 in saliva and urine following application of Heliaseal F were reported to be similar to baseline.

23
 24 **Table 17. Saliva and Urinary Levels of Total Bisphenol A in Adults Receiving Dental Sealants**

Collection time	Mean \pm SD Bisphenol A level (ng/mL) ^a		
	Both sealants	Delton LC	Heliaseal F
<i>Saliva</i>			
Pretreatment	0.30 \pm 0.17	0.34 \pm 0.19	0.22 \pm 0.03
Immediately after treatment	26.5 \pm 30.7	42.8 \pm 28.9	0.54 \pm 0.45
1 hour post-treatment	5.12 \pm 10.7	7.86 \pm 12.73	0.21 \pm 0.03
<i>Urine (creatinine-adjusted)</i>			
Pretreatment	2.41 \pm 1.24	2.6 \pm 1.4	2.12 \pm 0.93
1 hour post-treatment	20.1 \pm 33.1	27.3 \pm 39.1	7.26 \pm 13.5
24 hours post-treatment	5.14 \pm 3.96	7.34 \pm 3.81	2.06 \pm 1.04

^aSamples were treated with β -glucuronidase.
 From Joskow et al. (52).

25
 26 The European Union (2) reviewed unpublished preliminary data from a human dermal absorption study.
 27 Skin samples obtained from 3 human donors (6 samples/donor/dose) were exposed to 5 or 50 mg/cm²
 28 (3.18 or 31.8 mg/mL) ¹⁴C-bisphenol A in ethanol vehicle. Following evaporation of the vehicle, bisphenol
 29 A was resuspended in artificial sweat. Radioactivity was measured in receptor fluid at various time
 30 intervals over a 24-hour period. Radioactivity was measured in the stratum corneum and “lower” skin
 31 layer at 24 hours. Authors of the European Union report noted that tritiated water was not used as a
 32 marker for skin integrity. However, based on the patterns of results, they concluded that skin integrity was
 33 likely lost after 4–8 hours. The European Union authors therefore concluded that the only reliable data
 34 from the study were those for the cumulative percentage of the dose in receptor fluid at 8 hours, which
 35 was reported at 0.57–1.22% at 5 mg/cm² and 0.491–0.835% at 50 mg/cm². Because radioactivity in skin
 36 was not measured at 8 hours, the percentage of the applied dose remaining on skin and available for
 37 future absorption could not be determined. Based on ratios of receptor fluid levels and lower skin levels
 38 (1:2 to 1:8) at 24 hours, and assuming that the higher ratio applies to skin at 8 hours, the authors of the

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1 European Union report predicted that 10% of the dose would be present in “lower” skin layers. Therefore,
2 dermal absorption of bisphenol A was estimated at 10%.

3 2.1.1.2 Distribution

4 In humans, bisphenol A was measured in cord blood and amniotic fluid, demonstrating distribution to the
5 embryo or fetus. Studies reporting bisphenol A levels in fetal and/or maternal compartments are
6 summarized in Table 18. Detailed descriptions of those studies are also presented below.
7

8 **Table 18. Levels of Bisphenol A in Maternal and Fetal Samples**

Study description; analytical method	Bisphenol A concentrations, µg/L, median (range) or mean±SD			Reference
	Serum or plasma		Amniotic fluid	
	Maternal	Fetal		
21 samples collected in women in the US before 20 weeks gestation; LC with electrochemical detection			≤0.5 (≤0.5–1.96)	Engel et al. (83)
37 German women, 32–41 weeks gestation; GC/MS	3.1 (0.3–18.9); 4.4 ± 3.9	2.3 (0.2–9.2); 2.9 ± 2.5		Schönfelder et al. (84)
37 Japanese women in early pregnancy; ELISA ^a	1.5 ± 1.2			Ikezuki et al. (63)
37 Japanese women in late pregnancy; ELISA ^a	1.4 ± 0.9			Ikezuki et al. (63)
32 Japanese infants at delivery; ELISA ^a		2.2 ± 1.8		Ikezuki et al. (63)
32 Japanese amniocentesis samples at 15–18 weeks gestation; ELISA ^a			8.3 ± 8.9	Ikezuki et al. (63)
38 samples obtained at full-term cesarean section; ELISA ^a			1.1 ± 1.0	Ikezuki et al. (63)
200 Japanese women carrying fetuses with normal karyotype at 16 weeks mean gestation; ELISA	2.24 (0.63– 14.36)		0.26 (0–5.62)	Yamada et al. (85)
48 Japanese women carrying fetuses with abnormal karyotypes at a 16 weeks mean gestation; ELISA	2.97 [~0.7–18.5] ^b		0 [~0–7.5] ^b	Yamada et al. (85)
9 sets of maternal and umbilical cord blood samples obtained at birth in Japanese patients; HPLC	0.43 (0.21–0.79)	0.64 (0.45– 0.76)		Kuroda et al. (68)
180 Malaysian newborns; GC/MS		Non- detectable (<0.10) to 4.05		Tan and Mohd (86)

^aSome samples were verified by HPLC.

^bEstimated from a graph.

9
10 Engel et al. (83) reported levels of bisphenol A in human amniotic fluid. Twenty-one samples were
11 obtained during amniocentesis conducted before 20 weeks gestation in women who were referred to a US
12 medical center for advanced maternal age. Bisphenol A concentrations in amniotic fluid were measured
13 using LC with electrochemical detection. Bisphenol A was detected in 10% of samples at levels
14 exceeding the LOD (0.5 µg/L). Bisphenol A concentration ranges of 0.5–1.96 µg/L were reported.
15

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1 Schönfelder et al. (84) examined bisphenol A levels in maternal and fetal blood and compared bisphenol
2 A concentrations in blood of male and female fetuses. In a study conducted at a German medical center,
3 blood samples were obtained from 37 Caucasian women between 32 and 41 weeks gestation. At
4 parturition, blood was collected from the umbilical vein after expulsion of the placenta. Bisphenol A
5 levels in plasma were measured by GC/MS. Control experiments were conducted to verify that bisphenol
6 A did not leach from collection, storage, or testing equipment. Bisphenol A was detected in all samples
7 tested, and concentrations measured in maternal and fetal blood are summarized in Table 18. Mean
8 bisphenol A concentrations were higher in maternal (4.4 ± 3.9 [SD] $\mu\text{g/L}$) than fetal blood (2.9 ± 2.5
9 $\mu\text{g/L}$). Study authors noted that in 14 cases fetal bisphenol A plasma concentrations exceeded those
10 detected in maternal plasma. Among those 14 cases, 12 fetuses were male. Analysis by paired *t*-test
11 revealed significantly higher mean bisphenol A levels in the blood of male than female fetuses (3.5 ± 2.7
12 versus 1.7 ± 1.5 ng/mL, $P = 0.016$). Bisphenol A levels were measured in placenta samples at 1.0–104.9
13 $\mu\text{g/kg}$.

14
15 Ikezuki et al. (63) measured levels of bisphenol A in serum from 30 healthy premenopausal women, 37
16 women in early pregnancy, 37 women in late pregnancy, and 32 umbilical cord blood samples. Levels of
17 bisphenol A were also measured in 32 samples of amniotic fluid obtained during weeks 15–18 of
18 gestation, 38 samples of amniotic fluid obtained at full-term cesarean section, and 36 samples of ovarian
19 follicular fluid collected during in vitro fertilization procedures. **[It was not stated if different samples**
20 **types were obtained from the same subjects.]** An ELISA method was used to measure bisphenol A
21 levels and results were verified by HPLC. The mean \pm SD level of bisphenol A in follicular fluid was
22 reported at 2.4 ± 0.8 $\mu\text{g/L}$. As summarized in Table 5 for nonpregnant women and Table 18 for maternal
23 and fetal samples, levels of bisphenol A in follicular fluid were similar to those detected in the serum of
24 fetuses and pregnant and non-pregnant women and in amniotic fluid collected in late pregnancy (~ 1 – 2
25 $\mu\text{g/L}$). Bisphenol A levels in amniotic fluid samples collected in early pregnancy were ~ 5 -fold higher
26 than in other samples, and the difference achieved statistical significance ($P < 0.0001$). Study authors
27 postulated that the higher levels of bisphenol A in amniotic fluid collected during gestation weeks 15–18
28 may have resulted from immature fetal liver function. They noted that according to unpublished data from
29 their laboratory, the percentage of glucuronidated bisphenol A in mid-term amniotic fluid was $\sim 34\%$,
30 which is much lower than reported values for other human fluids ($>90\%$).

31
32 Yamada et al. (85) measured bisphenol A levels in maternal serum and amniotic fluid from Japanese
33 women. Samples were collected between 1989 and 1998 in women undergoing amniocentesis around
34 gestation week 16. One group of samples was obtained from 200 women carrying fetuses with normal
35 karyotypes, and a second group of samples was obtained from 48 women carrying fetuses with abnormal
36 karyotypes. An ELISA method was used to measure bisphenol A concentrations. Levels of bisphenol A
37 measured in maternal plasma and amniotic fluid are summarized in Table 18. Median levels of bisphenol
38 A in maternal serum (~ 2 – 3 $\mu\text{g/L}$) were significantly higher [**~ 10 -fold**] than levels in amniotic fluid (~ 0 –
39 0.26 $\mu\text{g/L}$) in the groups carrying fetuses with normal and abnormal karyotypes. However, in 8 samples
40 from women carrying fetuses with normal karyotypes, high levels (2.80 – 5.62 $\mu\text{g/L}$) of bisphenol A were
41 measured in amniotic fluid. The study authors interpreted the data as indicating that bisphenol A does not
42 accumulate in amniotic fluid in most cases but accumulation is possible in some individuals. Bisphenol A
43 levels in maternal blood were significantly higher [**by $\sim 33\%$**] in woman carrying fetuses with abnormal
44 versus normal karyotypes. However, the study authors noted that the effect may not be related to
45 bisphenol A exposure because there was no adjustment for maternal age, and levels in amniotic fluid did
46 not differ between groups. In the group carrying fetuses with normal karyotypes, data obtained from 1989
47 to 1998 were summarized by year. Median bisphenol A levels in serum significantly decreased over that
48 time from a level of 5.62 $\mu\text{g/L}$ detected in 1989 to 0.99 $\mu\text{g/L}$ in 1998.

49
50 Kuroda et al. (68) used an HPLC method to measure bisphenol A levels in 9 sets of maternal and cord
51 blood samples obtained from Japanese patients at the time of delivery. Bisphenol A levels were also

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1 measured in 21 sets of serum and ascitic fluid samples collected from sterile Japanese patients of
2 unspecified sexes and ages. Results for pregnant women are summarized in Table 18. Mean \pm SD levels
3 of bisphenol A were lower in maternal (0.46 ± 0.20 ppb [$\mu\text{g/L}$]) than cord blood (0.62 ± 0.13 ppb [$\mu\text{g/L}$]).
4 There was a weak positive correlation ($r = 0.626$) between bisphenol A levels in maternal and cord blood.
5 Levels of bisphenol A in the blood of sterile patients are summarized in Table 5. Mean \pm SD levels of
6 bisphenol A were higher in ascitic fluid (0.56 ± 0.19 ppb [$\mu\text{g/L}$]) than in serum (0.46 ± 0.20 ppb [$\mu\text{g/L}$]).
7 The correlation between bisphenol A level in serum and ascitic fluid was relatively strong ($r = 0.785$).
8

9 Tan and Mohd (86) used a GC/MS method to measure bisphenol A levels in cord blood at delivery in 180
10 patients at a Malaysian medical center. Bisphenol A was detected in 88% of samples. As noted in Table
11 18, levels ranged from <0.10 to 4.05 $\mu\text{g/L}$.
12

13 Calafat et al. (87) reported a median bisphenol A concentration of ~ 1.4 $\mu\text{g/L}$ **[as estimated from a**
14 **graph]** in milk from 32 women who participated in the NHANES III survey. **[No information was**
15 **provided on analytical methods or the form of bisphenol A (e.g., free or total).]**
16

17 Sun et al. (88) used an HPLC method to measure bisphenol A levels in milk from 23 healthy lactating
18 Japanese women. Bisphenol A levels ranged from 0.28 to 0.97 $\mu\text{g/L}$, and the mean \pm SD concentration
19 was reported at 0.61 ± 0.20 $\mu\text{g/L}$. No correlations were observed between bisphenol A and triglyceride
20 levels in milk. Values from 6 milk samples were compared to maternal and umbilical blood samples
21 previously reported in a study by Kuroda et al. (68). Bisphenol A values were higher in milk, and the
22 milk/serum ratio was reported at 1.3. Bisphenol A values in milk were comparable to those in umbilical
23 cord serum. **[It was not clear whether milk and serum samples were obtained from the same**
24 **volunteers in the two studies.]**
25

26 Schaefer et al. (89) measured concentrations of bisphenol A and other compounds in uterine endometrium
27 of women undergoing hysterectomy for uterine myoma at a German medical center. Endometrial and fat
28 samples were obtained between 1995 and 1998 from 23 women (34–51 years old) with no occupational
29 exposure. Samples were handled with plastic-free materials and stored in glass containers. Concentrations
30 of environmental chemicals were measured in samples by GC/MS. None of 21 fat samples had detectable
31 levels of bisphenol A. Bisphenol A was detected in 1 of 23 endometrial samples; the median
32 concentration was reported at <1 $\mu\text{g/kg}$ wet weight, and the range was reported at 0 – 13 $\mu\text{g/kg}$. **[It is not**
33 **known why a median value and range were reported when bisphenol A was only detected in 1**
34 **sample.]**
35

36 As part of a study to compare an ELISA and an LC/MS method for biological monitoring of bisphenol A,
37 Inoue et al. (90) measured levels of bisphenol A in semen samples obtained from 41 healthy Japanese
38 volunteers (18–38 years old). Analysis by the ELISA method indicated bisphenol A levels ranging from
39 levels below the detection limit (2.0 $\mu\text{g/L}$) to 12.0 $\mu\text{g/L}$. The LC/MS method indicated that the bisphenol
40 A level in all samples was <0.5 $\mu\text{g/L}$, the LOQ. The study authors concluded that the LC/MS method was
41 more accurate and sensitive and that the ELISA method overestimated bisphenol A concentrations,
42 possibly due in part to nonspecific antibody interactions.
43

2.1.1.3 Metabolism

45 Völkel et al. (62) measured bisphenol A and metabolite levels in human urine following exposure to a low
46 bisphenol A dose. The human volunteers consisted of 3 healthy females (25–32 years old) and 3 healthy
47 males (37–49 years old) who were asked to refrain from alcohol and medicine intake for 2 days prior to
48 and during the study. Volunteers received 25 μg D_{16} -bisphenol A in drinking water **[0.00028–0.00063**
49 **mg/kg bw based on reported body weights]**, a dose reported to represent a worst-case human exposure.
50 Urine samples were collected at 0, 1, 3, 5, and 7 hours following exposure. Analyses for D_{16} -bisphenol A
51 and D_{16} -bisphenol A-glucuronide were conducted by LC/MS and HPLC. Recovery of D_{16} -bisphenol A-

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1 glucuronide in urine within 5 hours of dosing was 85% of dose in males and 75% of dose in females.
2 Analysis following treatment of urine with glucuronidase resulted in recovery of 97% of the dose in males
3 and 84% of the dose in females. The highest concentrations of bisphenol A glucuronide in urine were
4 measured at 1 hour (221–611 pmol [**50–139 ng bisphenol A eq**]/mg creatine) and 3 hours (117–345 pmol
5 [**27–79 ng bisphenol A eq**]/mg creatinine) following exposure. Elimination half-life was estimated at 4
6 hours. Bisphenol A levels exceeding the detection limit were detected in only 2 urine samples at
7 concentrations of ~10 pmol [**2 ng**]/mg creatinine.

8
9 Völkel et al. (91) examined toxicokinetics and metabolism of bisphenol A in humans administered a low
10 dose. Volunteers in this study consisted of 3 healthy females (24–31 years of age) and 6 healthy males
11 (28–54 years of age) who were non- or occasional smokers; volunteers were asked to refrain from alcohol
12 and medicine intake for 2 days before and during the study. In two different studies, D₁₆-bisphenol A was
13 orally administered to volunteers via gelatin capsules at a dose of 5 mg (0.054–0.090 mg/kg bw). The
14 dose was reported to be ~10-fold higher than the estimated human exposure level of 0.6 mg/day. In the
15 first study, urine samples were collected at 6-hour intervals until 42 hours following exposure and blood
16 samples were collected at 4-hour intervals until 32 hours following exposure in 3 males and 3 females. In
17 a second, more detailed study conducted in 4 of the male volunteers, blood samples were collected at 30–
18 60-minute intervals until 381 minutes following exposure. Samples were analyzed by GC/MS and
19 LC/MS. In the first study, a terminal half-life of 5.3 hours was reported for D₁₆-bisphenol A glucuronide
20 clearance from blood. The half-life for urinary elimination was reported at 5.4 hours. D₁₆-Bisphenol A
21 glucuronide levels in plasma and urine fell below LOD at 24–34 hours post dosing. Complete urinary
22 recovery (100%) was reported for the D₁₆-bisphenol A glucuronide. In the second study, maximum
23 plasma concentration of D₁₆-bisphenol A glucuronide (~800 pmol [**183 ng bisphenol A eq**]/mL) was
24 obtained 80 minutes after oral administration. The half-life for initial decline in plasma was reported at 89
25 minutes. Free D₁₆-bisphenol A was not detected in plasma. According to study authors, the study
26 demonstrated rapid absorption of bisphenol A from the gastrointestinal tract, conjugation with glucuronic
27 acid in the liver, and rapid elimination of the glucuronide in urine. Study authors noted that the rapid and
28 complete excretion of bisphenol A glucuronide in urine suggested that in contrast to rats, enterohepatic
29 circulation did not occur in humans.

30
31 Table 6 in Section 1 provides information on bisphenol A and metabolites detected in human urine. A
32 study conducted in the US used an HPLC method to examine 30 urine samples collected from a
33 demographically diverse adult population in 2000–2004 (69). Mean urinary compound composition was
34 9.5% bisphenol A, 69.5% bisphenol A glucuronide, and 21% bisphenol A sulfate conjugate. A study
35 conducted in Korea used an HPLC method to examine urine collected from 15 men (mean age 42.6 years)
36 and 15 women (mean age 43.0 years) (70). Gender-related differences were observed for urinary
37 metabolic profiles. Mean urinary compound composition in men was reported at 29.1% bisphenol A,
38 66.2% bisphenol A glucuronide, and 4.78% bisphenol sulfate conjugate. The urinary metabolite profile in
39 females was 33.4% bisphenol A, 33.1% bisphenol A glucuronide, and 33.5% bisphenol A sulfate
40 conjugate. The study authors concluded that women had a greater ability for sulfation than men.

41 42 **2.1.1.4 Excretion**

43 As discussed in greater detail in Section 2.1.1.3, two studies in which human volunteers were
44 administered low doses of D₁₆-bisphenol A (~0.00028–0.090 mg/kg bw) demonstrated that most of the
45 dose (85–100%) was eliminated through urine (62) (91). In those studies, the half-lives for urinary
46 elimination were reported at 4–5.4 hours. As discussed in more detail in Section 2.1.1.3, examination of
47 human urine samples revealed that bisphenol A glucuronide and sulfate conjugates are present at higher
48 concentrations than is the parent compound (69, 70).

2.0 General Toxicology and Biological Effects

2.1.2 Experimental animal

Original animal studies that were potentially important for the interpretation of developmental and reproductive toxicity were reviewed thoroughly. Examples included:

- Studies examining toxicokinetics or metabolism in pregnant or lactating animals
- Studies examining toxicokinetic difference observed with different doses or exposure routes
- Studies looking at age-related differences in toxicokinetics or metabolism
- Studies in non-rodent species such as primates

Secondary sources were utilized for general information not considered critical to the interpretation of developmental and reproductive toxicity data.

2.1.2.1 Absorption

In rats orally exposed to bisphenol A at doses ≤ 100 mg/kg bw, maximum bisphenol A concentrations (C_{\max}) were generally measured in plasma within 0.083–0.75 hours following exposure (92, 93, 94, 95, 96). At doses of 1 or 10 mg/kg bw, time to maximum bisphenol A concentration (T_{\max}) in plasma was longer in postnatal day (PND) 21 rats (1.5–3 hours) than in PND 4 and 7 rats (0.25–0.75 hours) (92). In a limited number of studies in which rats were subcutaneously (sc) dosed with up to 100 mg/kg bw bisphenol A, time (0.5–4 hours) to reach C_{\max} was longer than with oral dosing, although the findings were not always consistent (93,96). In one study, T_{\max} was comparable in oral and intraperitoneal (ip) dosing of rats (93). Another study reported that C_{\max} was attained at 0.7 hours in monkeys orally exposed to 10 or 100 mg/kg bw bisphenol A and at 0.5 hours in chimpanzees orally exposed to 10 mg/kg bw bisphenol A (96). In the same study, a longer T_{\max} (2 hours) was observed following exposure of monkeys and chimpanzees to the same doses by sc injection compared to oral intake. Additional details for these studies are presented below.

As discussed in greater detail in Section 2.1.2.3, bisphenol A is glucuronidated in the liver and intestine, and most of the dose is absorbed as bisphenol A glucuronide following oral exposure of rats (92). In ovariectomized rats gavaged with bisphenol A, bioavailability of bisphenol A was reported at 16.4% at a 10 mg/kg bw dose and 5.6% at a 100 mg/kg bw dose (97). The findings are fairly consistent with a second study in which maximum plasma values of free bisphenol A represented low percentages [**<2–8%**] of the total radioactive dose in rats orally administered bisphenol A at 10 or 100 mg/kg bw (93); maximum values of free bisphenol A represented higher percentages of the radioactive dose in rats given 10 or 100 mg/kg bw sc [**64–82% free bisphenol A**] or ip [**19–54%**] (93). Percentages of parent bisphenol A in blood were also higher in monkeys exposed intravenously (iv; 5–29%) than orally (0–1%) (98). Similarly, HPLC analysis of plasma conducted 1 hour following sc or gavage dosing of 4 female 21-day-old Sprague Dawley rats/group with bisphenol A revealed higher bisphenol A plasma concentrations with sc than with gavage dosing (Table 19) (99). One study in male and female rats gavaged with 10 mg/kg bw bisphenol A demonstrated higher plasma levels of bisphenol A in immature animals than in adults (10.2–48.3 $\mu\text{g/g}$ [**mg/L**] plasma at 4 days of age; 1.1–1.4 $\mu\text{g/g}$ [**mg/L**] plasma at 7 days of age; 0.2 $\mu\text{g/g}$ [**mg/L**] plasma at 21 days of age; and 0.024–0.063 $\mu\text{g/g}$ [**mg/L**] plasma in adulthood) (92).

1 **Table 19. Plasma Bisphenol A Concentrations in 21-day-old Rats at 1 Hour Following Oral Gavage**
 2 **or SC Dosing**

Dose, mg/kg bw	Plasma concentration, µg/L	
	sc injection	Oral gavage
0 (sesame oil vehicle)	Not detected	Not detected
8	94.6 ± 58.0	Not examined
40	886.3 ± 56.4	Not detected
160	2948 ± 768.8	198.8 ± 88.2
800	Not examined	2879.0 ± 2328.3

Values presented as mean ± SD.

From: (99).

3
 4 A review by the European Union (2) noted that in the study by Pottenger et al. (93), fecal excretion
 5 represented the highest proportion of the eliminated dose (74–83% in males and 52–72% in females)
 6 following oral or parenteral exposure of rats to 10 or 100 mg/kg bw bisphenol A. The authors of the
 7 European Union report therefore concluded that absorption [**assumed to be of the radioactive dose**] is
 8 likely extensive following oral intake. Adding to the proof of extensive oral absorption is the observation
 9 that more than 50% of fecal elimination occurred at 24 hours post dosing, a time period beyond the
 10 average gastrointestinal transit time of 12–18 hours for rats. Possible explanations provided for the
 11 detection of parent compound in feces were cleavage of conjugates within intestines and enterohepatic
 12 circulation.

13 14 2.1.2.2 Distribution

15 16 2.1.2.2.1 Pregnant or lactating animals

17 Information on distribution in pregnant or lactating rats is presented first followed by other species.
 18 Studies including oral exposures are summarized before those with parenteral exposures.

19
 20 Takahashi and Oishi (94) examined disposition and placental transfer of bisphenol A in F344 rats. Rats
 21 were orally administered 1000 mg/kg bw bisphenol A (>95% purity) in propylene glycol on gestation day
 22 (GD) 18 (GD 0 = day of vaginal plug). Rats were killed at various time points between 10 minutes and 48
 23 hours after bisphenol A dosing. At each time point, 2–6 dams and 8–12 fetuses obtained from 2–3 dams
 24 were analyzed. Blood was collected from dams and kidneys, livers, and fetuses were removed for
 25 measurement of bisphenol A levels by HPLC. Results are summarized in Table 20. Study authors noted
 26 the rapid appearance of bisphenol A in maternal blood and organs and in fetuses. Levels of bisphenol A at
 27 6 hours following dosing were 2% of peak levels in maternal blood and 5% of peak levels in fetuses. It
 28 was noted that in fetuses, area under the time-concentration curve (AUC) was higher and mean retention
 29 time, variance of retention time, and terminal half-life were longer than in maternal blood.

30

1 **Table 20. Toxicokinetic Endpoints for Bisphenol A in Rats Dosed with 1000 mg/kg bw Bisphenol A**
 2 **on GD 18**

Endpoint	Maternal tissue			Fetus
	Blood	Liver	Kidney	
C _{max} , mg/L	14.7	171	36.2	9.22
T _{max} , minutes	20	20	20	20
AUC, mg·hour/L	13.1	700	84.0	22.6
Mean retention time, hours	10.6	29.3	12.0	20.0
Variance in retention time, hours squared	203	657	227	419
Half-life, hours				
From 20 to 40 minutes	0.0952	0.178	0.245	0.55
From 40 minutes to 6 hours	2.58	1.75	2.98	1.60
From 6 to 48 hours	4.65	No data	No data	173

From Takahashi and Oishi (94).

3
 4 Dormoradzki et al. (100) examined metabolism, toxicokinetics, and embryo-fetal distribution of bisphenol
 5 A in rats during 3 different gestation stages. Sprague Dawley rats were gavaged with bisphenol A (99.7%
 6 purity)/radiolabeled ¹⁴C-bisphenol A (98.8% radiochemical purity) at 10 mg/kg bw. Bisphenol A was
 7 administered to 1 group of non-pregnant rats and 3 different groups of pregnant rats on GD 6 (early
 8 gestation), 14 (mid gestation), or 17 (late gestation). GD 0 was defined as the day that sperm or a vaginal
 9 plug were detected. Blood, urine, and feces were collected at multiple time points between 0.25 and 96
 10 hours post dosing. It appears that most and possibly all samples were pooled. Four rats in each group
 11 were killed at 96 hours post dosing. Maternal organs, 6 embryos or fetuses/dam (when possible), and
 12 placentas were collected. Samples were analyzed for radioactivity and bisphenol A and/or bisphenol A
 13 glucuronide by HPLC/liquid scintillation spectrometry.

14
 15 In all groups, 90–94% of radioactivity was recovered. Elimination of bisphenol A and its metabolites is
 16 discussed in Section 2.1.2.4. At 96 hours following dosing, low percentages of the dose were present in
 17 carcass (~1–6%) and tissues such as brain, fat, liver, kidney, ovary, uterus, and skin. The only
 18 quantifiable data in placentas and fetuses at 96 hours were obtained in the GD 17 group, and those
 19 samples contained 0.01–0.07% of the bisphenol A dose. Standard deviations for maternal and fetal tissues
 20 generally exceeded 50% of the mean. Study authors concluded that disposition of radioactivity was
 21 similar in pregnant and non-pregnant rats.

22
 23 Toxicokinetic data obtained from plasma profiles are summarized in Table 21. The authors stated that
 24 there was high inter-animal variability. The presence of 2 C_{max} values was noted by the authors, and they
 25 stated that it was the result of enterohepatic circulation of radioactivity. Bisphenol A was not quantifiable
 26 in most plasma samples. Because bisphenol A glucuronide represented most (~95–99%) of the
 27 radioactivity, plasma profiles for that metabolite were nearly identical to profiles for radioactivity.

28

1 **Table 21. Toxicokinetic Data for Radioactivity in Pregnant and Non-pregnant Rats Gavaged with**
 2 **10 mg/kg bw ¹⁴C-bisphenol A**

Endpoint	Non-pregnant	GD 6–10	GD 14–18	GD 17–21
C _{max1} , mg eq/L	0.716	0.370	0.482	1.006
T _{max1} , hours	0.25	0.25	0.25	0.25
C _{max2} , mg eq/L	0.171	0.336	0.211	0.278
T _{max2} , hours	18	12	24	12
Time to non-quantifiable level, hours	72	Not determined	72	96
AUC				
¹⁴ C, mg-eq·hour/L	6.1	12.4	7.1	10.2
Bisphenol A glucuronide, mg-eq·hour/L	5.8	12.3	6.8	9.7
Percent as bisphenol A glucuronide	95.1	99.2	95.8	95.1

From Dormoradzki et al. (100).

3
 4 A second study was conducted by Dormoradzki et al. (100) to measure bisphenol A and bisphenol A
 5 glucuronide levels in maternal and fetal tissues. Rats were gavaged with radiolabeled bisphenol A at 10
 6 mg/kg bw on GD 11, 13, or 16. Blood was collected over a 24-hour period. Five rats/group/time period
 7 were killed at 0.25, 12, and 96 hours post dosing. Maternal blood and organs, yolk sacs/placentas, and
 8 embryos/fetuses were removed for measurement of bisphenol A and bisphenol A glucuronide. Yolk
 9 sacs/placentas and fetuses were pooled at most time periods. Results are summarized in Table 22.

10
 11 At 0.25 hours following dosing, bisphenol A glucuronide levels in maternal plasma were similar in
 12 groups dosed on GD 11 and 13 but levels were 1.7–2 times higher in the group dosed on GD 16. At 12
 13 hours post dosing in all exposure groups, bisphenol A glucuronide levels in maternal plasma were
 14 reduced 7- to 11-fold from values observed at 0.25 hours. Levels of radioactivity in plasma were not
 15 sufficient for analysis at 96 hours post dosing. Bisphenol A was detected in maternal plasma at 0.25 hours
 16 post dosing in rats that were exposed to a higher radioactive concentration (0.5 mCi compared to 0.2 mCi)
 17 on GD 16; bisphenol A levels were 26.5-fold lower than bisphenol A glucuronide levels.

18
 19 In animals dosed on GD 11, bisphenol A glucuronide was only detected in yolk sac/placenta at 0.25 hours
 20 post dosing and the concentration was ~17 times lower than the level detected in maternal blood for the
 21 same time period. With dosing on GD 11, bisphenol A glucuronide was not detected in embryos and
 22 bisphenol A was not detected in yolk sac/placenta or embryos. In animals dosed on GD 13, bisphenol A
 23 glucuronide was detected in yolk sac/placenta at 0.25 and 12 hours post dosing and levels were 9–24-fold
 24 lower than those detected in maternal plasma for the same time period. Bisphenol A was also detected in
 25 yolk sac/placenta at 0.25 and 12 hours after dosing on GD 13 and levels were similar to those detected in
 26 the blood of 2 dams. A lower level of bisphenol A was detected in embryos of dams at 0.25 hours
 27 following dosing on GD 13, and bisphenol A was the only moiety detected in embryos. Following dosing
 28 on GD 16, bisphenol A glucuronide and bisphenol A were detected in yolk sac/placenta at 0.25 and 12
 29 hours post dosing. Levels of bisphenol A glucuronide in yolk sac/placenta were 7- to 8-fold lower than
 30 levels detected in maternal plasma. From 0.25 to 12 hours, levels of bisphenol A decreased 4.9-fold and
 31 levels of bisphenol A glucuronide decreased 9-fold. Mean levels of bisphenol A in yolk/sac placenta
 32 following exposure on GD 16 were similar to the blood level detected in 1 of 2 dams.

33
 34 In yolk sac/placenta and fetuses of dams dosed with a higher level of radioactivity (0.5 mCi) on GD 16,
 35 bisphenol A glucuronide and bisphenol A were detected at 0.25 hours following dosing. Compared to
 36 concentrations detected in placenta, fetal levels of bisphenol A glucuronide were ~26-fold lower and
 37 bisphenol A levels were 5-fold lower. Bisphenol A levels were lower than bisphenol A glucuronide levels
 38 by 3.6-fold in yolk sac/placenta and by 0.7-fold in fetuses. Study authors concluded that there is no
 39 selective affinity for bisphenol A or bisphenol A glucuronide by the yolk sac/placenta or embryo/fetus.

Table 22. Maternal and Fetal Levels of Bisphenol A Following Gavage Dosing of Dams with 10 mg/kg bw Bisphenol A

Exposure	Bisphenol A level, mg/L or mg/kg					
	Maternal plasma		Yolk sac/placenta		Embryo/fetus	
	Glucuronide	Parent	Glucuronide	Parent	Glucuronide	Parent
GD 11, 0.2 mCi						
0.25 hours	1.060 ± 0.258	0.041	0.062	<LOD ^a	<LOD	<LOD
12 hours	0.099 ± 0.036	<LOD	<LOD	<LOD	<LOD	<LOD
96 hours	NA	NA	<LOD	<LOD	<LOD	<LOD
GD 13, 0.2 mCi						
0.25 hours	0.868 ± 0.189	0.078	0.036	0.019	<LOD	0.013
12 hours	0.117 ± 0.033	0.008	0.013	0.009	<LOD	<LOD
96	Not analyzed due to insufficient radioactivity					
GD 16, 0.2 mCi						
0.25 hours	1.768 ± 0.783	0.485, 0.129	0.223 ± 0.104	0.166 ± 0.069	0.031, 0.009	0.122, 0.020
12 hours	0.174 ± 0.045	<LOD	0.025 ± 0.005	0.034 ± 0.002	NA	NA
96 hours	Not analyzed due to insufficient radioactivity				0.016	0.008
GD 16, 0.5 mCi						
0.25 hours	1.699 ± 0.501	0.064 ± 0.025	0.342 ± 0.104	0.095 ±0.031	0.013 ± 0.008	0.018 ± 0.011

Data expressed as mean ± SD or single values for individual or pooled data.

^aLimit of detection (LOD) for bisphenol A reported at 0.005–0.029.

From Dormoradzki et al. (100).

Kurebayashi et al. (101) examined distribution of radioactivity in pregnant and lactating rats dosed with ¹⁴C-bisphenol A. Pregnant rats were orally dosed with 0.5 mg/kg bw ¹⁴C-bisphenol A on GD 12, 15, or 18. The rats were killed at 30 minutes or 24 hours following dosing (n = 1/time period) and examined by whole-body radioluminography. Study authors noted that the distribution of label was nearly identical in dams at each gestation time point. At 30 minutes following dosing, the concentration of radioactivity in dam blood was ~31–43 µg bisphenol A eq/L. The highest concentration of radioactivity was detected in maternal liver (~219–317 µg bisphenol A eq/kg) and kidney (~138–270 µg bisphenol A eq/kg); concentrations in other tissues (lung, ovary, placenta, skin, and uterus) were ~10-fold lower. Fetuses, fetal membranes, and yolk sacs did not contain quantifiable levels of radioactivity at 30 minutes following maternal exposure at any gestation time point. At 24 hours following exposure of dams, radioactivity concentrations in blood (~4–11 µg bisphenol A eq/L) were nearly 3–10-fold lower than values obtained at 30 minutes following exposure. Levels of radioactivity remained highest in liver. At 24 hours following exposure, radioactivity was only detected in fetuses and fetal tissues from dams dosed on GD 18. Radioactivity levels in fetuses or fetal tissues compared to maternal blood were ~30% in fetuses, nearly equal in fetal membranes, and ~5-fold higher in yolk sacs. Study authors concluded that there was limited distribution of radiolabel to fetuses.

In another study by Kurebayashi et al. (101), a lactating rat was orally dosed with 0.5 mg/kg bw ¹⁴C-bisphenol A on PND 11 and caged with 5 neonatal rats for 24 hours. One male and one female neonatal rat were killed at the end of the 24-hour period and examined by whole-body radioluminography. The 3 remaining neonates were caged for 24 hours with a dam that was not exposed to bisphenol A. One male and one female neonate were then killed and examined by whole-body radioluminography. In pups killed immediately after being nursed by the lactating dam exposed to ¹⁴C-bisphenol A, most of the radioactivity was detected in intestinal contents (~30–46 µg bisphenol A eq/kg) and lower levels were found in gastric

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1 contents and urinary bladder (< 10 µg bisphenol A eq/kg). After being nursed for 24 hours by a dam that
 2 was not exposed to bisphenol A, radioactivity was only detected in intestinal contents and the level was
 3 ~20–40% of that measured in pups examined immediately after being nursed by dams receiving ¹⁴C-
 4 bisphenol A.

5
 6 An additional 3 lactating dams were dosed with 0.5 mg/kg bw ¹⁴C-bisphenol A on PND 11 for
 7 examination of radioactivity in plasma and milk over a 48-hour period. Table 23 summarizes
 8 toxicokinetic endpoints for radioactivity in milk and plasma. Study authors concluded that there was
 9 significant secretion of ¹⁴C-associated radioactivity into milk.

10
 11 **Table 23. Toxicokinetic Endpoints for Radioactivity in Lactating Rats Orally Administered 0.5**
 12 **mg/kg bw ¹⁴C-bisphenol A on PND 11**

Endpoint	Milk	Maternal plasma
C _{max} , µg-eq/L	4.46	27.2
T _{max} , hours	8	4
Elimination half-life, hours	26	31
AUC (0–48 hours), µg-eq·hour/L	156	689

From Kurebayashi et al. (101).

13
 14 Miyakoda et al. (102) examined placental transfer of bisphenol A in rats. Wistar rats were administered an
 15 oral dose of bisphenol A (99% purity) at 10 mg/kg bw on GD 19. Blood was collected and fetuses were
 16 removed at 1, 3, and 24 hours following dosing. Bisphenol A levels were measured in plasma and fetuses
 17 by GC/MS. **[A statement in Figure 3 of the study indicated that values were the means of 5 or 7**
 18 **experiments; it is possible the authors meant that 5 or 7 dams were dosed.]** Concentrations of
 19 bisphenol A peaked in maternal plasma and fetuses within 1 hour of dosing, with bisphenol A levels
 20 measured at ~34 ppb [µg/L] in maternal plasma and 11 ppb [µg/kg] in fetuses. At 3 hours after dosing,
 21 bisphenol A levels were ~10% of peak levels in maternal plasma and 40% of peak levels in fetuses. At 24
 22 hours post dosing, bisphenol A levels in fetuses were detected at 70% of peak value and levels in fetuses
 23 were more than twice the levels in maternal plasma. Study authors concluded that bisphenol A is rapidly
 24 transferred to the fetus and tends to remain longer in fetuses than in maternal blood.

25
 26 Snyder et al. (103) examined the toxicokinetics of bisphenol A in lactating rats. On PND 14, lactating CD
 27 rats were gavaged with 100 mg/kg bw ¹⁴C-bisphenol A. Milk, blood, and organs were collected from 2–4
 28 dams/group at 1, 8, 24, or 26 hours after dosing. **[While the text indicates collection of samples at 26**
 29 **hours, Table 3 of the study indicates collection at 24 hours. The collection time reported in the**
 30 **study table was used when there were discrepancies between text and table.]** Animals were injected
 31 with oxytocin prior to milk collection. Radioactivity in pup carcasses was measured at 2, 4, 6, and 24
 32 hours following exposure of dams; 8–16 pups/time period were examined. Samples were analyzed by
 33 scintillation counting, HPLC, and/or nuclear magnetic resonance. At 1 and 8 hours following exposure,
 34 the highest percentage of the radioactive dose was detected in intestine with contents (75–83%). Among
 35 the other organs examined, the highest percentage of the radioactive dose was detected in liver (0.38–
 36 0.74%) and much lower percentages were detected in kidney and lung (≤0.02%). Low percentages of the
 37 radioactive dose were also detected in milk (≤0.0020%), blood (~0.006%), plasma (~0.01%), and fat
 38 (≤0.004%). Compared to earlier time periods, radioactivity levels were lower at 24 hours post dosing
 39 (26% of the dose detected in intestine and contents), but distribution was similar. At all 3 sampling time
 40 points, radioactivity levels were highest in plasma > blood > milk. The major radioactivity peak in plasma
 41 was represented by bisphenol A glucuronide at 1, 8, and 26 hours following exposure. Bisphenol A
 42 glucuronide also represented the major radioactive peak detected in milk. Radioactivity levels in pups
 43 amounted to <0.01% of the maternal dose. Radioactivity levels in pups tended to increase over time.

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1 From 2 to 24 hours following exposure, mean \pm SD radioactivity levels rose from 44 ± 24 to 78 ± 11 μg
 2 bisphenol A eq/pup.

3
 4 Yoshida et al. (104) compared bisphenol A levels in rats and their offspring during the lactation period.
 5 The main focus of the study was developmental toxicity, which is discussed in Section 3.2.3.2. In the
 6 distribution study, Donryu rats (12–19/group) were gavaged with bisphenol A at 0
 7 (carboxymethylcellulose solution), 0.006, or 6 mg/kg bw/day from GD 2 to the day before weaning (21
 8 days post-delivery). Bisphenol A levels were measured in maternal and pup serum, milk, and pup liver by
 9 GC/MS on PND 10, 14, and/or 21. Milk samples were obtained from pup stomachs. Pup serum and liver
 10 samples were pooled. Two to six dams/litters were examined in each dose group and time period.
 11 Samples of tap water, drinking water from plastic containers, and feed were measured for bisphenol A
 12 content by HPLC. Bisphenol A was not detected in fresh tap water but was detected at ~ 3 $\mu\text{g/L}$ following
 13 storage of that water in plastic containers. Bisphenol A concentration in feed was ~ 40 $\mu\text{g/kg}$. Results for
 14 maternal and fetal tissues are summarized in Table 24. Bisphenol A levels in the serum of high dose-dams
 15 were significantly elevated compared to the control group on PND 21. No other significant differences
 16 were observed in bisphenol A levels in samples between treated and control groups.

17
 18 **Table 24. Bisphenol A Concentrations in Maternal and Pup Samples During Lactation in Rats**
 19 **Gavaged with Bisphenol A**

Sample	Time of analysis	Sex	Dose group, mg/kg bw/day			
			0	0.006	6	
Bisphenol A concentration, ppb [$\mu\text{g/L}$ or $\mu\text{g/kg}$]						
<i>Dam^a</i>						
Serum	PND 21		3 ± 0	4 ± 0	11 ± 4	
Milk	PND 10		28 ± 9	8 ± 21	8 ± 3	
	PND 14		255 ± 78	205 ± 7	185 ± 50	
<i>Pup^b</i>						
Serum	PND 10	Female	4	10	23	
		Male	15	5	7	
	PND 14	Female	5	4	3	
		Male	4	5	4	
	PND 21	Female	9	3	9	
		Male	14	9	20	
	Liver	PND 10	Female	13	12	17
			Male	9	9	14
		PND 14	Female	22	100	18
			Male	45	14	16
		PND 21	Female	60	70	37
			Male	69	9	60

^aValues are presented as mean \pm SD.

^bPup samples were pooled.

From Yoshida et al. (104).

20
 21 Kim et al. (105) used an HPLC method to measure bisphenol A levels in rat dams and their offspring.
 22 Dams were gavaged with bisphenol A (>99.7% purity) at doses of 0 (corn oil vehicle), 0.002, 0.020,
 23 0.200, 2, or 20 mg/kg bw/day on GD 7–17. Dams and offspring were killed at 21 days following
 24 parturition, and serum was collected for measurement of bisphenol A. Development effects observed in
 25 this study are summarized in Section 3.2.1.1. Bisphenol A was not detected in the serum of dams at the
 26 two lowest doses. Respective concentrations of bisphenol A in the serum of dams at the 3 highest doses
 27 were 0.900, 0.987, and 1.00 mg/L. In offspring, bisphenol A was not detected in serum at the 3 lowest

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1 doses. At the 2 highest doses, the respective concentrations of bisphenol A in offspring were 0.69 and
2 0.74 mg/L in males and 0.71 and 0.82 mg/L in females.

3
4 Shin et al. (106) examined elimination of bisphenol A from maternal-fetal compartments of rats. On 1 day
5 between GD 17 and 19, four Sprague Dawley rats were iv injected with 2 mg/kg bw bisphenol A.
6 Amniotic fluid, placenta, and fetuses were collected at multiple intervals between 5 minutes and 8 hours
7 following injection. Bisphenol A concentrations in samples were measured by HPLC. Transfer rate
8 constants and clearance rates were determined using a 5-compartment model consisting of maternal
9 central, maternal tissue, placental, fetal, and amniotic fluid compartments. Toxicokinetic findings are
10 summarized in Table 25. Rapid distribution of bisphenol A was observed in placenta, fetus, and amniotic
11 fluid. Bisphenol A levels in placenta and fetus remained higher than those in maternal serum over most of
12 the sampling period. Amniotic fluid contained the lowest level of bisphenol A. Decay curves in amniotic
13 fluid, fetus, and placenta paralleled decay curves in maternal serum. Transfer rate constants and clearance
14 rates are summarized in Table 26. Transfer rate constants were greater in the direction of amniotic fluid to
15 fetus or placenta than in the opposite direction. The elimination rate constant and clearance rate from the
16 fetal compartment were much lower than for the maternal central compartment. The clearance rate from
17 placenta to fetus was higher than clearance rate from fetus to placenta. The authors calculated that 65.4%
18 of the bisphenol A dose was delivered to the fetus, 33.2% to the maternal central compartment, and 1.4%
19 to amniotic fluid. According to the study authors, the low transfer rate from the fetal to amniotic
20 compartment suggested minimal fetal excretion of unchanged bisphenol A through urine and feces into
21 the amniotic fluid. They also noted that the small fetal compartment transfer constant compared to the
22 relative fetal-placental transfer constant indicated minimal metabolism by the fetus. Authors estimated
23 that 100% of bisphenol A was eliminated from the fetus via the placental route and concluded that fetal
24 elimination represents 0.05% of total elimination from the maternal-fetal unit.

25
26 **Table 25. Toxicokinetic Endpoints for Bisphenol A in Pregnant Rats iv Dosed with 2 mg/kg bw**
27 **Bisphenol A**

Endpoint	Compartment			
	Maternal serum	Placenta	Fetus	Amniotic fluid
AUC, $\mu\text{g}\cdot\text{hour}/\text{L}$	905.5 \pm 275.8	4009 \pm 962.7	1964.7 \pm 678.5	180.4 \pm 102.0
Elimination half-life, hours	2.5 \pm 0.9	2.2 \pm 0.8	2.2 \pm 0.8	3.9 \pm 3.1
Mean residence time, hours	3.0 \pm 1.1	2.0 \pm 0.5	3.0 \pm 0.9	5.6 \pm 4.7
C_{max} , $\mu\text{g}/\text{L}$	927.3 \pm 194.3	1399.2 \pm 323.7	794 \pm 360.6	75.1 \pm 59.7
T_{max} , hours	No data	0.1 \pm 0.1	0.6 \pm 0.3	0.3 \pm 0.2

Values presented as mean \pm SD. From Shin et al. (106).

28
29 **Table 26. Intercompartmental Transfer and Clearances in Pregnant Rats After iv Bisphenol A**

Compartment	Transfer rate constant, hour^{-1}	Clearance rate, mL/minute
Maternal central to maternal tissue	3.4 \pm 2.6	38.2 \pm 26.5
Maternal tissue to maternal central	1.7 \pm 1.3	50.2 \pm 36.7
Maternal central to placental	0.7 \pm 0.5	8.3 \pm 5.4
Placental to maternal central	23.6 \pm 14.7	2.2 \pm 1.3
Placental to fetal	46.4 \pm 29.2	4.1 \pm 2.1
Fetal to placental	22.8 \pm 28.0	7.6 \pm 6.0
Fetal to amniotic fluid	0.00001 \pm 0.00002	0.00001 \pm 0.00001
Fetal	0.0062 \pm 0.0044	0.0024 \pm 0.0015
Amniotic fluid to fetal	14.0 \pm 21.0	0.8 \pm 1.1
Amniotic fluid to placental	7.9 \pm 6.7	0.7 \pm 0.7
Placental to amniotic fluid	1.0 \pm 1.3	0.1 \pm 0.1
Maternal central	0.9 \pm 0.6	9.7 \pm 5.3

Values presented as mean \pm SD. From Shin et al. (106)

2.0 General Toxicology and Biological Effects

1
2 Yoo et al. (95) examined mammary excretion of bisphenol A in rats. At 4–6 days postpartum, 4–6
3 lactating female Sprague Dawley rats/group were iv injected with bisphenol A at 0.47, 0.94, or 1.88
4 mg/kg bw and then infused with bisphenol A over a 4-hour period at rates of 0.13, 0.27, or 0.54 mg/hour.
5 Blood samples were collected at 2, 3, and 4 hours, and milk was collected at 4 hours following initiation
6 of infusion. Prior to collection of milk, rats were injected with oxytocin to increase milk production.
7 HPLC was used to measure bisphenol A levels in serum. Differences in data for mean systemic clearance
8 were analyzed by analysis of variance (ANOVA). Results are summarized in Table 27. The study authors
9 noted extensive excretion of bisphenol A into milk, with milk levels exceeding serum levels. No
10 significant differences were reported for systemic clearance rates between the 3 doses. Steady state
11 concentrations of bisphenol A in maternal serum and milk increased linearly according to dose.
12

13 **Table 27. Toxicokinetic Endpoints in Lactating Rats Infused with Bisphenol A**

Endpoint	Bisphenol A infusion rate, mg/hour		
	0.13	0.27	0.54
Systemic clearance, mL/minute/kg	119.2 ± 23.8	142.4 ± 45.3	154.1 ± 44.6
Steady state serum bisphenol A concentration, ng/mL	66.1 ± 15.5	120.0 ± 34.7	217.1 ± 65.0
Steady state milk bisphenol A concentration, ng/mL	173.1 ± 43.3	317.4 ± 154.4	493.9 ± 142.2
Milk/serum ratio	2.7 ± 0.9	2.6 ± 1.2	2.4 ± 0.6

Data presented as mean ± SD.

From Yoo et al. (95).

14
15 Kabuto et al. (107) reported bisphenol A concentrations in mice indirectly exposed to bisphenol A during
16 gestation and lactation. The focus of the study was oxidative stress; more details are presented in Section
17 3.2.7. Six ICR mouse dams were given drinking water containing 1% ethanol vehicle or bisphenol A at 5
18 or 10 µg/L. **[Based on the reported water intake of 5 mL/day and an assumed body weight of 0.02 kg**
19 **(81), it is estimated that bisphenol A intakes in mice at the start of pregnancy were 0.0013 and**
20 **0.0025 mg/kg bw/day.]** Mice gave birth about 3 weeks following mating and pups were housed with
21 dams for 4 weeks. **[Based on an assumed body weight of 0.0085 kg and assumed water intake rate of**
22 **0.003 L/day (81), it is estimated that intake of bisphenol A in weanling males was 0.0018 and 0.0035**
23 **mg/kg bw/day.]** At 4 weeks of age, male pups were killed and a GC/MS technique was used to measure
24 bisphenol A levels in brain, kidney, liver, and testis in an unspecified number of control pups and in 4
25 pups from the 10 µg/L group. Study authors reported that they could not detect bisphenol A in control
26 pups. In pups from the 10 µg/L group, the highest concentration of bisphenol A was detected in kidney
27 (~24 µg/kg wet weight), followed by testis (~20 µg/kg wet weight), brain (~18 µg/kg wet weight), and
28 liver (~11 µg/kg wet weight).
29

30 Zalko et al. (108) examined metabolism and distribution of bisphenol A in pregnant CD-1 mice. A series
31 of studies was conducted in which mice were treated with ³H-bisphenol A (>99.9% purity)/unlabeled
32 bisphenol A (>99% purity). Mice were exposed to different regimens; biological samples examined
33 included blood, liver, fat, gall bladder, uterus, ovaries, digestive tract and contents, urine, and feces. In the
34 first exposure scenario, mice were sc injected with 0.025 mg/kg bw labeled/unlabeled bisphenol A on GD
35 17; three animals/time period were examined at 0.5, 2, and 24 hours following dosing. In the second
36 exposure scenario, 2 mice/group were sc injected with 50 mg/kg bw bisphenol A on GD 17 and killed 24
37 hours following dosing. In the third scenario, 3 non-pregnant female mice/group were “force-fed” a single
38 oral dose of 0.025 mg/kg bw bisphenol A; urine and feces were collected over 24 hours, and animals were
39 killed at 24 hours. Biological samples were analyzed by scintillation analysis, HPLC, MS, and/or nuclear
40 magnetic resonance.
41

2.0 General Toxicology and Biological Effects

1 In pregnant mice injected with 0.025 mg/kg bw/day bisphenol A and examined 24 hours later, 85.68% of
 2 the radioactivity was recovered. The highest percentages of radioactivity were detected in the digestive
 3 tract and contents (~45%) and feces (~21%). Less radioactivity was detected in the litter (~4%), liver
 4 (~2%), bile (~2%), urine (~6%), and carcass (~3%). Blood, ovaries, uterus, placenta, amniotic fluid, fat,
 5 and cage washes each contained <1% of the radioactive dose. At 0.5 hours following dosing, levels of
 6 radioactivity were highest in uterus > liver > placenta > fetus > amniotic fluid > ovaries > carcass >
 7 blood. Radioactivity levels in tissues were lower by 8 and 24 hours following exposure. **[Compared to**
 8 **radioactive levels detected in tissues at 24 hours, levels detected at 0.5 hours were ~12-fold higher in**
 9 **uterus, 3-fold higher in liver, 8-fold higher in placenta, 3.5-fold higher in fetuses, 2-fold higher in**
 10 **amniotic fluid, and 3.5-fold higher in ovaries.]** The only information provided for mice sc dosed with
 11 50 mg/kg bw bisphenol A and examined 24 hours later was for radioactivity levels in organs; the highest
 12 levels were detected in liver > uterus > amniotic fluid > fetuses > ovary. Study authors stated that
 13 distribution of radioactivity was comparable in mice treated with 50 and 0.025 mg/kg bw bisphenol A. In
 14 the mice orally dosed with 0.025 mg/kg bw bisphenol A and examined 24 hours later, levels of
 15 radioactivity in blood, ovaries, and uterus were reported to be significantly lower **[by ~1–2 orders of**
 16 **magnitude]** than levels in animals exposed by sc injection, but the level in the liver was not significantly
 17 different. No qualitative differences in metabolites were observed following oral or sc exposure. **[Data**
 18 **wer not shown by study authors.]** Distribution of parent compound and metabolites detected in maternal
 19 and fetal tissues is summarized in Table 28. Further discussion on metabolites is included in Section
 20 2.1.2.3.
 21

22 **Table 28. Qualitative Analysis of Maternal and Fetal Tissues Following Injection of Mice with 0.025**
 23 **mg/kg bw Radiolabeled Bisphenol A on GD 17**

Hours after dose	Percentage of bisphenol A-associated compound detected					
	Hydroxylated glucuronide	Double glucuronide	Metabolite F ^a	Glucuronide	Parent	Others
Maternal plasma						
0.5	3	4	4	39	41	9
2	2	4	4	63	17	10
24	20	0	0	65	0	15
Placenta						
0.5	0	0	2	25	72	1
2	1	1	7	62	26	3
24	5	4	59	19	6	6
Fetus						
0.5	1	0	5	44	49	1
2	1	0	13	66	16	3
24	1	0	13	60	15	2
Amniotic fluid						
0.5	1	2	1	83	9	4
2	1	1	5	88	2	2
24	8	1	24	44	2	20
Maternal liver						
0.5	0	0	18	37	31	12
2	1	8	20	45	13	13
24	2	7	16	36	23	17

^aMost likely bisphenol A glucuronide conjugated to acetylated galactosamine or glucosamine.
 From Zalko et al. (108).

24
 25 Uchida et al. (109) examined distribution of bisphenol A in pregnant mice and monkeys. On GD 17 (GD
 26 0 = day of vaginal plug), ICR mice were sc injected with bisphenol A 100 mg/kg bw in sesame oil

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1 vehicle. More than 3 mice/time point were killed at various points between 0.5–24 hours following
2 injection. An untreated control group consisted of 6 mice. **[Data were not presented for controls.]**
3 Maternal and fetal serum and organs were collected. Among organs collected were fetal uteri and testes,
4 which were pooled. On GD 150, 2 Japanese monkeys (*Macaca fuscata*) were sc injected with 50 mg
5 bisphenol A/kg bw and at 1 hour following injection, fetuses were removed by cesarean section. Two
6 untreated fetuses were used as controls. Maternal and fetal serum and organs, not including reproductive
7 organs, were collected from monkeys. Bisphenol A concentrations were measured by GC/MS in mouse
8 and monkey samples.

9
10 In mice, bisphenol A was detected within 0.5 hours of exposure in all tissues examined, including
11 placenta; maternal and fetal serum, liver, and brain; and fetal uterus and testis. Bisphenol A
12 concentrations were higher in fetal than maternal serum and liver. **[Peak concentrations were observed**
13 **within 0.5–1 hour in most tissues, with the exception of fetal brain (2 hours), and levels remained**
14 **elevated for 1–6 hours, depending on tissue. More than 1 peak was observed in fetal serum, uterus,**
15 **and testis.]** In exposed monkeys, bisphenol A was found at the highest concentrations (15.6–72.50
16 mg/kg) in fetal heart, intestine, liver, spleen, kidney, thymus, muscle, cerebrum, pons, and cerebellum;
17 bisphenol A levels in the same organs from control monkeys were measured at 3.70–22.80 mg/kg. Lower
18 levels of bisphenol A were detected in umbilical cord and maternal and fetal serum of the exposed group
19 (1.70–6.10 mg/kg) and control group (0.02–0.25 mg/kg). The study authors stated that the most likely
20 source of bisphenol A in control monkeys was the feed, which was found to contain bisphenol A. The
21 study authors concluded that the placental barrier does not protect the fetus from bisphenol A exposure.

2.1.2.2.2 Non-pregnant and non-lactating animals

22
23 Domoradzki et al. (92), examined the effects of dose and age on toxicokinetics and metabolism of
24 bisphenol A in rats. Neonatal and adult male and female Sprague Dawley rats were gavaged with ¹⁴C-
25 bisphenol A (~99% radiochemical purity)/non-radiolabeled bisphenol A (99.7% purity). Three neonatal
26 rats/age/sex/time period were dosed on PND 4, 7, and 21 with 1 or 10 mg/kg bw bisphenol A. Adult rats
27 (11 weeks old) **[number treated not specified]** were dosed with 10 mg/kg bw bisphenol A. Blood
28 samples were collected at various time points from 0.25 to 24 hours post dosing in neonatal rats and from
29 0.25 to 96 hours in adult rats. Plasma samples were pooled on PND 4. Immature rats were killed at 24
30 hours post-dosing, and adult rats were killed at 96 hours post dosing. Brain, liver, kidneys, skin, and
31 reproductive organs were collected from neonatal rats. Levels of radioactivity, bisphenol A, and/or
32 metabolites were analyzed in blood and tissue samples using HPLC and liquid scintillation spectrometry.

33
34
35 In neonatal and adult rats, radioactivity levels in plasma generally peaked within 0.25–0.75 hours. With
36 the exception of 0.25 hours post dosing on PND 4, when plasma radioactivity levels were ~4-fold higher
37 in males than females, plasma radioactivity levels were generally similar in male and female rats. At 24
38 hours post dosing, plasma radioactivity levels were 4–100 times lower in all groups of neonatal rats.
39 Trends were noted for decreasing radioactivity levels with increasing age. Information related to dose-
40 and age-related effects on metabolism is presented in Section 2.1.2.3.

41
42 Toxicokinetic values for bisphenol A are listed in Table 29. C_{max} and AUC values for bisphenol A
43 decreased with increasing age, especially following dosing with 10 mg/kg bw. Bisphenol A
44 concentrations were lower in adults than neonates. No patterns were observed for half-lives, and the
45 authors stated that values in neonates may not have been reliable because bisphenol A concentrations
46 were near the LOD at the end of the 24-hour observation period. Ratios of C_{max} and AUC values for the
47 10 and 1 mg/kg bw doses were different at each age and generally decreased with age. Plasma bisphenol
48 A levels were very low in adults dosed with 10 mg/kg bw; therefore, few data were available.

49
50 Toxicokinetic values for bisphenol A glucuronide are listed in Table 30. Peak plasma concentrations of
51 bisphenol A glucuronide were 9–22 times higher in neonates than adult rats dosed with 10 mg/kg bw

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1 bisphenol A. AUC values for bisphenol A glucuronide were also higher in neonates than adults [**~2–6**
 2 **times higher**]. In neonates dosed with 1 mg/kg bw, AUC values and elimination half-lives for bisphenol
 3 A glucuronide decreased with age. Ratios of C_{max} and AUC values for the 10 and 1 mg/kg bw doses were
 4 nearly proportional. In adults dosed with 10 mg/kg bw, bisphenol A glucuronide levels peaked at 0.25
 5 hours and secondary peaks were observed at 18 and 24 hours. In neonates dosed with 10 mg/kg bw, levels
 6 of bisphenol A glucuronide peaked at 0.75–1.5 hours and then bisphenol A glucuronide was eliminated in
 7 an apparently monophasic manner. Half-lives of elimination were shorter in neonates compared to adults.
 8 In neonatal rats, the bisphenol A glucuronide represented 94–100% of the 1 mg/kg bw dose and 71–97%
 9 of the 10 mg/kg bw/day dose. In adult rats, ~100% of the dose was represented by bisphenol A
 10 glucuronide.

11
 12 Half-life and AUC data for bisphenol A-derived radioactivity in organs of neonatal rats are summarized
 13 in Table 31. Radioactivity was distributed to all organs and dose-related increases were observed. The
 14 study authors noted lower concentrations in brain than in other tissues. [**Levels of radioactivity in**
 15 **reproductive organs compared to those in plasma varied at each evaluation period but were usually**
 16 **within the same or 1 order of magnitude lower.**] With the exception of males dosed with 10 mg/kg bw
 17 bisphenol A, half-lives decreased with age. There were some disproportionate increases in ratios of AUC
 18 at 10 and 1 mg/kg bw.

19
 20 The study authors concluded:

- 21 • Metabolism of bisphenol A to its glucuronide conjugate occurred as early as PND 4 in rats,
- 22 • Dose-dependent differences occurred in neonatal rats, as noted by a larger fraction of the lower
 23 dose being metabolized to the glucuronide, and
- 24 • There were no major sex differences in metabolism or toxicokinetics of bisphenol A.

25
 26 **Table 29. Toxicokinetic Values for Bisphenol A in Rats Following Gavage Dosing with 1 or 10**
 27 **mg/kg bw**

Endpoint	Age at exposure and sex							
	PND 4		PND 7		PND 21		Adult	
	Male	Female	Male	Female	Male	Female	Male	Female
Bisphenol A Dose: 1 mg/kg bw								
T_{max} , hours	0.25	0.25	0.25	0.25	3	3		
C_{max} , mg/L	0.03	0.06	0.04	0.08	0.005	0.006		
Half-life, hours	7.2	7.3	21.8	8.8				
AUC, mg·hour/L	0.2	0.1	0.1	0.1				
Bisphenol A Dose: 10 mg/kg bw								
T_{max} , hours	0.25	0.25	0.25	0.25	1.5	1.5	0.25	0.75
C_{max} , mg/L	48.3	10.2	1.1	1.4	0.2	0.2	0.024	0.063
Half-life, hours	17	6.7	11.4	8.5	4.3	6.6	“0”	“0”
AUC, mg·hour/L	23.1	7.2	1.9	1.7	1.1	1	“0”	“0”
Ratio of value at 10 to 1 mg/kg bw/day								
C_{max}	1610	170	27.5	17.5				
AUC	115.2	72	19	17				

Data missing from table cells were not determined.

From Domoradzki et al. (92).

28
 29

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1 **Table 30. Toxicokinetic Values for Bisphenol A Glucuronide in Rats following Gavage Dosing with**
 2 **1 or 10 mg/kg bw Bisphenol A**

Endpoint	Age at exposure and sex							
	PND 4		PND 7		PND 21		Adult	
	Male	Female	Male	Female	Male	Female	Male	Female
Bisphenol A dose: 1 mg/kg bw								
T _{max} , hours	0.75	0.75	0.75	0.25	0.25	0.25		
C _{max} , mg/L	1.3	1.5	2	1.1	0.8	0.8		
Half-life, hours	26.1	24.2	6.6	6.4	4.2	4.1		
AUC, mg·hour/L	9	9.6	7.7	7.7	4.1	3.3		
AUC _{BPA-glucuronide} /AUC _{BPA}	45	96	77	77				
Bisphenol A dose: 10 mg/kg bw								
T _{max} , hours	1.5	1.5	1.5	0.75	0.75	0.75	0.25	0.25
C _{max} , mg/L	13.1	6.3	6.6	10.3	10.4	7.8	0.6	0.7
Half-life, hours	7.3	9.8	9.1	8.4	4.4	4.4	22.5	10.8
AUC, mg·hour/L	80	50.3	58.9	60.9	60.3	56.1	31.5	9.8
AUC _{BPA-glucuronide} /AUC _{BPA}	3.5	7	31	36	55	56		
Ratio of value at 10 to 1 mg/kg bw/day								
C _{max}	10.1	4.2	3.3	9.4	13	9.8		
AUC	8.9	5.2	7.6	7.9	14.7	17		

Data missing from table cells were not determined.
 From Domoradzki et al. (92).

3
 4 **Table 31. Distribution of Radioactivity to Tissues at 24 Hours Following Dosing with Radiolabeled**
 5 **Bisphenol A**

Tissue	PND 4			PND 7			PND 21		
	Half-life, hours	AUC, mg·h/kg	AUC ratio of doses	Half-life, hours	AUC, mg·h/kg	AUC ratio of doses	Half-life, hours	AUC, mg·h/kg	AUC ratio of doses
<i>Females, 1 mg/kg bw</i>									
Brain	11.7	0.4		6.7	0.2		3.6	0.1	
Liver	18	7.5		7.9	7.1		3.6	2.9	
Kidney	18.1	9.4		7.3	9.5		5.0	3.0	
Ovary	11.7	7.3		6.0	3.5		3.7	0.9	
Uterus	7.4	8.3		6.2	3.0		3.4	1.0	
Carcass	11.2	22.2		10.0	16.6		4.0	8.3	
Plasma	19.5	9.4		6.4	7.8		3.6	3.5	
<i>Females, 10 mg/kg bw</i>									
Brain	7.2	3.3	8.3	8.0	2.5	12.5	4.9	1.7	17.0
Liver	11.1	44.8	6.0	10.0	59.6	8.4	4.5	39.1	13.5
Kidney	15.2	43.9	4.7	8.6	66.6	7.0	5.3	36.5	12.2
Ovary	6.5	136.2	18.7	5.0	69.7	19.9	3.6	21.1	23.4
Uterus	15.2	127.0	15.3	4.8	108.5	36.2	3.4	30.6	30.6
Carcass	6.6	112.8	5.1	7.0	130.7	7.9	4.8	100.9	12.2
Plasma	9.2	61.0	6.5	8.1	67.0	8.6	3.7	59.0	16.9
<i>Males, 1 mg/kg bw</i>									
Brain	14.1	0.3		6.0	0.3		3.4	0.1	
Liver	19.7	6.1		6.6	7.3		3.7	3.2	
Kidney	19.3	8.5		7.0	8.6		4.6	3.4	
Testis	10.3	3.4		5.7	2.0		3.4	0.8	
Carcass	11.1	22.2		9.0	17.3		4.1	9.0	

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Tissue	PND 4			PND 7			PND 21		
	Half-life, hours	AUC, mg·h/kg	AUC ratio of doses	Half-life, hours	AUC, mg·h/kg	AUC ratio of doses	Half-life, hours	AUC, mg·h/kg	AUC ratio of doses
Plasma	24.0	9.2		6.6	7.7		3.4	4.2	
<i>Males, 10 mg/kg bw</i>									
Brain	3.1	4.7	15.7	8.0	2.9	9.7	4.7	1.7	17.0
Liver	11.6	48.4	7.9	11.8	62.0	8.5	5.1	40.9	12.8
Kidney	5.4	68.9	8.1	9.8	59.6	6.9	6.9	30.4	8.9
Testes	5.8	36.8	10.8	7.6	22.1	11.1	5.2	8.1	10.1
Carcass	8.3	111.7	5.0	8.6	135.5	7.8	4.8	95.2	10.6
Plasma	6.9	113.0	12.3	9.9	69.0	9.0	4.0	62.0	14.8

From Domoradzki et al. (92).

1
2 Pottenger et al. (93) examined the effects of dose and route on metabolism and toxicokinetics of
3 bisphenol A in rats. Information focusing on toxicokinetics is primarily summarized in this section, while
4 metabolic data are primarily summarized in Section 2.1.2.3. Adult male and female F344 rats were dosed
5 with ¹⁴C-bisphenol A (99.3% radiochemical purity)/non-radiolabeled bisphenol A (99.7% purity) at doses
6 of 10 or 100 mg/kg bw by oral gavage or ip or sc injection. Blood was collected at multiple time points
7 between 0.083 and 168 hours post dosing, and excreta were collected for 7 days. Animals were killed 7
8 days post dosing. Blood, brain, gonads, kidneys, liver, fat, skin, uterus, and carcass were analyzed by
9 liquid scintillation counting and HPLC. Some samples were analyzed by HPLC/electrospray
10 ionization/MS.

11
12 Toxicokinetic endpoints for bisphenol A in blood are summarized in Table 32. Study authors noted that
13 concentration-time profiles of bisphenol were dependent on dose, exposure route, and sex. The longest
14 T_{max} was observed with sc dosing. C_{max} and AUC values were lowest following oral administration. Time
15 to non-quantifiable levels of bisphenol A was longest following sc exposure. The only sex-related
16 difference was a higher C_{max} value in females than males following oral dosing. In most cases, bisphenol
17 A toxicokinetics were linear across doses within the same administration route, as noted by approximate
18 proportionate increases in C_{max} and AUC values from the low to the high dose. Toxicokinetics data for
19 radioactivity in plasma are summarized in Table 33. Levels of radioactivity were dependent on exposure
20 route and to a lesser extent, dose and sex. AUC values for radioactivity were lowest following oral
21 exposure. Time to non-quantifiable concentration was longest following sc dosing. For most groups, C_{max}
22 and AUC values were proportionate across doses within the same exposure route. A second part of the
23 study examined metabolites and is summarized in Section 2.1.2.3.

24

2.0 General Toxicology and Biological Effects

1 **Table 32. Toxicokinetic Endpoints for Bisphenol A in Blood Following Dosing of Rats by Gavage or**
 2 **Injection**

Endpoint	Exposure route and doses (mg/kg bw)					
	10 oral	100 oral	10 ip	100 ip	10 sc	100 sc
<i>Males</i>						
T _{max} , hours	N/A	0.083	0.5	0.25	0.75	0.5
C _{max} , mg/L, hours ^a	^b	0.22 ± 0.09	0.69 ± 0.08	9.7 ± 1.27	0.39 ± 0.16	5.19 ± 0.98
Time to non-quantifiable level, hours	0.083	0.75	8	12	18	24
AUC, mg·hour/L		0.1	1.1	16.4	2.6	24.5
<i>Females</i>						
T _{max} , hours	0.25	0.25	0.25	0.25	4	0.75
C _{max} , mg/L, hours ^a	0.04 ± 0.03	2.29 ± 1.82	0.87 ± 0.15	13.13 ± 4.13	0.34 ± 0.06	3.97 ± 0.6
Time to non-quantifiable level, hours	1		24	72	48	72
AUC, mg·hour/L	0.42	4.4	1.4	26.2	3.1	31.5

^aMean ± SD.

^bNon-quantifiable (0.01 µg/g at 10 mg/kg bw and 0.1 µg/g at 100 mg/kg bw).

Missing values were not determined.

From Pottenger et al. (93).

3
 4 **Table 33. Toxicokinetics for Radioactivity Following Dosing of Rats with Bisphenol A through**
 5 **Different Exposure Routes**

Endpoint	Exposure route and doses (mg/kg bw)					
	10 oral	100 oral	10 ip	100 ip	10 sc	100 sc
<i>Males</i>						
T _{max} , hours	0.25	0.25	0.5	0.25	1	0.75
C _{max} , mg eq/L, hours	0.73 ± 0.22	3.92 ± 1.93	1.26 ± 0.09	29.3 ± 11.7	0.61 ± 0.24	6.33 ± 0.43
Time to non-quantifiable level, hours	72	72	96	96	96	144
AUC, mg·eq-hour/L	8.1	66.5	16.9	170	15.5	218
<i>Females</i>						
T _{max} , hours	0.083	0.25	0.25	0.5	0.75	0.75
C _{max} , mg eq/L, hours	1.82 ± 0.66	28.33 ± 8.64	2.27 ± 0.19	67.81 ± 7.33	0.52 ± 0.06	5.66 ± 0.95
Time to non-quantifiable level, hours	72	72	72	120	120	168
AUC, mg·eq-hour/L	9.54	94.9	15.3	247	21.6	297

From Pottenger et al. (93).

6
 7 Upmeier et al. (97) examined toxicokinetics in rats exposed to bisphenol A through the oral or iv route.
 8 Ovariectomized DA/Han rats (130–150 g bw) were exposed to bisphenol A by iv injection with 10 mg/kg
 9 bw or oral gavage with 10 or 100 mg/kg bw. Blood was collected from treated rats at multiple time points
 10 until 2 hours following iv dosing and 3 hours following oral dosing. The number of rats sampled during
 11 each time period was 3–5. To reduce stress, only some of the rats were sampled at each time point. In
 12 control animals, blood was collected 2 hours following dosing with vehicle. Bisphenol A levels in plasma
 13 were measured by GC/MS. Dosing with 10 mg/kg bw iv resulted in a maximum plasma level of 15,000
 14 µg/L bisphenol A. Concentrations rapidly decreased to 700 µg/L within 1 hour, 100 µg/L within 2 hours,

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1 and non-detectable concentrations by 24 hours following exposure. The apparent final elimination half-
 2 life was estimated at 38.5 hours. In rats gavaged with 10 mg/kg bw, an initial maximum blood level of 30
 3 $\mu\text{g/L}$ was obtained at 1.5 hours. A decrease in bisphenol A blood level at 2.5 hours was followed by a
 4 second peak of 40 $\mu\text{g/L}$ at 6 hours, leading study authors to conclude that enterohepatic cycling was
 5 occurring. The same patterns of bisphenol A levels in blood were observed following gavage dosing with
 6 100 mg/kg bw. Peak levels were observed at 30 minutes (150 $\mu\text{g/L}$) and 3 hours (134 $\mu\text{g/L}$) following
 7 exposure. According to the study authors, the differences in peak levels observed between the 2 doses
 8 suggested lower bioavailability at the high dose than at the low dose. Oral bioavailability of bisphenol A
 9 was estimated at 16.4% at the low dose and 5.6% at the high dose.

10
 11 Yoo et al. (95) examined toxicokinetics of a low iv dose and a higher gavage dose of bisphenol A in male
 12 rats. Five adult male Sprague Dawley rats/group were administered bisphenol A by iv injection at a dose
 13 of 0.1 mg/kg bw or by gavage at a dose of 10 mg/kg bw. Multiple blood samples were collected until 3
 14 hours following iv dosing and 24 hours following gavage dosing. HPLC was used to measure bisphenol A
 15 levels in serum. Route-specific differences in mean systemic clearance were analyzed by Student *t*-test.
 16 Results are summarized in Table 34. The study authors noted bi-exponential decay of serum bisphenol A
 17 levels following iv dosing, significantly longer elimination half-life with oral than iv exposure, and low
 18 oral bioavailability of bisphenol A.

19
 20 **Table 34. Toxicokinetic Values for Bisphenol A in Adult Rats Exposed to Bisphenol A through the**
 21 **IV or Oral Route**

Endpoint	Bisphenol A dosing	
	0.1 mg/kg bw, iv	10 mg/kg bw, gavage
Distribution half-life, minutes	6.1 \pm 1.3	
Terminal elimination half-life, hours	0.9 \pm 0.3	21.3 \pm 7.4
AUC, $\mu\text{g}\cdot\text{hour/L}$	16.1 \pm 3.2	85.6 \pm 33.7
Systemic clearance, mL/minute/kg	107.9 \pm 28.7	
Steady-state volume of distribution, L/kg	5.6 \pm 2.4	
C_{max} , $\mu\text{g/L}$		14.7 \pm 10.9
T_{max} , hours		0.2 \pm 0.2
Apparent volume of distribution, L/kg		4273 \pm 2007.3
Oral clearance, mL/minute/kg		2352.1 \pm 944.7
Absolute oral bioavailability, %		5.3 \pm 2.1

Data presented as mean \pm SD.

From Yoo et al. (95).

22
 23 Kurebayashi et al. (110) conducted a series of studies to examine toxicokinetics and metabolism of
 24 bisphenol A in adult F344N rats exposed through the oral or iv route. In these studies, radioactivity levels
 25 were measured by scintillation counting. Bisphenol A or its metabolites were quantified by HPLC,
 26 electrospray ionization/ MS, or nuclear magnetic resonance. As discussed in greater detail in Section
 27 2.1.2.4, fecal excretion was the main route of elimination for radioactivity following oral or iv dosing of
 28 rats with 0.1 mg/kg bw ^{14}C -bisphenol A. A study describing biliary excretion and metabolites in bile is
 29 summarized in Section 2.1.2.3. Toxicokinetic endpoints were determined in a study in which blood was
 30 drawn from 3 male rats/group at various time points between 0.25 and 48 hours following oral gavage or
 31 iv dosing with 0.1 mg/kg bw bisphenol A. Results of the study are summarized in Table 35. Rapid
 32 absorption of radioactivity was observed following oral dosing. AUC values were significantly lower for
 33 oral than iv dosing. In a another study, rats were administered ^{14}C -bisphenol A by iv injection and blood
 34 was collected 30 minutes later for determination of blood/plasma distribution and protein binding. At a
 35 blood radioactivity level of 80 nM [**18 μg bisphenol A eq/L**], preferential distribution to plasma was
 36 observed, with the blood/plasma ratio reported at 0.67. At radioactivity levels of 6–31 $\mu\text{g}\cdot\text{eq/L}$ (27–135
 37 nM), plasma protein binding was reported at 95.4%. Additional studies reviewed by Teeguarden et al.

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1 (111) reported plasma protein binding of bisphenol A at ~90–95%. An additional study by Kurebayashi et
 2 al. (110) compared metabolic patterns and excretion following exposure to a higher bisphenol A dose;
 3 that study is discussed in Section 2.1.2.3.

4
 5 **Table 35. Toxicokinetic Endpoints for ¹⁴C-Bisphenol A-Derived Radioactivity in Rats Exposed to**
 6 **0.1 mg/kg bw ¹⁴C-Bisphenol A Through the Oral or IV Route**

Endpoint	IV exposure	Oral exposure
T _{max} , hour		0.38 ± 0.10
C _{max} , µg-eq/L		5.5 ± 0.3
Half-life-α, hours	0.59 ± 0.09	No data
Half-life-β, hours	39.5 ± 2.1	44.5 ± 4.1
Absorbance rate, hour ⁻¹		3.6 ± 1.0
Volume of distribution, L/kg	27.0 ± 0.7	No data
Total body clearance, L/hour/kg	0.522 ± 0.011	0.544 ± 0.049
Mean residence time, hour	51.7 ± 2.4	No data
AUC, µg-eq·hour/L		
0–6 hours	33.9 ± 1.6	18.4 ± 0.7 ^a
0–24 hours	79.3 ± 3.3	60.0 ± 7.1 ^a
0–48 hours	118 ± 4	102 ± 13 ^a
0–∞	192 ± 4	185 ± 16
Oral bioavailability ^b		
0–6 hours		0.54
0–24 hours		0.76
0–48 hours		0.86
0–∞		0.97

Data presented as mean ± SD.

Missing values are not applicable or were not reported.

^aP < 0.05 compared to iv exposure.

^bVariances not reported.

From Kurebayashi et al. (110).

7
 8 Kurebayashi et al. (101) administered ¹⁴C-bisphenol A to adult male and female F344 rats (3/dose/sex) at
 9 doses of 0.020, 0.1, or 0.5 mg/kg bw orally or 0.1 or 0.5 mg/kg bw by iv injection. Plasma samples were
 10 analyzed for radioactivity over a 72-hour period to determine toxicokinetic endpoints. Results are
 11 summarized in Table 36. Study authors noted that the AUC was almost linearly correlated with dose.
 12 Several peaks were observed with oral or iv exposure, indicating enterohepatic cycling, according to the
 13 study authors. Study authors noted that substantially lower AUC values in females than in males
 14 following oral exposure could have resulted from lower absorption and/or a higher elimination rate.
 15 Distribution of radioactivity was evaluated 0.5, 24, and 72 hours following oral administration of 0.1
 16 mg/kg bw bisphenol A to adult male and female Wistar rats (3/sex/time point). At 0.5 hours following
 17 exposure, most of the radioactivity (~12–51 µg bisphenol A eq/kg) was found in kidney and liver. **[A**
 18 **large amount of radioactivity was also reported for intestinal contents, but those data were not**
 19 **shown by the study authors.]** Lower amounts of radioactivity (~2–7 µg bisphenol A eq/kg or L) were
 20 detected in adrenal gland, blood, lung, pituitary gland, skin, and thyroid gland of both sexes; uterus; and
 21 bone marrow, brown fat, and mandibular gland of males. In males, <1 µg bisphenol A eq/kg was detected
 22 in skeletal muscle and testis. Radioactivity was non-quantifiable in brain and eye of both sexes;
 23 epididymis, prostate gland, and heart of males; and bone marrow, brown fat, skeletal muscle, and
 24 mandibular gland of females. At ≥24 hours following exposure, radioactivity was primarily detected only
 25 in kidney, liver, and intestinal contents, with the exception of ~3 µg bisphenol A eq/L detected in blood of
 26 males at 24 hours following dosing. Study authors noted that elimination of radioactivity from some

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1 tissues appeared to occur more rapidly in females than in males. Distribution in pregnant animals was also
 2 examined and is described in Section 2.1.2.2.1.

3
 4 **Table 36. Toxicokinetic Endpoints for Plasma Radioactivity in Rats Dosed with ¹⁴C Bisphenol A**

Endpoints	Route and dose (mg/kg bw)				
	Oral			IV	
	20	100	500	100	500
<i>Males</i>					
Elimination half-life, hours	78 ± 52	18 ± 3	21 ± 3	19 ± 2	21 ± 3
AUC, µg·eq·h/L	36 ± 6	178 ± 44	663 ± 164	266 ± 46	865 ± 97
Apparent absorption, %	82	81	60		
<i>Females</i>					
Elimination half-life, hours	20 ± 7	22 ± 13	18 ± 8	13 ± 3	16 ± 2
AUC, µg·eq·h/L	14 ± 5	99 ± 19	500 ± 43	190 ± 45	1029 ± 81
Apparent absorption, %	35	50	50		

Data presented as mean ± SD.

From: Kurebayashi et al. (101).

5
 6 Kabuto et al. (112) reported distribution of bisphenol A in mice. Male ICR mice were ip dosed with
 7 bisphenol A at 0, 25, or 50 mg/kg bw/day for 5 days and killed 6 hours following the last dose. Bisphenol
 8 A levels in tissues of animals from the high-dose group were determined by GC/MS. In mice of the high-
 9 dose group, the highest concentrations of bisphenol A were detected in kidney (~2.02 mg/kg wet weight)
 10 and body fat (~1.25 mg/kg wet weight). Lower concentrations of bisphenol A (≤0.42 mg/kg wet weight or
 11 mg/L) were detected in brain, lung, liver, testis, and plasma.

12
 13 Kurebayashi et al. (98) examined the toxicokinetics of a low bisphenol A dose in cynomolgus monkeys
 14 following gavage or iv dosing. Three adult male and female monkeys were dosed with 0.1 mg/kg bw ¹⁴C-
 15 bisphenol A (99% radiochemical purity)/non-radiolabeled bisphenol A [**purity not reported**]. Monkeys
 16 were dosed by iv injection on day 1 of the study and by gavage on day 15 of the study. Urine and feces
 17 were collected for 7 days post dosing. Blood samples were collected at various time points from 0.083 to
 18 72 hours following iv dosing and for 0.25 to 71 hours after oral dosing. Binding to plasma protein was
 19 determined at some time points over 0.25–4 hours. Samples were analyzed by liquid scintillation counting
 20 and HPLC. Following oral or iv exposure, the percentage of radioactivity recovered in excreta and cage
 21 washes was 81–88% over a 1-week period. As discussed in greater detail in Section 2.1.2.4, most of the
 22 radioactivity was excreted in urine and very little was excreted in feces. Toxicokinetic endpoints are
 23 summarized in Table 37. Based on the toxicokinetic values, study authors concluded that absorption of
 24 bisphenol A following oral exposure was rapid and high, and terminal elimination half-lives of bisphenol
 25 A/metabolites were longer following iv than oral exposure. As discussed in more detail in Section 2.1.2.3,
 26 glucuronide compounds were the major metabolites detected in urine, and higher percentages of the
 27 radioactive dose in plasma were represented by bisphenol A following iv than oral dosing.

28

1 **Table 37. Toxicokinetic Endpoints for Radioactivity in Male and Female Cynomolgus Monkeys**
 2 **Exposed to ^{14}C -Bisphenol A Through iv Injection or by Gavage**

Endpoint	Male	Female
<i>Intravenous exposure</i>		
AUC, $\mu\text{g}\cdot\text{eq}\cdot\text{hour/L}$	377 \pm 85	382 \pm 96
Volume of distribution, L/kg	1.58 \pm 0.11	1.82 \pm 0.41
Half-life, hours	13.5 \pm 2.6	14.7 \pm 2.1
Total body clearance, L/hours/kg	0.27 \pm 0.05	0.28 \pm 0.08
Mean residence time, hours	5.93 \pm 0.91	6.68 \pm 0.72
<i>Oral exposure</i>		
AUC, $\mu\text{g}\cdot\text{eq}\cdot\text{hour/L}$	265 \pm 74	244 \pm 21
T _{max} , hours	1.00 \pm 0.87	0.33 \pm 0.14
C _{max} , $\mu\text{g}\cdot\text{eq/L}$	104 \pm 85	107 \pm 37
Half-life, hours	9.63 \pm 2.74	9.80 \pm 2.15
Bioavailability	0.70 \pm 0.16	0.66 \pm 0.13

[Mean \pm SD assumed based on data presentations elsewhere in this paper.]

From Kurebayashi et al. (98).

3
 4 Negishi et al. (96) compared toxicokinetics of bisphenol A in female F344/N rats, cynomolgus monkeys,
 5 and Western chimpanzees. Bisphenol A was administered by oral gavage and sc injection at doses of 10
 6 or 100 mg/kg bw/day to rats and monkeys and 10 mg/kg bw to chimpanzees. Three rats/dose/time point
 7 were killed before and at various times between 0.5 and 24 hours following bisphenol A administration.
 8 Three monkeys/group and 2 chimpanzees were first exposed orally and 1 week later by sc injection. In
 9 monkeys, blood samples were drawn before and at various times from 0.5 to 24 hours after dosing. In
 10 chimpanzees, blood was drawn before and at multiple time points between 0.25 and 24 hours following
 11 dosing. Bisphenol A was measured in serum by ELISA, and toxicokinetics endpoints were determined.
 12 Results are summarized in Table 38. The study authors noted that the bioavailability of bisphenol was
 13 lowest in rats < chimpanzees < monkeys following exposure through either route. In most cases,
 14 bisphenol A was not detected in rat serum following oral administration of the 10 mg/kg bw dose. In all
 15 species, higher bioavailability was observed with sc than oral dosing.

16
 17 **Table 38. Toxicokinetic Endpoints for Bisphenol A in Rats, Monkeys, and Chimpanzees**

Endpoints	10 mg/kg bw		100 mg/kg bw	
	Oral	SC	Oral	SC
<i>Rat (data presented as mean \pm SD)</i>				
C _{max} , $\mu\text{g/L}$		872 \pm 164	580 \pm 398	3439 \pm 679
T _{max} , hours		1.0	0.5	1.0
AUC _{0-4h} , $\mu\text{g}\cdot\text{hour/L}$		1912 \pm 262	506 \pm 313	9314 \pm 2634
AUC _{0-24h} , $\mu\text{g}\cdot\text{hour/L}$		3377 \pm 334	1353 \pm 462	23,001 \pm 6387
<i>Monkey (data presented as mean \pm SD)</i>				
C _{max} , $\mu\text{g/L}$	2793 \pm 920	57,934 \pm 1902	5732 \pm 525	10,851 \pm 3915
T _{max} , hours	0.7 \pm 0.2	2.0 \pm 0.0	0.7 \pm 0.2	2.0 \pm 0.0
AUC _{0-4h} , $\mu\text{g}\cdot\text{hour/L}$	3209 \pm 536	15,316 \pm 5856	14,747 \pm 2495	48,010 \pm 11,641
AUC _{0-24h} , $\mu\text{g}\cdot\text{hour/L}$	3247 \pm 587	39,040 \pm 10,738	52,595 \pm 8951	189,627 \pm 21,790
<i>Chimpanzee (data presented for 2 animals)</i>				
C _{max} , $\mu\text{g/L}$	325; 96	2058; 1026	Dose not administered	
T _{max} , hours	0.5; 0.5	2.0; 2.0		
AUC _{0-4h} , $\mu\text{g}\cdot\text{hour/L}$	491; 235	5658; 3109		
AUC _{0-24h} , $\mu\text{g}\cdot\text{hour/L}$	1167; 813	21,141; 12,492		

Data were not reported in cases where table cells are empty.

From Negishi et al. (96).

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2.1.2.3 Metabolism

Information is arranged in this section according to species. In rats, study summaries are arranged in order of those providing general or route-specific information on metabolites, specifics on organs or enzyme isoforms involved in metabolism, and pregnancy-, sex-, or age-related effects on metabolism.

Pottenger et al. (93) examined the effects of dose and route on toxicokinetics of bisphenol A in rats. Disposition of bisphenol A and its metabolites in urine and feces is primarily described in this section, while results of the toxicokinetics study are primarily described in Section 2.1.2.2. Five adult F344 rats/sex/group were dosed with ¹⁴C-bisphenol A (99.3% radiochemical purity)/non-radiolabeled bisphenol A (99.7% purity) at doses of 10 or 100 mg/kg bw by oral gavage or ip or sc injection. Excreta were collected for 7 days. Samples were analyzed by HPLC or HPLC/electrospray ionization/MS. The percentage of radioactivity recovered from all groups was 84–98%. Fecal elimination represented the largest percentage of radioactivity in all exposure groups (52–83%). Eight peaks were identified in feces, and the largest peak (representing 86–93% of radioactivity) was for unchanged bisphenol A. Elimination of radioactivity through urine was ~2-fold higher in females (21–34%) than males (13–16%) in all dose groups. Fourteen different peaks were identified in urine. It was estimated that radioactivity in urine was represented by bisphenol A monoglucuronide (57–87%), bisphenol A (3–12%), and bisphenol A sulfate (2–7%). Some differences were noted for retention of radioactivity following dosing by gavage (0.03–0.26%), ip injection (0.65–0.85%), and sc injection (1.03–1.29%).

Metabolites associated with bisphenol A exposure were examined in a second study by Pottenger et al. (93). Three rats/sex/dose/route/time point were dosed with ¹⁴C-bisphenol A/non-radiolabeled bisphenol A at 10 or 100 mg/kg bw by oral gavage or ip or sc injection. Rats were killed at 2 different time points following dosing, T_{max}, and the time when bisphenol A levels were no longer quantifiable. Times at which rats were killed were determined by data obtained during the first study. Plasma samples were pooled at each time period and examined by HPLC or HPLC/electrospray ionization/MS. Qualitative and quantitative differences were observed for parent compound and metabolites in plasma following exposure through different routes. Following oral exposure, bisphenol A glucuronide was the most abundant compound detected in plasma at both time periods (C_{max} and time when parent compound was not quantifiable) and represented 68–100% of total radioactivity. Following ip or sc exposure, unmetabolized bisphenol A was the most abundant compound at T_{max}; levels of radioactivity represented by unmetabolized bisphenol A were 27–51% following ip exposure and 65–76% following sc exposure. Only 2–8% of radioactivity was represented by bisphenol A following oral exposure. Some compounds observed following ip or sc exposure were not observed following oral exposure. A compound tentatively identified as a sulfate conjugate was observed following ip exposure and represented a small portion of radioactivity. An unresolved peak of 3 compounds was observed following ip or sc exposure, at the time when parent compound was not quantifiable and represented that major percent of radioactivity for that time point. Three additional unidentified, minor peaks were observed following ip or sc but not oral exposure. The major sex differences observed were higher C_{max} values for bisphenol A and bisphenol A glucuronide in females than males, especially following ip administration. A review by the European Union (2) noted that the substantially higher concentrations of parent compound with ip and sc compared to oral exposure indicated the occurrence of first-pass metabolism following oral intake.

Elsby et al. (113) examined bisphenol A metabolism by rat hepatocytes. In the hepatocyte metabolism study, hepatocytes were isolated from livers of adult female Wistar rats and incubated in dimethylsulfoxide (DMSO) vehicle or bisphenol A 100 or 500 μM [23 or 114 mg/L] for 2 hours. Metabolites were identified by HPLC or LC/MS. Data were obtained from 4 experiments conducted in duplicate. At both concentrations, the major metabolite was identified as bisphenol A glucuronide, which was the only metabolite identified following incubation with 100 μM bisphenol A. Two additional minor metabolites identified at the 500 μM concentration included 5-hydroxy-bisphenol A-sulfate and bisphenol

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1 A sulfate. Another part of the study comparing metabolism of bisphenol A by rat and human metabolites
2 is discussed in Section 2.1.1.3. Another study (114) comparing metabolism of bisphenol A in humans,
3 rats, and mice is also summarized in Section 2.1.1.3.

4
5 In neonatal rats gavaged with 1 or 10 mg/kg bw ¹⁴C-bisphenol A on PND 4, 7, and 21 and adult rats
6 gavaged with 10 mg/kg bw bisphenol A, the major compounds detected in plasma were bisphenol A
7 glucuronide and bisphenol A (92). Up to 13 radioactive peaks were identified in neonatal rats dosed with
8 10 mg/kg bw and 2 were identified in neonates dosed with 1 mg/kg bw/day. At the 10 mg/kg bw dose, the
9 level of bisphenol A glucuronide detected in plasma increased with age. Metabolic profiles were
10 generally similar in males and females. The study authors noted that metabolism of bisphenol A to its
11 glucuronide conjugate occurs as early as PND 4 in rats. However, age-dependent differences were
12 observed in neonatal rats, as noted by a larger fraction of the lower dose being metabolized to the
13 glucuronide. More details from this study are included in Section 2.1.2.2.

14
15 Kurebayashi et al. (101) used a thin layer chromatography technique to examine metabolite profiles in
16 blood, urine, and feces of 3 male rats orally dosed with 0.5 mg/kg bw ¹⁴C-bisphenol A. **[The procedure
17 did not identify metabolites.]** Parent bisphenol A represented ~2% of the dose in plasma at 0.25 and 6
18 hours post dosing and ~0.3% of the dose at 24 hours after exposure. Unmetabolized bisphenol A
19 represented 1.6% of compounds in urine and 77.2% of compounds in feces collected over a 24-hour
20 period. Free bisphenol A represented 47.1% of compounds in urine following β-glucuronidase hydrolysis
21 of urine, and there was an almost equivalent decrease in a metabolite the study authors identified as
22 “M2.” Therefore, the study authors stated that M2 was most likely bisphenol A glucuronide. M2 was the
23 major metabolite identified in plasma (~74–77%) and urine (~40%).

24
25 The European Union (2) reviewed studies by Atkinson and Roy (115, 116) that reported two major and
26 several minor adducts in DNA obtained from the liver of CD-1 rats dosed orally or ip with 200 mg/kg bw
27 bisphenol A. Chromatographic mobility of the two major adducts was the same as that observed when
28 bisphenol A was incubated with purified DNA and a peroxidase or microsomal P450 activation system.
29 The profile closely matched that of adducts formed with the interaction between bisphenol O-quinone and
30 purified rat DNA deoxyguanosine 3'-monophosphate. Formation of the adduct appeared to be inhibited
31 by known inhibitors of cytochrome P (CYP) 450. It was concluded that bisphenol A is possibly
32 metabolized to bisphenol O-quinone by CYP450.

33
34 Biliary excretion of bisphenol A and its metabolites following oral or iv dosing with bisphenol A was
35 examined by Kurebayashi et al. (110). Bile ducts of 3 rats/sex/group were cannulated, and the rats were
36 dosed with 0.1 mg/kg bw ¹⁴C-bisphenol A (>99% radiochemical purity) in phosphate buffer vehicle by
37 oral gavage or iv injection. Biliary fluid was collected every 2 hours over a 6-hour period to determine
38 percent total biliary excretion and percent of dose represented by bisphenol A glucuronide. Results are
39 summarized in Table 39. The study authors noted that the importance of biliary excretion following oral
40 or iv dosing. ¹⁴C-bisphenol A-glucuronide was the predominant metabolite in bile.

1 **Table 39. Biliary Excretion in Male and Female Rats Exposed to 0.1 mg/kg bw ¹⁴C-Bisphenol A**
 2 **Through the Oral or iv Route**

Parameters	Male		Female	
	IV	Oral	IV	Oral
Biliary excretion, %				
0–2 hours	48	32	35	28
0–4 hours	61	44	50	39
0–6 hours	66	50	58	45
Radioactivity in bile represented by glucuronide, %	84	86	87	88
Dose excreted as glucuronide in bile, %	55	43	50	40

From Kurebayashi et al. (110).

3
 4 In another study by Kurebayashi et al. (110), biliary, fecal, and urinary metabolites were examined in
 5 male rats gavaged with 100 mg/kg bw bisphenol A or D₁₆-bisphenol A in corn oil. Bile was collected over
 6 an 18-hour period, and urine and feces were collected over a 72-hour period. The primary metabolite
 7 detected in urine was bisphenol A glucuronide, which represented 6.5% of the dose. Lower percentages of
 8 the dose (≤1.1%) were present in urine as bisphenol A and bisphenol A sulfate. In feces, the primary
 9 compound detected was bisphenol A, which represented 61% of the dose. No glucuronide or sulfate
 10 conjugated metabolites of bisphenol A were detected in feces. Most of the dose in bile consisted of
 11 bisphenol A glucuronide (41% of the dose). Bisphenol A represented 0.3% of the dose in bile. The study
 12 authors noted that as with oral or iv exposure to a smaller dose, feces was the main route of elimination
 13 for bisphenol A and bile was the main elimination route for bisphenol A glucuronide.

14
 15 A study by Yokota et al. (117) examined the hepatic isoform of uridine diphosphate
 16 glucuronosyltransferase (UDPGT) involved in the metabolism of bisphenol A and distribution of the
 17 enzyme in organs of Wistar rats. Using yeast cells genetically engineered to express single rat UDPGT
 18 enzymes, it was determined that UGT2B1 was the only isoform capable of glucuronidating bisphenol A.
 19 Microsomal UDPGT activity towards bisphenol A was demonstrated in liver, kidney, and testis, but
 20 activity was minimal in lung and brain. **[Minimal activity was also observed for intestine]**. Northern
 21 blot analyses revealed high expression of UGTB1 only in liver. It was demonstrated that 65% of
 22 glucuronidation activity was absorbed by binding with anti-UGTB1, indicating that additional isoforms
 23 are likely involved in glucuronidation of bisphenol A.

24
 25 The intestine was determined to play a role in the metabolism of bisphenol A in rats. Nine-week-old male
 26 Sprague Dawley rats were orally administered 0.1 mL of a solution containing 50 g/L bisphenol **[5 mg**
 27 **total or ~17 mg/kg bw assuming a body weight of ~0.3 kg (81)]** (118). Rats were killed at multiple time
 28 intervals between 15 minutes and 12 hours following exposure. The small intestine was removed and
 29 separated into upper and lower portions. Intestinal contents were removed from each section. Bisphenol A
 30 and metabolite levels were measured by HPLC. Activities and expression of β-glucuronidase were
 31 determined. A large amount of bisphenol A glucuronide was detected in the upper and lower portions of
 32 the small intestine, and a large amount of free bisphenol A was detected in the cecum. Less bisphenol A
 33 was detected in colon and feces. The observations lead the study authors to conclude that free bisphenol A
 34 generated in the cecum as a result of deconjugation was reabsorbed in the colon. The presence of large
 35 amounts of bisphenol A glucuronide in the small intestine at 12 hours following exposure suggested that
 36 bisphenol A was reabsorbed in the colon and re-excreted as the glucuronide. As determined in an assay
 37 using *p*-nitrophenol-β-*d*-glucuronide as a substrate, ~70% of total β-glucuronidase activity was present in
 38 the cecum and 30% in the colon. Western blot analysis revealed a large amount of bacterial β-
 39 glucuronidase protein in cecum and colon contents.

40
 41 Glucuronidation and absorption of bisphenol A in rat intestine were studied by Inoue et al. (119).
 42 Intestines were obtained from 8-week-old male Sprague Dawley rats, and the small intestine was divided

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1 into 4 sections. Small intestine and colon were everted and exposed to 40 mL of a solution containing
2 bisphenol A at 10, 50, or 100 μM [**2.3, 11, or 23 mg/L, resulting in delivery of 91, 456, or 913 μg**
3 **bisphenol A to the everted intestine**]. Every 20 minutes during a 60-minute time period, reaction
4 products were collected from serosal and mucosal sides and analyzed by HPLC. Optimal glucuronidation
5 was observed at 50 μM [**11 mg/L**]. At 60 minutes following exposure to 50 μM bisphenol A, ~37% of
6 bisphenol A was absorbed by the small intestine and ~83% was glucuronidated. Approximately 74.7% of
7 the glucuronide was excreted on the mucosal side and ~25.3% transported to the serosal side of small
8 intestine. Slightly greater absorption of bisphenol A in the colon (48.6%) compared to the proximal
9 jejunum (37.5%) was observed at 60 minutes following exposure to the 50 μM solution. Transport of both
10 bisphenol A and bisphenol A glucuronide to the serosal side of intestine increased distally and was
11 greatest in the colon. Minimal mucosal excretion was observed in the colon.

12
13 Inoue et al. (120) compared glucuronidation of bisphenol A in pregnant, non-pregnant, and male rats.
14 Livers of 4 male and non-pregnant Sprague Dawley rats/group were perfused via the portal vein for 1
15 hour with solutions containing bisphenol A at 10 or 50 μM [**2.3 or 11 mg/L**]. The total amount of
16 bisphenol A infused into livers was 1.5 or 7.5 μmol [**0.34 or 1.7 mg**]. On GD 20 or 21, livers of 4
17 pregnant Sprague Dawley rats were perfused for 1 hour with 10 μM [**2.3 mg/L**] bisphenol A. At the start
18 of perfusion, excreted bile and perfusate in the vein were collected every 5 minutes for 1 hour. Samples
19 were analyzed by HPLC. Statistical analyses were conducted by Student *t*-test and ANOVA. Bisphenol A
20 glucuronidation in the liver was 59% in male rats and 84% in non-pregnant female rats perfused with the
21 10 μM solution. The glucuronide was excreted primarily through bile in both males and females, but a
22 significantly higher amount was excreted through bile in non-pregnant females than in males. The total
23 amount of glucuronide excreted into bile and vein was ~1.4-fold higher in females than males following
24 perfusion with the 10 μM [**2.3 mg/L**] solution. At the 50 μM [**11 mg/L**] concentration, bisphenol A
25 glucuronidated within liver was 66% in males and 91% in females. In males the glucuronide was excreted
26 mainly in bile, and in females, a higher amount of glucuronide was excreted in the vein. In livers of
27 pregnant rats perfused with the 10 μM [**2.3 mg/L**] solution, 69% of bisphenol A was glucuronidated in
28 the liver. Percentages of glucuronide excretion were 54.5% through bile and 45.5% through the vein in
29 pregnant rats. In a comparison of pregnant rats and non-pregnant rats perfused with 10 μM [**2.3 mg/L**]
30 bisphenol A, biliary excretion in pregnant rats was half that observed in non-pregnant rats, and venous
31 excretion in pregnant rats was 3-fold higher than in non-pregnant rats. To determine the pathway of
32 bisphenol A glucuronide excretion, livers of 4 male Eisai hyperbilirubinemic rats, a strain deficient in
33 multidrug resistance-associated protein, were perfused with 50 μM [**11 mg/L**] bisphenol A. During and
34 after perfusion, nearly all of the bisphenol A was excreted into the vein, thus indicating that multidrug
35 resistance-associated protein mediates biliary excretion of bisphenol A glucuronide. The study authors
36 concluded that bisphenol A is highly glucuronidated and excreted into bile using a multidrug resistance-
37 associated protein-dependent mechanism, and that venous excretion increases and biliary excretion
38 decreases during pregnancy.

39
40 Miyakoda et al. (121) examined the production of bisphenol A glucuronide in fetal and adult rats.
41 Bisphenol A was orally administered at 10 mg/kg bw to pregnant Wistar rats on GD 19 and to 10-week-
42 old adult male Wistar rats. **[The numbers of animals exposed was not reported. In some legends for**
43 **study figures, it was stated that the data were from 4 experiments, suggesting that 4 pregnant rats**
44 **and adult males may have been exposed.]** Fetuses were removed at 1 hour following dosing. Blood was
45 drawn and testes were removed from adult males at 1, 3, and 8 hours following dosing. GC/MS was used
46 to measure bisphenol A levels in 19 fetuses and in testis of adult rats prior to and following
47 homogenization with β -glucuronidase. In fetal extracts, there were no differences in bisphenol A levels
48 before or after treatment with β -glucuronidase, suggesting that bisphenol A glucuronide was not present
49 at detectable levels. The study authors noted the possibility that bisphenol A glucuronide was not
50 transferred from dams to fetuses and stated that glucuronidation by the rat fetus is unlikely. At 1 hour
51 following dosing of adult male rats, 90% of bisphenol A was detected as glucuronide in plasma and testis.

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1 Bisphenol A glucuronide levels gradually decreased and bisphenol A levels increased slightly in testis
2 over the 8-hour sampling period. In plasma, bisphenol A-glucuronide decreased to 55% of the maximum
3 observed concentration at 3 hours following dosing and increased to 100% of maximum observed level at
4 8 hours following dosing. Based on levels of bisphenol A glucuronide in testis and blood (40 ppb [$\mu\text{g}/\text{kg}$]
5 and 600 ppb [$\mu\text{g}/\text{L}$]) at 8 hours, the study authors concluded that bisphenol A glucuronide passage
6 through the testicular barrier was unlikely. It was thought that bisphenol A passed through the testicular
7 barrier, was converted to the glucuronide within the testis, and was then gradually released following
8 digestion of the glucuronide by β -glucuronidase.

9
10 Matsumoto et al. (122), studied developmental changes in expression and activity of the UDPGT isoform
11 UGT2B towards bisphenol A in Wistar rats. Activity towards other compounds was also examined but
12 this summary focuses on bisphenol A. Microsomes were prepared from livers of fetuses, neonates on
13 PND 3, 7, 14, and 21, and pregnant rats on GD 10, 15, and 19. Activity towards the bisphenol A substrate
14 was measured using an HPLC method. Expression of UGT2B1 protein was examined by Western blot
15 and messenger ribonucleic acid (mRNA) expression was examined by Northern blot. Little-to-no UGT2B
16 activity towards bisphenol A was detected in microsomes of fetuses. Activity increased linearly following
17 birth and reached adult levels by PND 21. **[No data on UGT2B activity for non-pregnant adult rats
18 were shown and it was not clear if activity in adults was examined in this study.]** The same
19 developmental patterns were observed for expression of UGT2B1 protein and mRNA. Activity and
20 protein expression of UGT2B1 were also found to be reduced in pregnant rats.

21
22 The European Union (2) reviewed an unpublished study by Sipes that compared clearance of bisphenol A
23 by hepatic microsome from fetal ($n = 8/\text{sex}$), immature ($n = 4/\text{sex}$), and adult ($n = 4$) rats. The clearance
24 rate in microsomes from male and female GD 19 rat fetuses (0.7–0.9 mL/minute/mg) was lower than
25 clearance rates in microsomes from 4-day-old males and females (1.2–2.6 mL/minute/mg), 21-day-old
26 males and females (2.4–2.7 mL/minute/mg), and their dams (2.6 mL/minute/mg). The European Union
27 concluded that clearance rate was lower in fetuses but reached adult levels by 4 days of age.

28
29 In a qualitative study of bisphenol A metabolites in pregnant mice injected with 0.025 mg/kg bw
30 bisphenol A, 10 radioactive peaks were observed in urine by Zalko et al. (108). The major metabolites
31 detected in urine were bisphenol A glucuronide and a hydroxylated bisphenol A glucuronide. Unchanged
32 bisphenol A was the major compound detected in feces (>95%). Bisphenol A glucuronide represented
33 more than 90% of the compounds detected in bile. Additional compounds detected in urine, feces,
34 digestive tract, or liver included a double glucuronide of bisphenol A and sulfate conjugates. Unchanged
35 bisphenol A, bisphenol A glucuronide, and “metabolite F” were the major compounds detected in all
36 tissues. The most abundant compound in all tissues was bisphenol A glucuronide, except in placenta
37 where bisphenol A and metabolite F were the major compounds detected. Concentrations of bisphenol A
38 decreased rapidly in all tissues. It was determined that metabolite F was most likely bisphenol A
39 glucuronide conjugated to acetylated galactosamine or glucosamine. Distribution of bisphenol A and its
40 metabolites in maternal and fetal tissues is summarized in Table 28. Additional details of this study are
41 included in Section 2.1.2.2.

42
43 Jaeg et al. (123) reported metabolites observed following incubation of CD-1 mouse liver microsomes or
44 S9 fractions with bisphenol A at 20–500 μM [4.6–114 mg/L]. The metabolites included isopropyl-
45 hydroxyphenol, bisphenol A glutathione conjugate, glutathionyl-phenol, glutathionyl 4-isopropylphenol,
46 2,2-bis-(4-hydroxyphenyl)1-propanol, 5-hydroxy bisphenol A, and bisphenol A dimers.

47
48 Kurebayshi et al. (98) examined metabolism of bisphenol A in monkeys. Three adult male and female
49 cynomolgus monkeys were dosed with 0.1 mg/kg bw ^{14}C -bisphenol A/non-radiolabeled bisphenol A by iv
50 injection on study day 1 and by gavage on study day 15 (98). Additional details of the study are included
51 in Section 2.1.2.2. Up to five peaks were identified in urine. Analysis by radio-HPLC suggested that the

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1 major peaks in both sexes treated by either exposure route were mono- and diglucuronides. Five peaks
 2 were identified in plasma, and some differences were noted in comparisons of iv to oral exposure. In the 2
 3 hours following dosing, most of the radioactivity in plasma was represented by bisphenol A glucuronide
 4 after iv dosing (57–82%) and oral dosing (89–100%). The percentage of radioactivity represented by
 5 unchanged bisphenol A was higher following iv (5–29%) than oral (0–1%) dosing.

6
 7 Kang et al. (124) reviewed studies that provided some information about metabolism of bisphenol A in
 8 fish and birds. One study reported bisphenol A sulfate and bisphenol A glucuronide as the major
 9 metabolites detected in zebrafish exposed to bisphenol A. A second study conducted in carp reported an
 10 increase in UDPGT activity for bisphenol A in microsomes and metabolism of bisphenol A to bisphenol
 11 A glucuronide in intestine. In quail embryos, metabolism and excretion of bisphenol A was reported, but
 12 specific metabolites were not indicated. Another study reported that ¹⁴C-bisphenol A administered orally
 13 or iv to laying quail was rapidly removed via bile and excreted through feces.

14 2.1.2.4 Elimination

15 Elimination of bisphenol A and its metabolites was examined in Sprague Dawley rats that were gavaged
 16 with bisphenol A and ¹⁴C-bisphenol A at 10 mg/kg bw (100). One group of rats was not pregnant, and 3
 17 additional groups were treated on either GD 6 (early gestation), 14 (mid gestation), or 17 (late gestation).
 18 More details of this study are available in Section 2.1.2.2. Most of the radioactivity (65–78%) was
 19 eliminated in feces. Elimination in urine accounted for 14–22% of the dose, and considerable variability
 20 for urinary elimination among animals was evident by the large standard deviations, which were 50% of
 21 means. The authors stated that bisphenol A glucuronide represented 62–70% of radioactivity in urine and
 22 bisphenol A represented 19–23% of radioactivity in urine [data were not shown by authors]. A total of
 23 9 peaks were identified in urine. In feces, 83–89% of radioactivity was represented by bisphenol A and 2–
 24 3% was represented by bisphenol A glucuronide; 7 peaks were identified in feces. The study authors
 25 concluded that urinary elimination and fecal elimination of radioactivity were similar in pregnant and
 26 non-pregnant rats.

27
 28 Difference in excretion following oral or iv exposure of rats to a low bisphenol A dose were examined by
 29 Kurebayashi et al. (110). Three male rats/group were exposed to 0.1 mg/kg bw ¹⁴C-bisphenol A (> 99%
 30 radiochemical purity) in phosphate buffer vehicle by oral gavage or iv injection. Radioactivity levels were
 31 measured in urine and feces, which were collected over a 48-hour period. Additional details of the study
 32 are included in Section 2.1.2.2. Results of that study are summarized in Table 40. With both oral and IV
 33 dosing, fecal excretion was the main route of elimination.

34
 35 **Table 40. Excretion of Radioactivity Following Oral or iv Dosing of Rats with 0.1 mg/kg bw ¹⁴C-**
 36 **Bisphenol A**

Time post dosing, hours	Percent radioactive dose excreted		
	Urine	Feces	Total
<i>Oral</i>			
0–24	6.3 ± 1.1	49.3 ± 2.1	55.7 ± 2.8
24–48	3.8 ± 1.0	32.3 ± 2.1	36.1 ± 3.0
Total	10.1 ± 1.6	81.6 ± 3.7	91.8 ± 5.0
<i>iv</i>			
0–24	8.4 ± 1.8	46.2 ± 1.8	54.6 ± 3.4
24–48	4.1 ± 0.9	31.4 ± 1.5	35.4 ± 1.8
Total	12.5 ± 0.9	77.6 ± 1.8	90.1 ± 2.7

Values presented as mean ± SD.
 From Kurebayashi et al. (110).

37

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1 Kurebayashi et al. (101) examined elimination of radioactivity in 3 adult male and female F344 rats that
2 were orally dosed with 0.1 mg/kg bw ¹⁴C-bisphenol A. Urine and feces were collected over a 168-hour
3 period and analyzed by liquid scintillation counting. Total radioactivity excreted in urine and feces over
4 the 168-hour period was ~98% in males and females. In male rats, ~10% was excreted in urine and ~88%
5 was excreted in feces. Female rats excreted ~34% of the radioactivity in urine and ~64% in feces. **[The**
6 **majority of radioactivity, ~90%, was excreted over 48 hours by males and 72 hours by females.]**
7

8 Snyder et al. (103) compared toxicokinetics of bisphenol A in CD and F344 rats. Four CD and F344 rats
9 were gavaged with 100 mg/kg bw ¹⁴C-bisphenol A in propylene glycol vehicle. Disposition of
10 radioactivity in urine, feces, and carcass was examined over a 144-hour period. Samples were analyzed by
11 scintillation counting, HPLC, or nuclear magnetic resonance. Data were analyzed by ArcSin
12 transformation of the square root of the mean and using two-sample *t*-test. Recovery of radioactivity was
13 93% in both strains. The highest concentrations of radioactivity were detected in feces (70% of dose in
14 CD rat and 50% of dose in F344 rats) followed by urine (21% of dose in CD rat and 42% of dose in F344
15 rats). The percentages of the dose excreted in urine and feces differed significantly by strain. Much lower
16 percentages of radioactivity were detected in the carcass (~1%). Bisphenol A glucuronide, representing
17 81–89% of the dose, was the major urinary metabolite detected in both strains. A much lower percentage
18 (2.2–10%) of the dose was represented by urinary bisphenol A.
19

20 Kim et al. (125) reported urinary excretion of bisphenol A in 4-week-old male F344 rats given bisphenol
21 A in drinking water at 0 (ethanol vehicle), 0.1, 1, 10, or 100 ppm (equivalent to 0.011, 0.116, 1.094, or
22 11.846 mg/kg bw/day) for 13 weeks. Urine samples were collected for 24 hours following administration
23 of the last dose and analyzed by HPLC before and after digestion with β-glucuronidase. The focus of the
24 study was male reproductive toxicity; the study is described in detail in Section 4.2.2.1. Bisphenol A was
25 not detected in the urine of rats from the control and 2 lowest dose groups. **[At the 2 highest doses, free**
26 **bisphenol A represented 60 and 30% of the total urinary bisphenol A levels.]**
27

28 In rats exposed to 10 or 100 mg/kg bw/day ¹⁴C-bisphenol A through the oral, ip, or sc routes, fecal
29 elimination represented the highest percentage of radioactivity in all exposure groups (52–83%) (93).
30 Elimination of radioactivity through urine was ~2-fold higher in females (21–34%) than males (13–16%)
31 in all dose groups. Additional details of this study are included in Section 2.1.2.3.
32

33 Elimination of bisphenol A and metabolites was examined in 3 adult male and female cynomolgus
34 monkeys dosed with 0.1 mg/kg bw ¹⁴C-bisphenol A/non-radiolabeled bisphenol A by iv injection on
35 study day 1 and by gavage on study day 15 (98). Additional details of the study are included in Section
36 2.1.2.2. Following oral or iv exposure, the percentage of radioactivity recovered in excreta and cage
37 washes was 81–88% over a 1-week period. Most of the radioactivity was recovered in urine (combination
38 of urine and cage washes), with most of the radioactivity excreted in urine within 12 hours and essentially
39 all of the dose excreted within 24 hours following treatment. Percentages of radioactive doses recovered
40 in urine within 1 week after dosing were ~79–86% following iv dosing and 82–85% following oral
41 dosing. Much smaller amounts were recovered in feces during the week following iv or oral exposure
42 (~2–3%). The study authors concluded that because fecal excretion was very low following oral exposure,
43 absorption was considered to be complete. The authors also noted that there were no obvious route or sex
44 differences in excretion of radioactivity. The study authors concluded that terminal elimination half-lives
45 were longer following iv than oral exposure. A limited amount of information was presented for the fast
46 phase, defined as the 2 hours following iv injection. Fast-phase elimination half-life of bisphenol A
47 following iv exposure was significantly lower in females (0.39 hours) than males (0.57 hours). There
48 were no sex-related differences in fast-phase half-life for bisphenol A glucuronide (0.79–0.82 hours) or
49 total radioactivity (0.61–0.67 hours).
50

2.1.3 Comparison of humans and experimental animals

Studies comparing toxicokinetics and metabolism of bisphenol A in humans and laboratory animals were reviewed and are summarized below. In most cases the data were from original sources, but information from secondary sources was included if the information was not new or critical to the evaluation of developmental or reproductive toxicity.

Elsby et al. (113) compared bisphenol A metabolism by rat and human microsomes. Microsomes were obtained from 8 immature Wistar rats (21–25 days old) and histologically normal livers from 4 male (25–57 years old) and 4 female (35–65 years old) Caucasian donors who were killed in accidents. Human microsomes were pooled according to sex of the donor. Glucuronidation was examined following exposure of microsomes to bisphenol A concentrations of 0–1000 μM [0–228 mg/L] for 30 minutes with human microsomes and 10 minutes with rat microsomes. Metabolites were identified by HPLC or LC/MS. Data were obtained from 4 experiments conducted in duplicate. Data were analyzed by Mann-Whitney test. Maximum velocity (V_{max}) and the rate constant (K_m) values are summarized in Table 41. The study authors reported a significant difference between the V_{max} for glucuronidation in immature rats and humans. No sex-related difference was reported for glucuronidation by human microsomes. As a result of less extensive glucuronidation by human than rat microsomes, the study authors noted that estrogen target tissues in humans may receive higher exposure to bisphenol A than tissues of immature female rats used in estrogenicity studies. Lastly, oxidation of bisphenol A by female rat or human microsomes was examined following incubation with 200 μM [46 mg/L] bisphenol A and NADPH. The only metabolite identified was 5-hydroxybisphenol A.

Table 41. Glucuronidation Kinetics in Microsomes From Immature Rats and Adult Humans

Sex/Species	V_{max} , nmol/minute/mg protein	K_m , μM
Male/human	5.9 \pm 0.4	77.5 \pm 8.3
Female/human	5.2 \pm 0.3	66.3 \pm 7.5
Female/immature rat	31.6 \pm 8.1	27.0 \pm 1.2

Data presented as mean \pm SEM.

From Elsby et al. (113).

The European Union (2) reviewed a series of studies by Sipes that compared metabolism of bisphenol A in microsomes from male and female humans (15 pooled samples/sex and 3–5 individual samples/sex), rats (4/sex), and mice (4/sex). It was concluded that the studies generally agreed with the findings of Elsby et al. (113). Clearance rates (V_{max}/K_m) in human microsomes (0.4–0.9 mL/minute/mg for pooled samples and 0.3–0.5 mL/minute/mg in individual samples) were lower than those observed in rats (1.0–1.7 mL/minute/mg) and mice (1.3–3.0 mL/minute/mg).

Pritchett et al. (114) compared metabolism of bisphenol A in hepatocyte cultures from humans, rats, and mice. Cell cultures were prepared from adult male and female F344 rats, Sprague Dawley rats, and CF1 mice. Human hepatocyte cultures were obtained from 3 females and 2 males. [No information was provided about the age of human donors.] Cells were exposed to ^{14}C -bisphenol A (99.3% purity)/bisphenol A (>99% purity) in a DMSO vehicle. In a cytotoxicity assessment, lactate dehydrogenase activity was measured in rat cells following incubation for 18 hours in 5–100 μM [1.1–23 mg/L] bisphenol A, and cytotoxicity was observed at ≥ 75 μM bisphenol A. Bisphenol A concentrations tested and times of exposure were 5–20 μM [1.1–4.6 mg/L] for up to 6 hours in time-dependent metabolism studies and 2.5–30 μM [0.57–6.8 mg/L] for 10 minutes in concentration-dependent metabolism studies. Metabolites in cell media were analyzed by HPLC and LC-tandem MS.

Analysis of media from human hepatocytes incubated with bisphenol A indicated that the major metabolite was bisphenol A glucuronide, and compounds found at lower concentrations were bisphenol A glucuronide/sulfate diconjugate, and bisphenol A sulfate conjugate. Table 42 summarizes percentages of

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1 each type of metabolite detected in media following incubation with 20 μM [4.6 mg/L] bisphenol A for 3
 2 hours in human cells and 6 hours in rodent cells. In cells from all sexes and species except male F344 rats,
 3 bisphenol A glucuronide was the major metabolite detected. The glucuronide/sulfate diconjugate was the
 4 major metabolite detected in cells from male F344 rats. In concentration-dependent studies conducted in
 5 F344 rat hepatocytes, a biphasic curve was obtained following a 10-minute incubation, with a V_{max} of
 6 0.36 nmol/min at bisphenol A concentrations of 20–30 μM [4.6–6.8 mg/L] and a V_{max} of \sim 0.15 nmol/min
 7 at bisphenol A concentrations of 2.5–10 nM [0.57–2.3 mg/L]. Table 43 summarizes the higher V_{max}
 8 values obtained with cells from human, rat, and mouse livers. Total hepatic capacity was determined by
 9 multiplying V_{max} by total number of hepatocytes/liver in vivo. [The only graphical data presented were
 10 for male F344 rats]. The authors noted that V_{max} values were highest in mice > rats > humans. However,
 11 when adjusted for total hepatocyte number in vivo, the values were predicted to be highest in humans >
 12 rats > mice.

13
 14 **Table 42. Metabolites Obtained from Incubation of Human, Rat, and Mouse Hepatocyte Cultures**
 15 **with 20 μM [4.6 mg/L] Bisphenol A**

Sex and species	Percentage of parent compound or metabolites			
	Glucuronide/sulfate	Sulfate	Glucuronide	Bisphenol A
Human samples				
Female-1	4	0	93	0
Female-2	2	0	84	2
Female-3	43	2	55	0
Male-1	1	0	85	0
Male-2	0	7.5	75	0
Rodent samples				
Male F344 rat	70	0	30	0
Female F344 rat	10	0	86	0
Male Sprague Dawley rat	30	2	58	0
Female Sprague Dawley rat	0	0	100	0
Male CF1 Mouse	0	0	100	0
Female CF1 mouse	0	0	93	0

Human cells were incubated for 3 hours, and animal cells were incubated for 6 hours.

From Pritchett et al. (114).

16
 17 **Table 43. Rates of Bisphenol A Glucuronide Formation Following Incubation of Human, Rat, and**
 18 **Mouse Hepatocytes with Bisphenol A**

Species and sex	V_{max} , nmol/min/ 0.5×10^6 hepatocytes	Hepatic capacity, $\mu\text{mol/hours}^a$
Human female	0.27	8000
F344 rat female	0.46	46.5
F344 rat male	0.36	61.8
Sprague Dawley female	0.39	54.5
Sprague Dawley male	0.45	79.9
CF1 mouse female	0.50	13.8
CF1 mouse male	0.82	23.6

^aHepatic capacity was estimated by multiplying V_{max} by total numbers of hepatic cells in vivo.

From Pritchett et al. (114).

19
 20 Data from Pritchett et al. (114) appeared to be included in a series of unpublished studies by Sipes that
 21 were reviewed by the European Union (2). In their review, the European Union noted that metabolic
 22 patterns appear to be similar in humans, rats, and mice. It was stated that the biphasic kinetic profile
 23 indicated involvement of a high-affinity glucuronidase enzyme at low concentrations and a high-capacity

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1 enzyme at high concentrations. In the interpretation of kinetic profiles in humans and experimental
2 animals, the authors of the European Union report noted that the study calculations did not consider in
3 vivo conditions such as varying metabolic capacity of hepatic cells, relationship of hepatic size to body
4 size, and possibly important physiological endpoints such as blood flow. In addition, it was noted that
5 calculations were based on limited data that did not address inter-individual variability in enzyme
6 expression.

7
8 Cho et al. (126) examined toxicokinetics of bisphenol A in mouse, rat, rabbit, and dog and used that
9 information to predict toxicokinetic values in humans. Bisphenol A was administered by iv injection at 2
10 mg/kg bw to 5 male ICR mice and at 1 mg/kg bw to 7 male Sprague Dawley rats, 7 male New Zealand
11 White rabbits, and 5 male beagle dogs. Blood samples were drawn before dosing and at multiple time
12 points between 2 minutes and 6 hours following injection. Serum bisphenol A levels were measured by
13 HPLC. Toxicokinetic endpoints in animals are summarized in Table 44. The study authors noted that
14 clearance and volume of distribution increased with increasing animal weight but that terminal half-life
15 remained relatively constant across the different species. Simple allometric scaling and species-invariant
16 time methods were used to predict values for a 70 kg human, and those values are summarized in Table
17 45. Regression analyses of estimates using the species-invariant time methods demonstrated that data
18 from the 4 animal species was superimposable ($r = 0.94-0.949$).

19
20 **Table 44. Toxicokinetic Endpoints for Bisphenol A in Mice, Rats, Rabbits, and Dogs iv Dosed with**
21 **2 mg/kg bw Bisphenol A**

Endpoint	Mouse ^a	Rat	Rabbit	Dog
Systemic clearance, L/hour	0.3	1.9 ± 0.4	12.6 ± 4.9	27.1 ± 8.0
Volume of distribution, L	0.1	1.3 ± 0.4	7.1 ± 2.3	20.0 ± 5.4
Half-life, minute	39.9	37.6 ± 12.8	40.8 ± 17.1	43.7 ± 21.9

Data are presented as mean ± SD.

^aVariances not reported.

From Cho et al. (126).

22
23 **Table 45. Predicted Bisphenol A Toxicokinetic Endpoints in Humans Based on Results from**
24 **Experimental Animal Studies**

Endpoint	Prediction method			
	Allometric scaling	Kallynochrons	Apolysichrons	Dienetichrons
Systemic clearance, L/hour	127.1	123	120.7	46.0
Volume of distribution, L	125.3	229.7	138.0	149.3
Half-life, minute	43.6	110.4	67.8	196.2

From Cho et al. (126).

25
26 Teeguarden et al. (111) developed a physiologically based pharmacokinetic (PBPK) model for bisphenol
27 A. Rat toxicokinetic data for the model were obtained from the studies by Pottenger et al. (93) and
28 Upmeier et al. (97). Human toxicokinetic data were obtained from the study by Völkel et al. (91). The
29 model was developed to simulate blood and uterine levels of bisphenol A following exposure of humans
30 through relevant routes. Correlations were determined for simulated bisphenol A binding to uterine
31 receptors and increases in uterine wet weight, as determined by an unpublished study by Twomey.
32 Although intestinal metabolism of bisphenol A to the glucuronide metabolite had been recently
33 demonstrated, the model attributed bisphenol A metabolism entirely to the liver. Plasma protein binding
34 was considered in both the rat and human model. The model accurately simulated plasma bisphenol A
35 glucuronide levels in humans orally administered 5 mg bisphenol A, with the exception of
36 underpredicting bisphenol A glucuronide levels at the 24–48 hour period following exposures.
37 Cumulative urinary elimination of bisphenol A glucuronide in human males and females was accurately
38 simulated. Less accurate simulations were observed for toxicokinetics in orally exposed rats, and the

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1 study authors indicated that a likely cause was oversimplification of the rat gastrointestinal compartment.
2 Comparisons in metabolic clearance rates for iv and oral exposure suggested significant intestinal
3 glucuronidation of bisphenol A. Enterohepatic recirculation strongly affected terminal elimination in rats
4 but not humans. Consideration of bound versus unbound bisphenol A was found to be important in
5 simulating occupancy of the estrogen receptor (ER) and uterine weight response. No increase in uterine
6 weight was reported with simulated receptor occupancy of ~1–15%. An increase in uterine weight was
7 reported with ~25% receptor occupancy, and doubling of uterine weight was reported with 63% receptor
8 occupancy.

9 **2.2 General Toxicity, Estrogenicity, and Androgenicity**

10 This section includes information on general toxicity as well as information on estrogenicity and
11 androgenicity; however, results of estrogenicity and androgenicity testing are not automatically
12 interpreted as evidence of toxicity.

13 *2.2.1 General toxicity*

14 The European Union (2) reported there were no adequate studies for assessing acute toxicity of bisphenol
15 A in humans.

16 In an acute toxicity study in rats orally dosed with bisphenol A at ≥ 2000 mg/kg bw, clinical signs
17 included lethargy, prostration, hunched posture, and piloerection (reviewed by the European Union (2)).
18 Gross signs in animals that died included pale livers and hemorrhage in the gastrointestinal tract. In a
19 study in which male and female rats were subjected to whole body inhalation exposure to 170 mg/m³
20 bisphenol A dust for 6 hours, there were no gross signs of toxicity (reviewed by the European Union (2)).
21 Effects observed in the respiratory tract at 2 but not 14 days following exposure included slight
22 inflammation of nasal epithelium and slight ulceration of the oronasal duct. LD_{50s} reported in studies with
23 oral, dermal, inhalation, or ip exposure are summarized in Table 46. The European Union (2) concluded
24 that bisphenol A is of low acute toxicity through all exposure routes relevant to humans.
25
26
27
28

29 **Table 46. LD_{50s} for Bisphenol A**

Species	Exposure route	LD ₅₀ (mg/kg bw)
Rat	Oral	3300–4100 ^a
		5000 ^b
		3250 ^c
Mouse	Inhalation	>170 mg/m ³ b
	Oral	4100–5200 ^a
Guinea pig	ip	2400 ^c
	Oral	150 ^c
Rabbit	Oral	4000 ^c
	Dermal	2230 ^{b,c}
		> 2000 ^b
		3 mL/kg ^c

^aNational Toxicology Program (NTP) (127).

^bReviewed by the European Union (2).

^cReviewed in ChemIDplus (1).

30 The European Union (2) noted limited anecdotal data reporting skin, eye, and respiratory tract irritation in
31 workers exposed to bisphenol A, but concluded that the reports were of uncertain reliability. It was noted
32 that a recent, well-conducted study in rabbits demonstrated that bisphenol A is not a skin irritant. Other
33 studies conducted in rabbits demonstrated eye irritation and damage, and it was concluded the bisphenol
34 A can potentially cause serious eye damage. Slight respiratory tract inflammation occurred in rats
35 inhaling ≥ 50 mg/m³ bisphenol A, and it was concluded that bisphenol A had limited potential for
36

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1 respiratory irritation. Based on the results of the studies described above, the European Union concluded
2 that bisphenol A is not corrosive.

3
4 The European Union (2) reviewed studies examining possible sensitization reactions in humans exposed
5 to products containing bisphenol A, and those studies reported mixed results. In studies reporting positive
6 findings, it was unclear if bisphenol A or epoxy resins were the cause of hypersensitivity. Cross-
7 sensitization responses in individuals exposed to compounds similar to bisphenol A were also reported.
8 Animal studies were determined unreliable for assessing sensitization. Based on the results of human
9 studies, it was concluded that bisphenol A may have potential for sensitization in individuals exposed to
10 resins. Human studies suggested that bisphenol A can induce dermal photosensitization responses.
11 Photosensitization studies in mice resulted in reproducible positive results. Mechanistic studies in mice
12 suggested that sensitization occurs through an immune-mediated process. The overall conclusion of the
13 European Union was that it was somewhat unclear if bisphenol A induces orthodox skin sensitization,
14 photosensitization, or responses in individuals previously sensitized to another substance, such as epoxy
15 resins. No information was available on potential respiratory sensitization by bisphenol A.

16
17 The European Union (2) summarized systemic toxicity reported in subchronic, chronic, and reproductive
18 toxicity studies of rats, mice, and dogs. CERHR also reviewed the studies that examined reproductive
19 organs, and those studies are summarized in detail in the appropriate section of this report. A relevant
20 study by Yamasaki et al. (128) was published subsequent to the European Union review and was
21 reviewed in detail by CERHR.

22
23 In studies reviewed by the European Union (2) and in a study by Yamasaki et al. (128), rats were orally
24 exposed to bisphenol A for periods of 28 days to 2 years. Cecal enlargement occurring at doses ≥ 25
25 mg/kg bw/day was the most frequently observed effect in those studies. Histological alteration in the
26 cecum consisting of mucosal hyperplasia was only reported in one study at doses ≥ 200 mg/kg bw/day.
27 Histopathological changes in liver and kidney were reported at doses ≥ 500 mg/kg bw/day. The changes in
28 liver were characterized by prominent hepatocyte nuclei or inflammation. Histopathology in kidney was
29 characterized by renal tubule degeneration or necrosis. Testicular toxicity (degeneration of seminiferous
30 tubules and arrested spermatogenesis) was observed in 1 study at doses ≥ 235 mg/kg bw/day.

31
32 The European Union (2) found subchronic and chronic studies conducted by the NTP (127) to be the only
33 reliable studies for assessing systemic toxicity in mice orally exposed to bisphenol A. The liver was found
34 to be the target organ of toxicity, with multinucleated giant hepatocytes observed in male mice exposed to
35 ≥ 120 mg/kg bw/day and female mice exposed to 650 mg/kg bw/day.

36
37 In a 90-day dietary study in dogs reviewed by the European Union (2), an increase in relative liver weight
38 with no accompanying histopathological alterations was found to be the only effect at doses ≥ 270 mg/kg
39 bw/day.

40
41 In a subchronic inhalation exposure study in rats reviewed by the European Union (2), cecal enlargement
42 as a result of distention by food was observed at ≥ 50 mg/m³. Also observed at ≥ 50 mg/m³ were slight
43 hyperplasia and inflammation of epithelium in the anterior nasal cavity.

44
45 A limited number of repeat-dose systemic toxicity studies were summarized in detail by CERHR because
46 they included examination of reproductive organs. Those studies are summarized in detail below.

47
48 NTP (127), conducted acute, subacute, and subchronic bisphenol A toxicity studies in F344 rats and
49 B6C3F₁ mice. Animals were randomly assigned to treatment groups. Purity of bisphenol A was $<98.2\%$.
50 Concentration and stability of bisphenol A in feed were verified. In acute studies, single doses of
51 bisphenol A in a 1.5% acacia vehicle were administered by gavage to 5 rats/group/sex at doses of 2150,

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1 3160, 4640, or 6810 mg/kg bw/day and 5 mice/group/sex at 1470, 2150, 3160, 4640, 6810, or 10,000
2 mg/kg bw. LD₅₀ values for that study are summarized in Table 46.

3
4 In a 14-day repeat dose study, survival and body weight gain were evaluated in 5 rats and mice/sex/group
5 that were fed diets containing bisphenol A at 0, 500, 1000, 2500, 5000, or 10,000 ppm. Survival was
6 unaffected by treatment. Weight gain was reduced by 60% or more in male rats exposed to ≥ 2500 ppm
7 and 40% or more in female rats exposed to ≥ 5000 ppm bisphenol A. Survival and weight gain in mice
8 were not affected by Bisphenol A exposure.

9
10 In subchronic studies, 10 rats and mice/sex/group were exposed to bisphenol A in diet for 13 weeks.
11 Dietary doses were 0, 250, 500, 1000, 2000, or 4000 ppm for rats and 0, 5000, 10,000, 15,000, 20,000, or
12 25,000 ppm for mice. A review by the European Union (2) estimated bisphenol A intake at 0, 25, 50, 100,
13 200, and 400 mg/kg bw/day for rats, 0, 600, 1200, 1800, 2400, and 3000 mg/kg bw in male mice, and 0,
14 650, 1300, 1950, 2600, and 3250 mg/kg bw/day in female mice. Animals were observed and weighed
15 during the study and killed and necropsied on the 91st day of the study. **[Histopathological evaluations
16 were conducted but it was not clear if all dose groups and all animals/dose group were examined.
17 There was no mention of statistical analyses.]** In rats, the only deaths occurred in 2/10 males of the
18 1000 ppm group. Weight gain was reduced by 18% or more in male rats and 10% or more in female rats
19 exposed to ≥ 1000 ppm. There were no effects on feed intake. Hyaline masses in the bladder lumen were
20 not observed in control male rats but were observed in 5 of 10 males exposed to 250 ppm, 3 of 10
21 exposed to 500 ppm, 3 of 10 exposed to 1000 ppm, 6 of 10 exposed to 2000 ppm, and 4 of 10 exposed to
22 4000 ppm. Cecal enlargement, which was observed in rats at a rate of 60–100% in each dose group with
23 the exception of females exposed to 250 ppm was considered to be treatment-related. No histological
24 alterations were observed in the cecum. Death in mice was limited to 2 of 10 females in the 5000 ppm
25 group. Weight gain was reduced by at least 14% in male mice exposed to $\geq 15,000$ ppm. Non-dose-related
26 decreases in weight gain of 17% or more occurred in female mice of all dose groups. A dose-related
27 increase in multinucleated giant hepatocytes was observed in all dose groups of male mice; the only
28 incidence data reported for multinucleated giant hepatocytes were 0 of 10 female controls and 9 of 10
29 male mice of the 25,000 ppm group. **[A complete set of data for histopathological findings was not
30 presented for rats or mice.]**

31
32 Yamasaki et al. (128) examined the effects of bisphenol A exposure on male and female CD rats in a
33 study conducted according to Good Laboratory Practices (GLP). **[Because this study included a
34 number of reproductive organ and hormone endpoints, it is also discussed in Sections 4.2.1.1 and
35 4.2.2.1.]** Rats were fed a commercial diet (MF Oriental Yeast Co.) and housed in stainless steel wire-
36 mesh cages. Rats were grouped according to body weight and then randomly assigned to treatment groups.
37 Ten 7-week-old rats/sex/group were gavaged with bisphenol A at 0 (olive oil vehicle), 40, 200, or 1000
38 mg/kg bw/day for 28 days. Due to the death of 1 animal exhibiting clinical signs in the 1000 mg/kg
39 bw/day group, the high dose was reduced to 600 mg/kg bw/day on study day 8. In an additional study,
40 rats were exposed to ethinyl estradiol at 0, 10, 50, or 200 $\mu\text{g}/\text{kg}$ bw/day for 28 days. Endpoints examined
41 during the study were clinical signs, body weight gain, and food intake. Estrous cyclicity was examined in
42 females for 2 weeks beginning on study day 15. Males were killed on study day 29 and females were
43 killed in diestrus on study day 30, 31, or 32. Hematology and clinical chemistry endpoints were assessed,
44 and blood hormone levels were measured by immunoassay systems. Sperm motility and viability were
45 evaluated. Organs, including those of the reproductive system, were weighed and subjected to
46 histopathological evaluation. With the exception of the testis and epididymis, which were fixed in Bouin
47 solution, the organs were fixed in 10% neutral buffered formalin. Statistical analyses included Bartlett test
48 for homogeneity of variance, ANOVA, Dunnett test, and/or Kruskal-Wallis test.

49
50 One female and 3 males from the high-dose group died; clinical signs observed in those animals included
51 soft stools, decreased mobility, reduced respiration rate, and decreased body temperature. Soft stools were

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1 also observed in surviving males and females of the mid- and high-dose groups. Results of the study are
 2 summarized in Table 47. Terminal body weights were lower in females of the mid- and high-dose groups
 3 and males of the high-dose group. During the first week of study, food intake was decreased in both sexes
 4 of the mid- and high-dose group. **[Data wer not shown by study authors.]** As noted in Table 47
 5 , some alterations in hematological and clinical chemistry endpoints were observed, mainly at the high
 6 dose. **[Data were not shown by study authors.]** There were no treatment-related abnormalities in sperm
 7 or alterations in blood levels of thyroid hormones, follicle stimulating hormone (FSH), luteinizing
 8 hormone (LH), 17 β -estradiol, prolactin, or testosterone. Number of females with diestrus lasting 4 or
 9 more days was increased in the high-dose group. Changes in relative organ weights **[assumed to be**
 10 **relative to body weight]** included decreased heart weight in females from the mid- and high-dose groups.
 11 At the high dose, there were decreases in relative weight of ventral prostate and increases in relative
 12 weights of testis and adrenals in males and thyroid and liver in females. Gross signs observed in animals
 13 that died included enlarged kidney, elevated mucosa in the forestomach, and atrophied spleen and
 14 thymus. In surviving animals, the cecum was enlarged in the mid- and high-dose group and forestomach
 15 mucosa was elevated in the high-dose group. As described in more detail in Table 47, histopathological
 16 alterations were observed in the intestine, cecum, and colon of males and intestine and cecum of females
 17 in the mid and high dose groups. Additional histopathological alterations were observed in the high-dose
 18 group in the kidney, forestomach, and adrenals of males and females and livers of females. The study
 19 authors concluded that change in estrous cyclicity was the only useful endpoint for evaluating the
 20 endocrine-mediated effects of bisphenol A.

21
 22 Male rats from the mid- and high-dose ethinyl estradiol groups experienced decreased prostate, seminal
 23 vesicle, and pituitary weights, increased testis weight, and histopathological alterations in prostate,
 24 seminal vesicle, mammary gland, and testis. Females from the mid- and high-dose ethinyl estradiol group
 25 experienced alterations in estrous cyclicity. Females from the high-dose group experienced decreased
 26 ovary weight, increased uterine weight, and histopathological changes in ovary, uterus, and vagina.
 27

28 **Table 47. Toxicological Effects in Rats Gavaged With Bisphenol A for 28 Days**

Endpoint	Bisphenol A dose (mg/kg bw/day)		
	40	200	600–1000 ^b
Males			
Terminal body weight	↔	↔	↓7%
Relative testes weight	↔	↔	↑21%
Ventral prostate weight	↔	↔	↓28%
Relative adrenal weight	↔	↔	↑19%
Feed intake ^a	↔	↓	↓
Prothrombin time ^a	↔	↔	↑
Glutamic-oxaloacetic transaminase ^a	↔	↑	↑
Triglyceride ^a	↔	↔	↓
Alkaline phosphatase ^a	↔	↔	↑
γ -Glutamyl transpeptidase ^a	↔	↔	↑
Chloride ^a	↔	↔	↑
Renal tubular degeneration and necrosis	0/10	0/10	7/7
Forestomach squamous epithelial cell hyperplasia	0/10	0/10	6/7
Lacteal dilatation in duodenum	0/10	10/10	2/7
Lacteal dilation in jejunum	0/10	0/10	2/7
Mucosal hyperplasia in cecum	0/10	3/10	6/7
Mucosal hyperplasia in colon	0/10	2/10	7/7
Adrenal cortical vacuolization	0/10	0/10	3/7
Females			

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Endpoint	Bisphenol A dose (mg/kg bw/day)		
	40	200	600–1000 ^b
Terminal body weight	↔	↓7%	↓5%
Relative thyroid weight	↔	↔	↑22%
Relative liver weight	↔	↔	↑10%
Relative heart weight	↔	↓9%	↓15%
Feed intake ^a	↔	↓	↓
Hemoglobin and hematocrit values ^a	↔	↔	↓
Cholinesterase ^a	↔	↓	↓
Glutamic-oxaloacetic transaminase ^a	↔	↔	↑
Albumin and albumin:globulin rats ^a	↔	↔	↓
Diestrus ≥ 4 days	0/10	0/10	3/9
Prominent hepatocyte nuclei	0/10	0/10	4/9
Renal tubular degeneration and necrosis	0/10	0/10	9/9
Forestomach squamous epithelial cell hyperplasia	0/10	0/10	5/9
Lacteal dilatation in duodenum	0/10	7/10	6/9
Mucosal hyperplasia in cecum	0/10	6/10	4/9
Adrenal cortical vacuolization	0/10	0/10	3/9

↑,↓ Statistically significant increase, decrease compared to controls; ↔ no statistically significant effects compared to controls.

^aData were not shown by study authors.

^bThe dose was 1000 mg/kg bw/day at the beginning of the study, but was decreased to 600 mg/kg bw/day in the second week of the study due to excessive toxicity.

From Yamasaki et al. (128).

1
2 General Electric (129) conducted a subchronic toxicity study in Beagle dogs orally dosed with bisphenol
3 A [**purity not reported**]. Dogs weighing 6.5–13.4 kg were housed in metal metabolism cages and fed
4 Purina Dog Chow. During a 90-day period, 4 dogs/sex/group were given feed containing bisphenol A at
5 0, 1000, 3000, or 9000 ppm. The European Union (2) estimated bisphenol A intake at 0, 28, 74, or 261
6 mg/kg bw/day in males and 0, 31, 87, or 286 mg/kg bw/day in females. Dogs were observed for body
7 weight gain, food, intake, and clinical signs. Ophthalmoscopic examination was conducted prior to and
8 following the treatment period. Hematology, clinical chemistry, and urinalysis evaluations were
9 conducted prior to treatment and at 1, 2, and 3 months into the study. Dogs were killed at the end of the
10 treatment period. Organs were weighed and fixed in 10% neutral buffered formalin. Histopathological
11 evaluations were conducted in organs from the control and high-dose groups; prostate, uterus, testis, and
12 ovary were among organs evaluated. [**Procedures for statistical analyses were not described.**] No
13 treatment-related clinical signs, ophthalmological changes, or death were observed during the study.
14 Bisphenol A treatment did not affect body weight gain or food intake. There were no treatment-related
15 effects on hematology, biochemistry, or urinalysis. Relative liver weight was significantly increased [**by**
16 **18% in males and 26% in females**] in the high-dose group, and the study authors considered the effect
17 to be treatment-related. No treatment-related gross or histopathological lesions were observed in the high-
18 dose group.

19
20 Nitschke et al. (130) conducted a subchronic inhalation toxicity test with bisphenol A in F344 rats. Rats
21 were fed Purina Certified Rodent Chow #5002 and housed in stainless steel wire cages. At 7 weeks of
22 age, rats were stratified according to body weight and randomly assigned to treatment groups. Thirty
23 rats/sex/group received whole-body exposures to polycarbonate grade bisphenol A dust (99.7% purity) at
24 0, 10, 50, or 150 mg/m³ for 6 hours/day, 5 days/week, for 13 weeks. Mass median aerodynamic diameter
25 of bisphenol A dust was measured at ≤5.2 microns. Stability and concentrations of bisphenol A were
26 verified. Rats were observed for clinical signs, body weight gain, and food intake. Ten rats/sex/group in
27 each time period were killed and necropsied on the day following and at 4 and 12 weeks following

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1 exposure. At each necropsy period, hematological and clinical chemistry endpoints were examined. The
2 lungs, brain, kidneys, and testes were weighed. Numerous organs were preserved in 10% phosphate-
3 buffered formalin. In most cases, histological examinations were conducted in organs from the control
4 and high-dose groups. Respiratory organs and organs with lesions or signs of toxicity were histologically
5 examined at all dose levels. Included among organs undergoing histopathological examination
6 immediately after the exposure period were the epididymis, mammary gland, ovary, oviduct, prostate,
7 seminal vesicles, testis, uterus, and vagina. No reproductive organs were examined following the recovery
8 periods. Statistical analyses included Bartlett's test, ANOVA, Dunnett test, Wilcoxon Rank-Sum test, and
9 Bonferroni correction for multiple comparisons. Gross pathology and histopathology data did not appear
10 to have been statistically analyzed.

11
12 During the exposure period, a reddish material around the nose (most likely porphyrin) was observed in
13 2–10 of 10 animals/sex in the 50 and 150 mg/m³ groups. Perineal soiling was observed in 2 of 10 females
14 in the 10 mg/m³ group and 9–10 of 10 animals/sex in the 50 and 150 mg/m³ groups. Decreased body
15 weight gain during treatment was observed in males from all dose groups and females in the 50 and 150
16 mg/m³ groups. Immediately following the treatment period, terminal body weights were reduced by ~5%
17 in males and ~11% in females from the 150 mg/m³ group. **[Body weights were ~4% lower in males
18 from the 50 mg/m³ group.]** No differences in feed intake were observed at this or any other time period
19 in the study. The only hematological effect observed was slightly increased hemoglobin in males exposed
20 to 10 mg/m³, but the study authors did not consider the effect to be biologically significant. Clinical
21 chemistry observations in the 150 mg/m³ group included decreased serum glutamic pyruvic transaminase
22 activity, serum glutamic oxaloacetic transaminase activity, and glucose in males and decreased total
23 protein and albumin and increased alkaline phosphatase activity in females. Alkaline phosphatase activity
24 was also increased in females exposed to 50 mg/m³. The study authors did not consider any of the clinical
25 chemistry changes to be biologically significant. Absolute liver weight was decreased in males exposed to
26 ≥10 and 150 mg/m³, and relative brain weight was increased in females exposed to ≥50 mg/m³. Additional
27 organ weight changes observed in females from the 150 mg/m³ group included decreased absolute liver
28 and kidney weights and increased relative lung weights. Because the organ weight changes were not
29 associated with microscopic changes in organs, the study authors concluded that the effects reflected
30 decreases in body weight and were not toxicologically significant. Cecal size was increased as a result of
31 distention by food in all (10/dose/sex) males and females exposed to ≥50 mg/m³, and the effect was
32 considered to be treatment-related. No histopathological alterations were observed for cecal wall
33 morphology. Hemolyzed blood was observed in the stomachs of 3–7 of 10 males/group exposed to 50 and
34 150 mg/m³, but there were no signs of histopathological alterations in the gastrointestinal tract. Slight
35 histopathological alterations, consisting of hyperplasia in stratified squamous and ciliated epithelium
36 lining and inflammation of submucosal tissues was observed in the anterior nasal cavities of all
37 (10/dose/sex) males and females exposed to ≥50 mg/m³. Slight-to-moderate hyperplasia of goblet cells
38 was also observed in the lateral nasal wall. No other treatment-related histopathological alterations were
39 observed, including in reproductive organs.

40
41 During the 4-week recovery period, body weights remained lower in males and females of the 50 and 150
42 mg/m³ groups. At 4 weeks following exposure, terminal body weights of males and females in the 150
43 mg/m³ group were ~6% lower than control values. A decrease in white blood cell count in females from
44 the 10 and 150 mg/m³ groups was the only hematological effect observed. The clinical chemistry effects
45 that were somewhat consistent with effects observed immediately following treatment were increased
46 alkaline phosphatase activity in females exposed to 10 and 150 mg/m³ and decreased serum glutamic
47 pyruvic activity transaminase activity in females exposed to 150 mg/m³; the study authors did not
48 consider the clinical chemistry changes to be treatment related. The study authors concluded that an
49 increase in relative brain weight in males of the 150 mg/m³ group was related to decreased body weights
50 in those animals. Enlarged cecal size was observed in 5 of 10 males of the 150 mg/m³ group, a decreased
51 incidence compared to the period immediately following treatment. Nasal histopathology was observed in

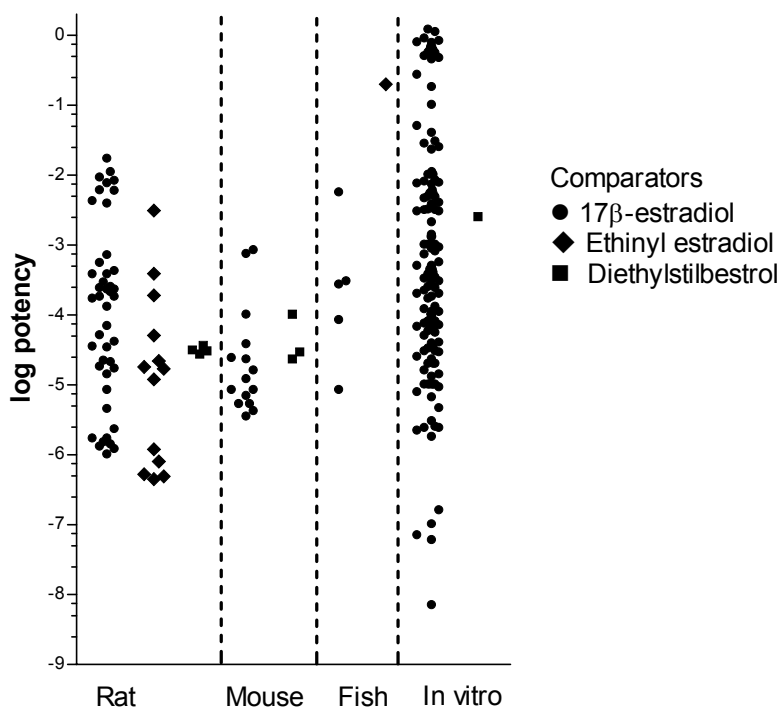
2.0 General Toxicology and Biological Effects

1 the 150 mg/m³ but was reduced in magnitude and severity compared to rats observed immediately
2 following exposure.

3
4 In rats examined following 12 weeks of recovery, body weights of males in the 150 mg/m³ group
5 remained lower than controls, and terminal body weight was decreased by ~6%. An increase in white
6 blood cell counts but not differential counts was observed in male rats of the 10 and 150 mg/m³ group.
7 The only clinical chemistry finding consistent with earlier observations was decreased total protein and
8 globulin in females from the 150 mg/m³ group, but the study authors did not consider the effect to be
9 biologically significant. Organ weight changes in the 150 mg/m³ group included decreased absolute
10 kidney and lung weights in males and decreased absolute and relative kidney weights in females. No
11 histopathological alterations were observed in kidney or lung. No other gross or histopathological
12 alterations were observed, including cecal enlargement and nasal histopathology, which were observed at
13 earlier time periods.

15 2.2.2 Estrogenicity

16 The first identification of bisphenol A as an estrogen has been attributed to Dodds and Lawson (131), who
17 reported that 100 mg injected by an unspecified route twice daily for 3 days resulted in maintenance of 5
18 of 5 rats in vaginal estrus for 40 days. The estrogenicity of bisphenol A has since been evaluated using
19 several different kinds of assays. In vitro studies are summarized in Table 48, and in vivo studies are
20 summarized in Table 49 using comparisons with 17 β -estradiol, ethinyl estradiol, diethylstilbestrol, and, in
21 one study, estrone. There is considerable variability in the results of these studies with the estrogenic
22 potency of bisphenol A ranging over about 8 orders of magnitude (Figure 2).

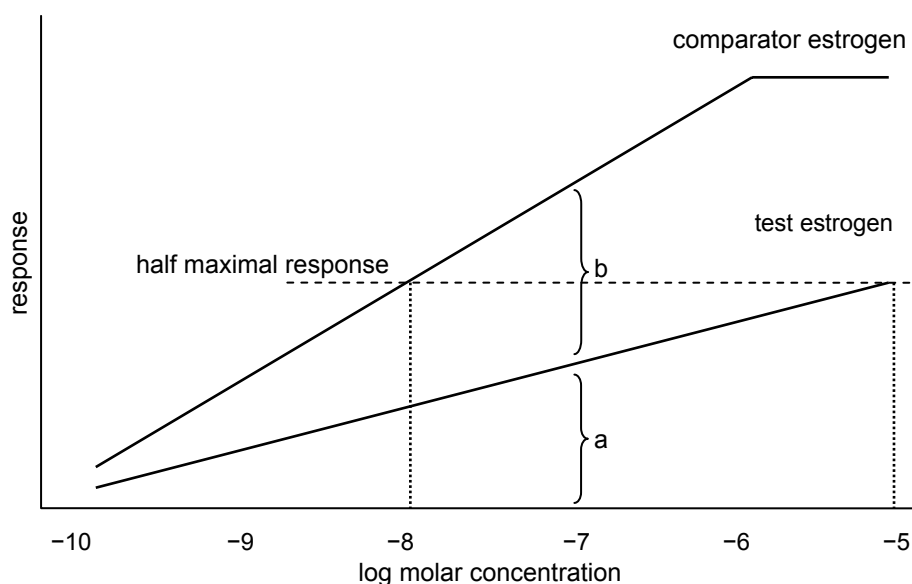


26
27 **Figure 2. Estrogenic potency of bisphenol a compared to other estrogens.**

28 Each data point represents 1 bisphenol A study in which bisphenol A was compared to a reference
29 estrogen in rats, mice, fish, or in vitro. Data summarized from Table 48 and Table 49.
30

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1 The most common method of comparing potency is to test responses over a range of concentrations and
2 to compare the concentrations producing the half-maximal (or other fractional) response of the
3 comparator estrogen. An alternative is to compare the magnitude of the response at an equimolar
4 concentration of the 2 estrogens. The difference in these two methods is illustrated in Figure 3. An
5 example of the difference in potency estimations according to comparison method is the study of
6 Vivacqua et al. (47), in which the fold-increase in reporter activity for an estrogen-responsive gene was
7 compared over a range of concentrations for bisphenol A and for 17 β -estradiol. This study's Figure 3
8 presents curves analogous to Figure 3, but also presents a bar graph comparing response of the reporter at
9 a 10⁻⁷ M concentration of each estrogen. Based on the half-maximal response to 17 β -estradiol, bisphenol-
10 A appeared 1000 times less potent than 17 β -estradiol, but based on the fold-difference in reporter activity
11 at 10⁻⁷ M, bisphenol A was about half as potent. Data for other estrogenicity comparisons in this paper
12 and in many other papers are presented only using bar graphs comparing responses at the same molar
13 concentrations of the 2 estrogens, thereby overestimating the estrogenic potency of bisphenol A compared
14 to studies in which comparisons are based on the half-maximal response.
15
16
17
18



19
20 **Figure 3. Alternative approaches to comparing estrogenic potency.**

21 In this example, the half-maximal response to the comparator estrogen occurs at 10⁻⁸ M. A
22 similar response occurs with the test estrogen at 10⁻⁵ M, suggesting a 1000-fold difference
23 in potency. If the magnitudes of response at equimolar concentrations are compared, the
24 apparent potency may be much different. The response to the test estrogen at 10⁻⁷ M (a) is
25 about half the response to the comparator estrogen at 10⁻⁷ M (a + b).
26
27

28 Competitive binding assays, which evaluate the concentration at which bisphenol A displaces labeled
29 17 β -estradiol from ER, are summarized in the top part of Table 48. The receptor binding of bisphenol A
30 in these assays varies over 3 orders of magnitude. Bisphenol A competes for human ER binding at molar
31 concentrations 20–10,000 times that of the native ligand. When bisphenol A binding to ER α and ER β
32 was compared in the same study, 2 reports found little difference by receptor subtype (132, 133), and 3
33 studies found binding to ER β to be 4, 10, 47, and 254 times greater than binding to ER α (134–138). Yeast

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1 reporter systems, which reflect activation of post-receptor pathways, show less variability; these studies
2 show bisphenol A activity to be 10,000–26,000 times less than that of 17β-estradiol.

3
4 Some variability in estimating bisphenol A potency appears to be due to differences between laboratories.
5 Andersen et al. (139) reported results from 3 laboratories that evaluated the proliferative response of
6 MCF-7 breast cancer cells to bisphenol A. The laboratories, which were in the US, Spain, and Denmark,
7 were sent samples of the same stock of bisphenol A, 17β-estradiol, and MCF-7 cells. Procedures were
8 similar in the labs, although 2 different counting methods were used. The bisphenol A potencies relative
9 to 17β-estradiol were 5×10^{-7} , 3×10^{-6} , and 1×10^{-5} . Laboratory variability may underlie some of the
10 large differences in cell-based assays for ER activation; in those studies bisphenol A molar potency
11 compared to 17β-estradiol were reported to vary by over 7 orders of magnitude (Table 48). Another
12 explanation for this wide range of reported values is the difference in defining relative potency in some
13 assays, as discussed above.

14
15 A study using ERα- and ERβ-reporting systems in 3 human cell lines found that bisphenol A had a small
16 antagonistic effect on ERα activation in the presence of 17β-estradiol in human embryonal kidney and
17 endometrial carcinoma cells (140). There were no significant interactions between bisphenol A and 17β-
18 estradiol on ERα activation in human osteosarcoma cells or on ERβ activation in any tested cell type. By
19 contrast, a study using a recombinant yeast assay for ERα activation found 17β-estradiol and bisphenol A
20 to have additive effects (141), and a study using MCF-7 cell proliferation found 17β-estradiol and
21 bisphenol A to have synergistic effects (142).

22
23 The data in Table 48 are applicable only to unconjugated bisphenol A. Estrogenic activity has not been
24 identified for bisphenol A glucuronide (138) or sulfate (143).

25
26 **Table 48. In Vitro Estrogenicity Testing of Bisphenol A**

Endpoint	Molar potency relative to 17β-estradiol	Reference
<i>Binding assays</i>		
Frog liver cytosol binding	$[1.4 \times 10^{-3}]$	Lutz and Kloas (144)
Carp liver cytosol binding	$[1.3 \times 10^{-3}]$	Segner et al. (145)
Rainbow trout ER binding	5.8×10^{-5}	Olsen et al. (146)
Rainbow trout ER binding	2.1×10^{-3}	Matthews et al. (147)
Anole ER binding	1.3×10^{-3}	Matthews et al. (147)
Chicken ER binding	4.4×10^{-4}	Matthews et al. (147)
Mouse ERα binding	8.6×10^{-5}	Matthews et al. (147)
Mouse uterine cytosol binding	$[1.2 \times 10^{-4}]$	Matthews et al. (138)
Rabbit uterine ER binding	$[1.3 \times 10^{-5}]$	Andersen et al. (139)
Rat uterine cytosol binding	$\sim 5 \times 10^{-4}$	Krishnan et al. (148)
Rat uterine cytosol binding	8×10^{-5}	Blair et al. (149)
Rat uterine cytosol binding	$1-2 \times 10^{-4}$	Kim et al. (150)
Rat ERα binding	$[2.5 \times 10^{-4}]$	Strunck et al. (151)
ER binding in rat lactotrophs	$1-10 \times 10^{-5}$	Chun and Gorski (152)
Rat ERα binding	5×10^{-4}	Kuiper et al. (153)
Rat ERβ binding	3.3×10^{-4}	Kuiper et al. (153)
Rat uterine ERα and β binding	6.2×10^{-5}	Washington et al. (154)
Rat uterine Type II estrogen-binding site	4×10^{-3}	Washington et al. (154)
ER binding in MCF-7 lysates	1×10^{-2}	Dodge et al. (155)

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Endpoint	Molar potency relative to 17β-estradiol	Reference
Human ERα binding	4×10^{-4}	Bolger et al. (156)
Human ERα binding	1×10^{-4}	Kuiper et al. (132)
Human ERβ binding	1×10^{-4}	Kuiper et al. (132)
Human ER binding	5.6×10^{-4}	Perez et al. (157)
Human ER binding	$[1.3 \times 10^{-4}]$	Andersen et al. (139)
ER binding in ECC-1 cells	3×10^{-3}	Bergeron et al. (158)
Human ERα binding	8×10^{-5}	Matthews et al. (147)
Human ERα binding	$[2.5 \times 10^{-3}]$	Nakagawa and Suzuki (159)
diethylstilbestrol		
Human ERα binding	7.3×10^{-4}	Routledge et al. (135)
Human ERβ binding	7.5×10^{-3}	Routledge et al. (135)
Human ER binding	$[7.1 \times 10^{-5}]$	Sheeler et al. (160)
Human ERα binding	$[8 \times 10^{-5}]$	Matthews et al. (138)
Human ERβ binding	$[3.8 \times 10^{-3}]$	Matthews et al. (138)
Human ERα binding	5×10^{-2}	Paris et al. (133)
Human ERβ binding	4×10^{-2}	Paris et al. (133)
Human ER binding	$[3 \times 10^{-4}]$	Strohecker et al. (161)
Human ERα binding	$[2.4 \times 10^{-4}]$	Seidlová-Wuttke et al. (136, 137)
Human ERβ binding	$[2.8 \times 10^{-2}]$	
Human ERα binding	$[1.1 \times 10^{-4}]$	Takemura et al. (134)
Human ERβ binding	$[4.4 \times 10^{-4}]$	Takemura et al. (134)
Human ER binding	3.15×10^{-3}	Olsen et al. (146)
<i>Recombinant yeast reporter systems</i>		
Human ER activation	5×10^{-5}	Coldham et al. (162)
Human ER activation	6.7×10^{-5}	Gaido et al. (163)
Human ER activation	$[2.5 \times 10^{-5}]$	Harris et al. (164)
Human ER activation	$[4-8 \times 10^{-5}]$	Andersen et al. (139)
Human ER activation	$[3.9 \times 10^{-5}]$	Sheeler et al. (160)
Human ER activation	$\sim 1 \times 10^{-4}$	Sohoni and Sumpter (165)
Human ER activation	3.7×10^{-5}	Metcalf et al. (166)
ERα activation	6.2×10^{-5}	Silva et al. (167)
ERα activation	$[1 \times 10^{-4}]$	Nishihara et al. (168)
ERα activation	$[\sim 1 \times 10^{-4}]$	Beresford et al. (169)
Human ERα	$[3.3 \times 10^{-5}]$	Rajapakse et al. (141)
Human ERα, no microsomes	$[5.5 \times 10^{-5}]$	Elsby et al. (113)
Human ERα, human liver microsomes	$[6.6 \times 10^{-6}]$	Elsby et al. (113)
ER activation	$\sim 10^{-5}$	Chen et al. (170)
Human ER activation	$[8.1 \times 10^{-5}]$	Segner et al. (145)
Human ER activation	9×10^{-5}	Li et al. (171)
ERα activation	$[4 \times 10^{-5}]$	Singleton et al. (172)
Human ERα, with denatured rat S9	$[2.4 \times 10^{-6}]$	Yoshihara et al. (173)
Human ERα, with active rat S9	$[9.2 \times 10^{-6}]$	Yoshihara et al. (173)
Human ERα, with denatured mouse S9	$[3.0 \times 10^{-6}]$	Yoshihara et al. (173)
Human ERα, with active mouse S9	$[7.8 \times 10^{-6}]$	Yoshihara et al. (173)
Human ERα, with denatured monkey S9	$[2.4 \times 10^{-6}]$	Yoshihara et al. (173)

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Endpoint	Molar potency relative to 17 β -estradiol	Reference
Human ER α , with active monkey S9	[6.0 $\times 10^{-6}$]	Yoshihara et al. (173)
Human ER α , with denatured human S9	[2.2 $\times 10^{-6}$]	Yoshihara et al. (173)
Human ER α , with active human S9	[4.6 $\times 10^{-6}$]	Yoshihara et al. (173)
Human ER α activity	[2.3 $\times 10^{-5}$]	Terasaki et al. (174)
Medaka ER α activity	[3.3 $\times 10^{-4}$]	Terasaki et al. (174)
“Estrogenic activity”	3.4×10^{-5}	Kawagoshi et al. (175)
<i>Other cell-based recombinant reporter systems</i>		
ER activation in trout gonad cell line	5.4×10^{-3}	Ackerman et al. (176)
Mouse ER α in HeLa cells	[<1 $\times 10^{-5}$]	Ranhotra et al. (177)
Mouse ER β in HeLa cells	[~1 $\times 10^{-2}$]	Ranhotra et al. (177)
HepG2 cells, human ER α	[3.0 $\times 10^{-3}$]	Snyder et al. (103)
HepG2 cells, human ER β	[1.1 $\times 10^{-2}$]	Snyder et al. (103)
Rat ER α in HeLa cells	[1.6 $\times 10^{-7}$]	Yamasaki et al. (178)
ER activation in HeLa cells	[8.8 $\times 10^{-4}$]	Takahashi et al. (179)
ER α activation in HeLa cells	[2.5 $\times 10^{-2}$]	Hiroi et al. (180)
ER β activation in HeLa cells	[2.3 $\times 10^{-2}$]	Hiroi et al. (180)
ER α activation in HeLa cells	[6.1 $\times 10^{-1}$]	Vivacqua et al. (47)
ER β activation in HeLa cells	[5.6 $\times 10^{-1}$]	Vivacqua et al. (47)
ER α activation in HeLa cells	[7.7 $\times 10^{-1}$]	Recchia et al. (181)
ER β activation in HeLa cells	[1.2]	Recchia et al. (181)
ER α activation in T47D cells	[6.2–7.9 $\times 10^{-1}$]	Recchia et al. (181)
Proliferation in T47D cells	[6.6 $\times 10^{-1}$]	Recchia et al. (181)
Human ER in hepatoma cells	[3 $\times 10^{-2}$]	Gould et al. (182)
Human ER α , human embryonal kidney	[4.8 $\times 10^{-3}$]	Kurosawa et al. (140)
Human ER β , human embryonal kidney	[4.6 $\times 10^{-3}$]	Kurosawa et al. (140)
Human ER α , endometrial carcinoma	[5.4 $\times 10^{-3}$]	Kurosawa et al. (140)
Human ER β , endometrial carcinoma	[4.9 $\times 10^{-3}$]	Kurosawa et al. (140)
Human ER α , osteosarcoma	[7.3 $\times 10^{-3}$]	Kurosawa et al. (140)
Human ER β , osteosarcoma	[7.7 $\times 10^{-3}$]	Kurosawa et al. (140)
Human ER α , human hepatoma cells	[2.7 $\times 10^{-1}$]	Gaido et al. (183)
Human ER β , human hepatoma cells	[1.8 $\times 10^{-1}$]	Gaido et al. (183)
<i>MCF-7 cells</i>		
G6PD activity	[1 $\times 10^{-1}$]	Kim et al. (184)
Expression of proteins	[1 $\times 10^{-3}$]	Perez et al. (157)
Progesterone receptor mRNA	not increased at 10^{-6} M ^a	Diel et al. (185)
Androgen receptor mRNA	not decreased at 10^{-6} M ^a	Diel et al. (185)
Progesterone receptor	~2 $\times 10^{-4}$	Krishnan et al. (148)
ER binding, serum-free	3.3×10^{-4}	Samuelsen et al. (186)
ER binding, 100% human serum	1.7×10^{-4}	Samuelsen et al. (186)
ER binding	3.2×10^{-3}	Olsen et al. (187)
ER activation	[1.4 $\times 10^{-5}$]	Kitamura et al. (188)
ER α expression	[7.5 $\times 10^{-5}$]	Matthews et al. (138)
ER β expression	[1.8 $\times 10^{-4}$]	Matthews et al. (138)
ER α activation	[4.7–6.9 $\times 10^{-1}$]	Vivacqua et al. (47)
ER α activation	[5.5–6.7 $\times 10^{-1}$]	Recchia et al. (181)

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Endpoint	Molar potency relative to 17 β -estradiol	Reference
pS2 induction	[1.8 $\times 10^{-6}$]	Leffers et al. (189)
ER production	[7 $\times 10^{-8}$]	Olsen et al. (187)
Progesterone receptor production	[6.8 $\times 10^{-8}$]	Olsen et al. (187)
pS2 production	[10 ⁻⁷]	Olsen et al. (187)
pS2 mRNA	[1.1]	Vivacqua et al. (47)
pS2 mRNA	[8.9 $\times 10^{-1}$]	Recchia et al. (181)
Cathepsin D mRNA	[8.2 $\times 10^{-1}$]	Recchia et al. (181)
Transcription of human telomerase reverse transcriptase	[$\sim 10^{-2}$]	Takahashi et al. (179)
Proliferation	[3.8 $\times 10^{-4}$]	Krishnan et al. (148)
Proliferation	1 $\times 10^{-3}$	Brotons et al. (40)
Proliferation	1 $\times 10^{-4}$	Soto et al. (190)
Proliferation	[$\sim 1 \times 10^{-3}$]	Dodge et al. (155)
Proliferation	[1 $\times 10^{-4}$]	Perez et al. (157)
Proliferation	[9.8 $\times 10^{-4}$]	Schafer et al. (191)
Proliferation (3 different laboratories)	5–100 $\times 10^{-7}$	Andersen et al. (139)
Proliferation	6 $\times 10^{-5}$	Körner et al. (192)
Proliferation	3 $\times 10^{-5}$	Kim et al. (150)
Proliferation	[2.5 $\times 10^{-6}$]	Suzuki et al. (142)
Proliferation	2 $\times 10^{-5}$	Samuelsen et al. (186)
Proliferation	[9.2 $\times 10^{-4}$]	Nakagawa and Suzuki (159)
Proliferation	[$\sim 1 \times 10^{-3}$]	Shimizu et al. (143)
Proliferation	[7 $\times 10^{-9}$]	Diel et al. (185)
Proliferation	1.6 $\times 10^{-5}$	Olsen et al. (187)
Proliferation	[4.5–5 $\times 10^{-1}$]	Vivacqua et al. (47)
Proliferation	[1.1 $\times 10^{-4}$]	Strohecker et al. (161)
Proliferation	[6 $\times 10^{-1}$]	Recchia et al. (181)
Proliferation	2 $\times 10^{-5}$	Olsen et al. (146)
Proliferation, with denatured rat S9	[6.5 $\times 10^{-5}$]	Yoshihara et al (193)
Proliferation, with active rat S9	[3.4 $\times 10^{-4}$]	Yoshihara et al (193)
<i>Rat pituitary cells</i>		
Proliferation	1–10 $\times 10^{-6}$	Chun and Gorski (152)
Proliferation	[$\sim 8.4 \times 10^{-3}$]	Steinmetz et al. (194)
Prolactin release	1 $\times 10^{-5}$	Chun and Gorski (152)
Prolactin release (GH ₃ cell)	[6 $\times 10^{-3}$]	Steinmetz et al. (194)
Prolactin release (F344 pituitary)	2–10 $\times 10^{-4}$	Steinmetz et al. (194)
Prolactin gene expression	[$\sim 1 \times 10^{-3}$]	Steinmetz et al. (194)
<i>Rat uterine adenocarcinoma cells</i>		
Induction of complement C3 mRNA	[8 $\times 10^{-3}$]	Strunck et al. (151)
<i>Human uterine adenocarcinoma cells</i>		
Progesterone receptor mRNA/protein	[$\sim 1 \times 10^{-2}$]	Bergeron et al. (158)
Proliferation	no effect at 10 ⁻⁵ M	Bergeron et al. (158)
<i>Vitellogenin production, fish hepatocytes</i>		
Carp	1 $\times 10^{-4}$	Smeets et al. (195)
Carp	[3.1 $\times 10^{-3}$]	Segner et al. (145)
Carp	[1 $\times 10^{-5}$]	Letcher et al. (196)

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Endpoint	Molar potency relative to 17β-estradiol	Reference
Carp	$[3 \times 10^{-4}]$	Rankouhi et al. (197)
Trout	2×10^{-5}	Shilling et al. (198)
Trout	$[8 \times 10^{-4}]$	Segner et al. (145)
Trout	2.9×10^{-5}	Olsen et al. (146)
<i>Frog hepatocytes</i>		
Vitellogenin mRNA expression	$[\sim 1 \times 10^{-3}]$	Kloas et al. (199)
Vitellogenin production	no effect at 100 μM	Rankouhi et al. (200)
ER mRNA expression	$\sim 10^{-2}$	Lutz et al. (201)

^aProgesterone receptor was increased and androgen receptor was decreased by 17β-estradiol 10^{-10} M.

1
2 In vivo tests (Table 49) have been conducted principally in rats and mice. Most endpoints in these studies
3 involved the uterus, and effects on uterine weight in immature or ovariectomized animals are the most
4 commonly reported uterine endpoints. The potency of bisphenol A in increasing uterine weight varies
5 over ~4 orders of magnitude. Some of this variation may be related to the short half-life of bisphenol A.
6 Uterotrophic evaluations are typically performed 24 hours after the last dose of the test agent is
7 administered. Laws et al. (202) showed no significant effect of bisphenol A at doses ≤400 mg/kg bw/day
8 given orally on uterine wet weight assessed 24 hours after administering the last dose. When assessed 6
9 hours after the last oral dose, bisphenol A 200 mg/kg bw/day increased uterine wet weight to ~2.5 times
10 the control **[estimated from a graph]**, which was about the same as the increase produced by
11 administering 17β-estradiol 0.005 mg/kg bw/day sc. Increase in uterine weight 6 hours after treatment
12 represents fluid inhibition and not true tissue growth. A dose-related decrease in blotted uterine weight
13 and body weight, with no effect on weight-adjusted uterine weight, was shown in pubertal rats treated on
14 PND 22–42/43 with bisphenol A by gavage at 400 or 600 mg/kg bw/day (203).

15
16 For studies showing an increase in uterine weight after bisphenol A treatment, dose route affects
17 response; bisphenol A given by gavage produced approximately half the uterine weight increase as did
18 the same dose given sc (202). A greater response by the sc than oral route was also shown by Yamasaki et
19 al. (99), who showed a lowest effective bisphenol A dose of 8 mg/kg bw/day by the sc route and 160
20 mg/kg bw/day by the oral route. The greater activity of sc than oral bisphenol A is presumably due to
21 glucuronidation of the orally administered compound with consequent loss of estrogenicity (138). Not all
22 studies confirmed this greater effect of sc compared to oral bisphenol A on uterine weight. Ashby and
23 Tinwell (204) concluded that the magnitude of uterine weight response was similar for sc and oral routes.
24 **[The Expert Panel notes a greater numerical magnitude of response after sc than oral exposure in**
25 **most of the experiments reviewed in this report, and that statistical comparison of the dose routes**
26 **was not reported.]** Matthews et al. (138) found a similar increase in uterine weight in rats given sc or
27 oral bisphenol A at 800 mg/kg bw/day.

28
29 Nagel et al. (205, 206) noted that 17β-estradiol is extensively protein-bound in vivo and bisphenol A is
30 minimally protein-bound. They suggested that estrogenicity can be more accurately predicted by
31 considering the free fraction of a chemical in serum. **[The Expert Panel notes that Figure 2 does not**
32 **suggest that bisphenol A is more potent than 17β-estradiol in vivo than in vitro. The Expert Panel**
33 **also notes that Nagel et al. appeared to be referring primarily to prediction of developmental effects**
34 **in the prostate rather than the estrogenic endpoints discussed in this section. The developmental**
35 **effects of bisphenol A in the prostate are discussed in Section 3.2.]**

36
37 Inter-strain variability in rats has been evaluated as a source of variability in estrogenicity assays.
38 Inspection of Table 48 does not suggest large sensitivity differences between Sprague Dawley, Wistar,

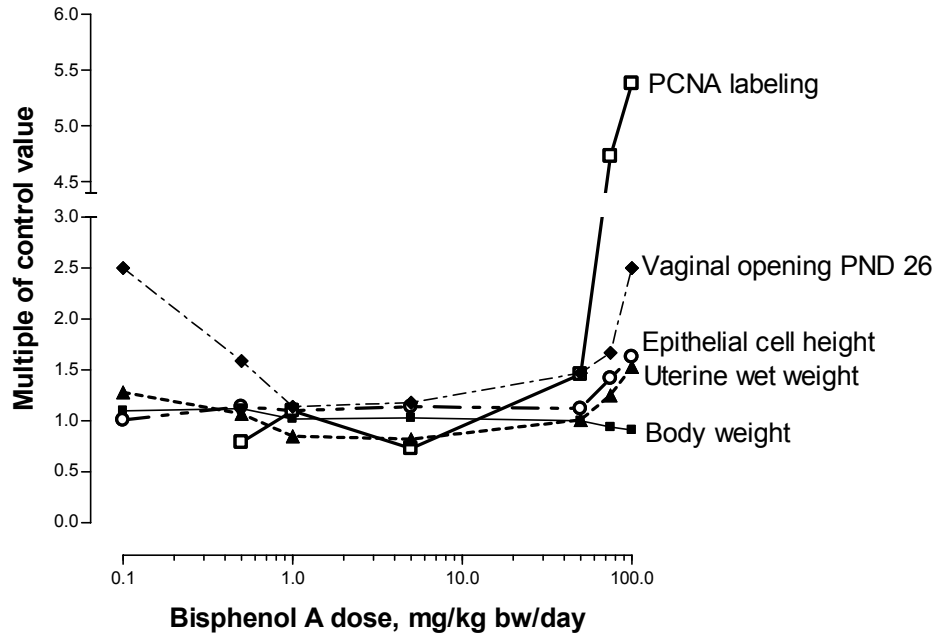
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1 and Long Evans rats. Greater sensitivity of F344 than Sprague Dawley rats has been shown with respect
2 to uterine weight and epithelial cell height (207), where 17 β -estradiol-adjusted potencies differed by 20–
3 37% between the strains. BrdU labeling of vaginal epithelium was 3 times greater in F344 than Sprague
4 Dawley rats in another study (208), and a third study (194) showed that both bisphenol A and 17 β -
5 estradiol increase serum prolactin in ovariectomized F344 but not ovariectomized Sprague Dawley rats.
6 Diel et al. (209) evaluated estrogenic response to bisphenol A in juvenile ovariectomized DA/Han,
7 Sprague Dawley, and Wistar rats. After 3 days of treatment with bisphenol A 200 mg/kg bw/day, there
8 were small statistically significant increases in uterine weight in DA/Han and Sprague Dawley rats but
9 not in Wistar rats. There were no alterations in uterine or vaginal epithelium or in uterine clusterin mRNA
10 expression in any of the strains after bisphenol A treatment.

11
12 Intra-laboratory variability has been noted for the bisphenol A uterotrophic assay in immature mice (210).
13 Of 8 studies performed over a 2-year period at sc bisphenol A dose levels up to 200 or 300 mg/kg bw/day,
14 4 showed a significant increase in uterine weight at 200 mg/kg bw/day. The other 4 studies, including the
15 2 studies that went to 300 mg/kg bw/day, showed no effect of bisphenol A treatment on uterine weight
16 despite the expect response to diethylstilbestrol. Study authors noted that reducing the permissible body
17 weight of the mice selected for study resulted in lower and less variable control uterine weights and
18 greater likelihood of bisphenol A effect (210, 211). **[The Expert Panel notes that these studies all used**
19 **high sc doses of bisphenol A.]**

20
21 It has been proposed that the rodent uterotrophic assay is relatively insensitive to the estrogenic effects of
22 bisphenol A (212). These authors treated immature CD-1 mice with bisphenol A and evaluated uterine
23 weight, relative area of uterine compartments, epithelial height, expression of lactoferrin and proliferating
24 cell nuclear antigen (PCNA), and induction of vaginal opening. Dose-response curves for the endpoints
25 that showed significant changes from control are illustrated in Figure 4. Uterine wet weight at the highest
26 dose was the only endpoint for which statistical significance was demonstrated. **[The Expert Panel notes**
27 **that the dose response curves for epithelial cell height and uterine wet weight appear parallel in the**
28 **upper-dose range.]** The study authors also noted that significant alterations in some endpoints were
29 observed at much lower doses (0.1 mg/kg bw/day for vaginal opening and 5 mg/kg bw/day for epithelial
30 cell height), giving rise to a U-shaped dose-response curve. **[The assertions of some investigators**
31 **notwithstanding, the Expert Panel notes that oral bisphenol A does not consistently produce**
32 **estrogenic responses and, when seen, estrogenic effects after oral treatment occur at high dose**
33 **levels.]**

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Figure 4. Dose-response curves for endpoints of estrogenic activity

On pair-wise testing, body weight was increased at 0.5 mg/kg bw/day and decreased at 100 mg/kg bw/day; vaginal opening was advanced at 0.1 and 100 mg/kg bw/day; epithelial cell height was increased at 5, 75, and 100 mg/kg bw/day; PCNA labeling was increased at 75 and 100 mg/kg bw/day; and uterine wet weight was increased at 100 mg/kg bw/day. Data from Markey et al. (212).

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1 **Table 49. In Vivo Estrogenicity Tests of Bisphenol A**

Model and exposure	Husbandry ^a	Endpoint	Molar potency/comparator ^b	Reference
Adult ovariectomized Sprague Dawley, gavage × 4 days	TD89222 diet, metal cage	<i>Rat uterus</i> Uterine wet weight	[3.9 × 10 ⁻³]/ethinyl estradiol	Dodge et al. (155)
Immature Sprague Dawley, bisphenol A given “orally” × 3 days; 17β-estradiol ip × 3 days	not indicated	Uterine weight	Not affected by bisphenol A at up to 150 mg/kg bw/day; 17β-estradiol was positive at 0.005 mg/day [-0.089 mg/kg bw/day]	Gould et al. (182)
Adult ovariectomized Crl:CD BR, gavage × 4 days	Purina 5002 diet, steel cage	Progesterone receptor Peroxidase activity Uterine weight Stromal cell proliferation <i>cfos</i> expression	[5.9 × 10 ⁻³]/17β-estradiol [7.6 × 10 ⁻³]/17β-estradiol [3.5 × 10 ⁻⁵]/17β-estradiol [4.1 × 10 ⁻⁵]/17β-estradiol [2.1 × 10 ⁻⁴]/17β-estradiol	Cook et al. (213)
Adult ovariectomized F344, ip × 1	Not indicated	Uterine wet weight: F344 Sprague Dawley	[8.2 × 10 ⁻³]/17β-estradiol [6.0 × 10 ⁻³]/17β-estradiol	Steinmetz et al. (207)
Adult ovariectomized F344 or Sprague Dawley, silastic implant × 3 days	Not indicated	Uterine cell height: F344 Sprague Dawley	[1.1 × 10 ⁻²]/17β-estradiol [9.2 × 10 ⁻³]/17β-estradiol	Steinmetz et al. (207)
Juvenile ovariectomized DA/Han, Wistar, or Sprague Dawley, gavage × 3 days	Not indicated	Uterine wet weight: DA/Han Wistar Sprague Dawley Uterine epithelium Vaginal epithelium Clusterin mRNA	[1.8 × 10 ⁻⁵]/ethinyl estradiol No response to 200 mg/kg/d [1.7 × 10 ⁻⁵]/ethinyl estradiol No response to 200 mg/kg/day No response to 200 mg/kg/day No response to 200 mg/kg/day	Diel et al. (209)
Immature Alpk:AP, sc × 3 days	RM3 diet, wire cage	Uterine wet weight	[2.6–2.7 × 10 ⁻⁵]/diethylstilbestrol	Ashby and Tinwell (204)
Immature Alpk:AP, gavage × 3 days	RM3 diet, wire cage	Uterine dry weight	[2.5–3.0 × 10 ⁻⁵]/diethylstilbestrol	
Immature Long Evans,	Purina 5001 diet	Uterine wet weight	[2.3–3.1 × 10 ⁻⁵]/diethylstilbestrol	
		Uterine dry weight	[2.7–3.6 × 10 ⁻⁵]/diethylstilbestrol	
		Uterine wet weight 6	[1.4 × 10 ⁻⁵]/17β-estradiol	Laws et al. (202)

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Model and exposure	Husbandry ^a	Endpoint	Molar potency/comparator ^b	Reference
gavage × 3 days		hours after dosing Uterine wet weight 24 hours after dosing	No effect at bisphenol A at ≤ 400 mg/kg bw/day	
Adult ovariectomized Long Evans	Purina 5001 diet	Uterine wet weight	No effect of bisphenol A at ≤ 100 mg/kg bw/day	Laws et al. (202)
Juvenile ovariectomized DA/Han, gavage × 3 days	Ssniff R-10 diet	Uterine wet weight relative to bw <i>Expression of:</i> Androgen receptor <i>ER</i> Progesterone receptor	[1.2 × 10⁻⁵] /ethinyl estradiol [3.9 × 10⁻⁴] /ethinyl estradiol [1.9 × 10⁻⁴] /ethinyl estradiol bisphenol A and ethinyl estradiol produced opposite effects	Diel et al. (214)
		Complement C3 Clusterine Glyceraldehyde phosphate dehydrogenase	[2.2 × 10⁻⁵] /ethinyl estradiol No bisphenol A effect at 200 mg/kg bw/day; ethinyl estradiol showed an effect at 0.1 mg/kg bw/day.	
Adult ovariectomized Alpk:ApfSD, sc × 3 days	Not indicated	Uterine wet weight Uterine dry weight	[1.7 × 10⁻⁴] /17β-estradiol [1.8 × 10⁻⁴] /17β-estradiol	Ashby et al. (215)
Immature Crj:CD (SD), sc × 3 days	MF diet, steel cage	Wet and blotted uterine weight	Effect noted at ≥8 mg/kg bw/day bisphenol A/no comparator	Yamasaki et al. (99)
Immature Crj:CD (SD), gavage × 3 days	MF diet, steel cage	Wet and blotted uterine weight	Effect noted at ≥160 mg/kg bw/day bisphenol A/no comparator	
Adult ovariectomized Wistar, sc × 7 days	Not indicated	Blotted uterine weight	Increased relative weight compared to placebo at ≥11 mg/kg bw/day; uterus reached 83% of weight of sham- ovariectomized control at bisphenol A dose of 250 mg/kg bw/day.	Goloubkova et al. (216)
Adult ovariectomized Sprague Dawley, exposed in drinking water × 3 days	Glass water bottles, plastic cage (negative E-Screen of ethanol cage washes)	Uterine wet weight	No effect of bisphenol A at up to 16.9 mg/kg bw/day; estrone positive at 0.12 mg/kg bw/day	Rubin et al. (217)
Adult ovariectomized	PMI Certified Rodent Diet,	Uterine wet weight	[1.7 × 10⁻⁶] /17β-estradiol	Kim et al. (150)

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Model and exposure	Husbandry ^a	Endpoint	Molar potency/comparator ^b	Reference
Sprague Dawley, sc × 3 days	polycarbonate cage, elm bedding	Uterine dry weight	[2.3 × 10 ⁻⁶]/17β-estradiol	
Immature Alpk:ApfSD, sc × 3 days	RM1 diet	Uterine wet weight Uterine dry weight	[2.9 × 10 ⁻⁴]/17β-estradiol No effect of bisphenol A at 800 mg/kg bw/day; 17β-estradiol positive at 0.4 mg/kg bw/day	Matthews et al. (138)
Immature Alpk:ApfSD, gavage × 3 days	RM1 diet	Uterine wet weight Uterine dry weight	[2.3–5.5 × 10 ⁻⁴]/17β-estradiol [2.4–7.1 × 10 ⁻⁴]/17β-estradiol	
Immature Sprague Dawley, sc × 3 days	Soy-free diet, polycarbonate cage	Uterine wet weight	No effect of bisphenol A at ≤ 1000 mg/kg bw/day; 17β-estradiol was positive at 0.04 mg/kg bw/day	An et al. (218)
		Calbindin D _{9k} expression	[8.4 × 10 ⁻⁶]/17β-estradiol	
		ERα expression	[3.4 × 10 ⁻⁵]/17β-estradiol	
Immature Crj:CD (SD), sc × 3 days	MF diet, steel cage	Uterine wet weight	[5.1 × 10 ⁻⁵]/ethinyl estradiol	Yamasaki et al. (178)
Immature Sprague Dawley, sc × 3 days	Soy-free diet, polycarbonate cage, corncob bedding	Blotted uterine weight Epithelial cell height	[8 × 10 ⁻⁷]/ethinyl estradiol [1.2 × 10 ⁻⁶]/ethinyl estradiol	Wade et al. (219)
Pubertal Sprague Dawley, gavage PND 22–42/43	Purina 5002 diet, polycarbonate cage, chip bedding	Blotted uterine weight	Absolute organ weight decreased with increase dose (400 and 600 mg/kg bw/day); no effect on relative organ weight	George et al. (203)
		Vaginal opening	No effect at 400 and 600 mg/kg bw/day	
Pregnant Sprague Dawley, sc bisphenol A on GD 17–19 (17β-estradiol sc × 1)	Soy-free diet, polycarbonate cage	Maternal uterine weight	[1.8 × 10 ⁻⁵]/17β-estradiol	Hong et al. (220)
		Maternal uterine calbindin D _{9k} protein	[1.7 × 10 ⁻⁵]/17β-estradiol	
Lactating Sprague Dawling, sc bisphenol A × 5 days (17β-estradiol sc × 1)	Soy-free diet	Maternal uterine calbindin D _{9k} mRNA	[2.2 × 10 ⁻⁵]/17β-estradiol	Hong et al. (221)
		calbindin D _{9k} protein	[6.9 × 10 ⁻⁵]/17β-estradiol	
Immature and adult	AO4C diet, wire cage	Uterine wet and dry	No effect in either model of	Strohecker et al. (222)

2.0 General Toxicology and Biological Effects

Model and exposure	Husbandry ^a	Endpoint	Molar potency/comparator ^b	Reference
ovariectomized Wistar, gavage × 4 days		weight	bisphenol A at ≤ 200 mg/kg bw/day/17β-estradiol positive at 0.025–0.035 mg/kg bw/day	
Immature Sprague Dawley, sc × 3 days	Soy-free feed, polycarbonate cage	Calbindin D _{9k} protein	[5.1 × 10 ⁻⁵]/17β-estradiol	An et al. (223)
Immature Sprague Dawley, sc × 3 days	Shinchon diet	Uterine wet weight Uterine wet weight relative to bw Glutathione peroxidase activity	[1.5 × 10 ⁻⁶]/17β-estradiol [1.3 × 10 ⁻⁶]/17β-estradiol [4.2 × 10 ⁻³]/17β-estradiol	Kim et al. (184)
Immature AlpK:ApfSD, gavage × 3 days	RM1 diet, polycarbonate cage	Blotted uterine weight <i>Expression of:</i> Progesterone receptor A Progesterone receptor B Complement C3 Lipocalcin	[2.5 × 10 ⁻⁴]/17β-estradiol [3.8 × 10 ⁻⁴]/17β-estradiol [4.2 × 10 ⁻⁴]/17β-estradiol [1.8 × 10 ⁻⁴]/17β-estradiol [2.3 × 10 ⁻⁴]/17β-estradiol	Ashby and Odum (224)
Immature AP, sc × 3 days	RM1 diet, polypropylene cages, sawdust and shredded paper bedding	Uterine wet weight Uterine dry weight	[1.0 × 10 ⁻⁶]/ethinyl estradiol [1.2 × 10 ⁻⁶]/ethinyl estradiol	Tinwell and Ashby (225)
Adult ovariectomized Sprague Dawley, diet × 3 months	Phytoestrogen-free diet	Uterine weight, endometrial thickness, <i>ERα</i> , <i>ERβ</i> expression Complement C3 expression	No bisphenol A effect at 0.37 mg/kg bw/day; estradiol benzoate positive control Bisphenol A and estradiol benzoate produced opposite effects	Seidlová-Wuttke et al. (136)
Immature Sprague Dawley, sc × 3 days	PMI Certified Rodent Diet	Uterine wet weight Uterine dry weight	[4.5 × 10 ⁻⁷]/ethinyl estradiol [4.9 × 10 ⁻⁷]/ethinyl estradiol	Kim et al. (226)
Adult ovariectomized Crj:CD (SD), sc × 3 days	Estrogen-free NIH-07PLD diet, aluminum cage, paper bedding	Uterine wet weight, relative to bw Blotted uterine weight, relative to bw	[2.1 × 10 ⁻⁵]/17β-estradiol [1.7 × 10 ⁻⁶]/17β-estradiol	Koda et al. (227)

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Model and exposure	Husbandry ^a	Endpoint	Molar potency/comparator ^b	Reference
Adult Holzman, progesterone-treated to delay implantation, given test agent sc on GD 7	Unspecified Purina rodent chow, plastic cage, pine shavings	Implantation	[4-34 × 10 ⁻⁶]/estrone	Cummings et al. (228)
<i>Rat vagina</i>				
Adult ovariectomized F344, ip × 1	Not indicated	BrdU labeling	Increased at bisphenol A dose of 37.5 but not 18.5 mg/kg bw/no comparator	Steinmetz et al. (207)
Adult ovariectomized Long Evans, bisphenol A by gavage × 11 days; 17β-estradiol by sc	Purina 5001 diet	<i>cfos</i> expression Vaginal cytology	[1.3 × 10 ⁻⁴]/17β-estradiol No effect at bisphenol A dose of 100 mg/kg bw/day; 17β-estradiol 0.005 mg/kg bw/day resulted in persistent estrus.	Laws et al. (202)
Long Evans treated PND 21–35 by gavage	Purina 5001 diet	Vaginal opening	No effect at bisphenol A dose ≤ 400 mg/kg bw/day; ethinyl estradiol was active at 0.01 mg/kg bw/day.	Laws et al. (202)
Adult ovariectomized F344 and Sprague Dawley, ip × 1	Not indicated	BrdU labeling	F344: [4.5 × 10 ⁻⁶]/17β-estradiol Sprague Dawley: [1.4 × 10 ⁻⁶]/17β-estradiol	Long et al. (208)
Immature Wistar, gavage × 4 days	AO4C diet, wire cage	Vaginal cornification	[3.8 × 10 ⁻⁴]/17β-estradiol	
Adult ovariectomized Wistar, gavage × 4 days	AO4C diet, wire cage	Vaginal cornification	No effect at bisphenol A dose of 100 mg/kg bw/day; 17β-estradiol was positive at 0.1 mg/kg bw/day	Strohecker et al. (222)
Immature Sprague Dawley, sc × 3 days	PMI Certified Rodent Diet	Vaginal weight	[5.3 × 10 ⁻⁷]/ethinyl estradiol	Kim et al. (226)
<i>Other rat organs</i>				
Ovariectomized Sprague Dawley, daily gavage for 5 weeks	TD89222 diet, metal cage	Prevention of bone mineral density decline	No effect at bisphenol A dose up to 10 mg/kg bw/day; no standard estrogen comparator.	Dodge et al. (155)
Adult ovariectomized	Phytoestrogen-free diet	Prevention of bone	No effect at bisphenol A dose ≤	Seidlová-Wuttke et al. (136)

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Model and exposure	Husbandry ^a	Endpoint	Molar potency/comparator ^b	Reference
Sprague Dawley, treated in feed		mineral density decline	370 µg/kg bw/day; estradiol benzoate was effective at 1.18 mg/kg bw/day.	
Adult ovariectomized Sprague Dawley and F344, by sc implant × 3 days	Not indicated	Serum prolactin	F344: $[1.7 \times 10^{-2}]$ /17β-estradiol Sprague Dawley: no effect of bisphenol A at 40–45 µg/day or 17β-estradiol at 1.2–1.5 µg/day.	Steinmetz et al. (194)
Adult ovariectomized Wistar, sc × 7 days	Not indicated	Pituitary weight	Increased compared to vehicle control at 128 but not 78 mg/kg bw/day	Goloubkova et al. (216)
		Serum prolactin	Increased compared to vehicle control at 128 mg/kg bw/day	
		<i>Mouse uterus</i>		
Immature CFLP, sc × 3 days	Not indicated	Relative uterine weight	No response at up to 0.5 mg [~50 mg/kg bw/day]	Coldham et al. (162)
Adult ovariectomized CD-1, sc × 1	Not indicated	<i>IGF1</i> expression	$[8.4 \times 10^{-4}]$ /17β-estradiol	Klotz et al. (229)
Juvenile-adult ovariectomized B6C3F ₁ , sc × 4 days	Purina 5001, polypropylene cage, chip bedding	Uterine wet weight	$[2.3 \times 10^{-5}]$ /17β-estradiol	Papconstantinou et al. (230)
		Endothelial proliferation	$[6.9 \times 10^{-6}]$ /17β-estradiol	
Juvenile-adult ovariectomized B6C3F ₁ , sc × 4 days	Purina 5001, polypropylene cage, cellulose fiber bedding	Induction of grp94	$[2.4 \times 10^{-5}]$ /17β-estradiol	Papconstantinou et al. (231)
		Induction of hsp72	$[3.5 \times 10^{-6}]$ /17β-estradiol	
		Induction of hsp90	$[5.3 \times 10^{-6}]$ /17β-estradiol	
Juvenile-adult ovariectomized B6C3F ₁ , sc × 4 days	Purina 5001, polypropylene cage, cellulose fiber bedding	Uterine weight	$[5.3 \times 10^{-6}]$ /17β-estradiol	Papconstantinou et al. (232)
		Induction of hsp90α	$[1.2 \times 10^{-5}]$ /17β-estradiol	
		Induction of grp24	$[8.4 \times 10^{-6}]$ /17β-estradiol	
Juvenile-adult ovariectomized B6C3F ₁ , sc × 1	Purina 5001, polypropylene cage, cellulose fiber bedding	Blotted uterine weight, 6 hours after dose	$[8.4 \times 10^{-6}]$ /17β-estradiol	Papconstantinou et al. (233)
		Blotted uterine weight, 12 hours after dose	$[4.2 \times 10^{-6}]$ /17β-estradiol	
Adult ovariectomized transgenic ER-reporter, sc × 1	Purina 5001, polystyrene cage	Uterine wet weight	$[2.9 \times 10^{-5}]$ /diethylstilbestrol	Nagel et al. (234)
		ER activation	$[1.0 \times 10^{-4}]$ /diethylstilbestrol	
Immature AP, sc × 3 days	RM1 diet, plastic cage,	Blotted uterine weight	$[2.3 \times 10^{-5}]$ /diethylstilbestrol in 4	Tinwell and Joiner (210)

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Model and exposure	Husbandry ^a	Endpoint	Molar potency/comparator ^b	Reference
	sawdust and shredded paper bedding		of 8 trials; other trials showed no effect at bisphenol doses up to 300 mg/kg bw/day.	
Immature AP, gavage × 3 days	RM1 diet, plastic cage, sawdust and shredded paper bedding	Blotted uterine weight	No effect at bisphenol A doses up to 300 mg/kg bw/day; diethylstilbestrol produced response at 10 µg/kg bw/day.	Tinwell and Joiner (210)
Immature CD-1, sc × 3 days	RM1 diet	Lactoferrin expression	No effect at bisphenol A doses up to 1000 mg/kg bw/day; diethylstilbestrol showed effect at 0.1 µg/kg bw/day.	Mehmood et al. (235)
		Uterine weight, BrdU incorporation, peroxidase production	No effect at bisphenol A doses up to 100 mg/kg bw/day; diethylstilbestrol showed effect at 1–5 µg/kg bw/day.	
Immature CD-1, sc minipump × 3 days	RMH 3000 diet, cage, and bedding estrogen-negative by E-Screen	Uterine wet weight	$[1.6 \times 10^{-5}]$ /17β-estradiol	Markey et al. (212)
		Epithelial cell height	$[3.8 \times 10^{-5}]$ /17β-estradiol	
		Lactoferrin expression	$[3.9 \times 10^{-5}]$ /17β-estradiol	
Ovariectomized adult B6C3F ₁ , ip × 3 days	Not indicated	Relative uterine to body weight	$[3.6–74 \times 10^{-5}]$ /17β-estradiol	Kitamura et al. (188)
Ovariectomized adult Swiss, sc × 1	Economy Rodent Maintenance diet	Increased uterine vascular permeability	$\sim 1 \times 10^{-4}$ /17β-estradiol	Milligan et al. (236)
<i>Other mouse organs</i>				
Juvenile-adult aromatase knock-out, diet × 4 months	NMF diet	Uterine and ovarian histology, bone mineral density	Dietary bisphenol A (0.1%) exerted estrogenic effects. Mean ± SD serum bisphenol A 84.3 ± 8.7 µg/L. No comparator estrogen was used for these endpoints.	Toda et al. (237)
<i>Fish</i>				
Immature rainbow trout, injected		Plasma vitellogenin	$[3 \times 10^{-4}]$ /17β-estradiol	Christiansen et al. (238)
Juvenile rainbow trout,		Plasma vitellogenin	$[5.6 \times 10^{-3}]$ /17β-estradiol	Andersen et al. (139)

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Model and exposure	Husbandry ^a	Endpoint	Molar potency/comparator ^b	Reference
injected Juvenile rainbow trout, exposed in water		Plasma vitellogenin	[~8.4 × 10 ⁻⁵]/17β-estradiol	Lindholst et al. (239)
Male medaka, exposed in feed		Plasma vitellogenin	[1.4 × 10 ⁻⁴]/ethinyl estradiol	Chikae et al. (240)
Male medaka, exposed in water		Hepatic vitellogenin and ERα mRNA	[8.4 × 10 ⁻⁶]/17β-estradiol	Yamaguchi et al. (241)
Male killfish, injected		Plasma vitellogenin	[2.7 × 10 ⁻⁴]/17β-estradiol	Pait et al. (242)
Male zebrafish, juvenile rainbow trout, exposed in water		Plasma vitellogenin	[~0.2]/ethinyl estradiol	Van den Belt et al.(243)
<i>Invertebrates</i>				
Mudsnail, exposed in water		New embryo production	[1.5 × 10 ⁻⁴]/ethinyl estradiol	Jobling et al. (244)
Ramshorn snail, exposed in water		Egg production	Increased (EC ₁₀ 13.9 ng/L); blocked by faslodex and tamoxifen. No comparison to reference estrogen	Oehlmann et al. (245)

^aHusbandry information for rodent studies includes caging and bedding materials and diet when indicated by the authors.

^bEstimates include comparison of administered dose, magnitude of effect, and molecular weight.

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1 Nagel et al. (234) developed a transgenic mouse with a thymidine kinase-*lacZ* reporter linked to 3
2 copies of the vitellogenin estrogen response element. This model showed an increase in ER
3 activity after a single sc bisphenol A dose of 25 µg/kg bw ($P = 0.052$), with further increases in
4 activity after 0.8 and 25 mg/kg bw. Uterine weight was only increased at the 25 mg/kg bw dose
5 level. Normalized to the diethylstilbestrol response, uterine weight response to bisphenol A 25
6 mg/ kg bw was less than one-third the response in ER activity [estimated from a graph].
7

8 Gene expression profiles have been performed to compare the presumably ER-mediated response
9 to bisphenol A with the response to reference ER agonists. Naciff et al. (246) evaluated
10 expression in the uteri and ovaries of Sprague Dawley fetuses after sc dosing of dams on GD 11–
11 20 with ethinyl estradiol 0, 0.5, 1 or 10 µg/kg bw/day or bisphenol A 0, 5, 50, or 400 mg/kg
12 bw/day. The high dose of both compounds induced nipples and areolae in male and female
13 fetuses. There were 366 genes in which expression was altered by ethinyl estradiol and 397 genes
14 in which expression was altered by bisphenol A. Expression of 66 genes was changed in the same
15 direction with high doses of ethinyl estradiol, bisphenol A, and genistein (which was also tested
16 in this model). Of the 40 genes with at least a 1.8-fold change in expression, 17 responded
17 similarly to ethinyl estradiol and bisphenol A. The authors identified 50 mg/kg bw/day as the
18 lowest dose level at which estrogen-like gene expression activity could be identified, which is
19 lower than the 400–800 mg/kg bw/day dose range at which uterotrophic activity is typically
20 reported in rats (204).
21

22 Terasaka et al. (247) used expression of 120 estrogen-responsive genes (based on previous work)
23 in MCF-7 cells to compare the profiles of bisphenol A and 17β-estradiol. Response was highly
24 correlated ($R = 0.92$) between the 2 compounds. Another gene array study (248) used MCF-7
25 cells that had lost ER and were re-engineered to express ERα. Among 40 estrogen-responsive
26 genes, 12 responded to both bisphenol A and 17β-estradiol, 9 responded only to bisphenol A, and
27 19 responded only to 17β-estradiol. In the ER-deficient MCF-7 cell line from which these cells
28 had been engineered, 1 gene responded to both bisphenol A and 17β-estradiol and 14 responded
29 to bisphenol A alone, suggesting ER-independent activity. The same group reported the response
30 of an additional 31 genes, associated with growth and development, from the same chip (172). In
31 the ERα-containing cells, 5 of these genes showed regulation with both 17β-estradiol and
32 bisphenol A, 13 were regulated only by bisphenol A, and 13 were regulated only by 17β-
33 estradiol.
34

35 Differences in the estrogenic activity of bisphenol A and reference estrogens may be due to
36 differences in recruiting by the liganded receptor of co-regulatory proteins. Singleton et al. (172)
37 used a co-regulator-independent yeast reporter system to evaluate the estrogenicity of bisphenol
38 A and 17β-estradiol. Bisphenol A activity was more than 3 orders of magnitude less than 17β-
39 estradiol in the yeast system, compared to about a 2-order-of-magnitude difference in an MCF-7
40 cell assay, leading the authors to postulate that mammalian co-activators may be involved in
41 enhancing bisphenol A activity. In a comparison of ER binding and co-activator recruitment,
42 Routledge et al. (135) showed bisphenol A to bind the receptor more avidly than the liganded
43 receptor recruited 2 co-activator proteins, normalized to 17β-estradiol (Table 50).
44

1 **Table 50. Bisphenol A Receptor Binding and Recruitment of Co-Activator Proteins**

Assay	Activity relative to 17 β -estradiol	
	ER α	ER β
Receptor binding	7.3×10^{-4}	7.5×10^{-3}
TIF2 recruitment	$< 1 \times 10^{-6}$	5×10^{-4}
SRC-1a recruitment	3×10^{-4}	2×10^{-4}

From Routledge et al. (135).

2
3 The classical ERs are cytosolic receptors that, when bound, translocate to the nucleus where they
4 produce their activity through alterations in genomic transcription. In contrast, a membrane-
5 bound ER has been described in murine pancreatic islet cells (249-252). This membrane-bound
6 receptor regulates calcium channels and modulates insulin and glucagon release. Bisphenol A has
7 been shown to activate this receptor in vitro at a concentration of 1 nM, which is similar to the
8 active concentration of diethylstilbestrol (249, 251). Treatment of mice with bisphenol A or 17 β -
9 estradiol sc at 10 μ g/kg bw acutely or daily for 4 days resulted in decreased plasma glucose and
10 increased insulin (253). By contrast, Adachi et al. (254) reported that exposure of rat pancreatic
11 islets to 0.1–1 μ g/L [0.4–4.4 nM] bisphenol A did not alter insulin secretion over a 1-hour period.
12 Exposure of islets to bisphenol A 10 μ g/L [44 nM] for 24 hours increased insulin release. This
13 response was prevented by actinomycin D and by ICI 182,780, supporting the conclusion that
14 bisphenol A insulin release occurs through interaction with the cytoplasmic ER rather than the
15 membrane-bound receptor.

16
17 In addition to the studies reviewed for this section, there are studies in which the putative
18 estrogenicity of environmental samples or synthetic products were evaluated using one or another
19 assay. For example, Olea et al. (48) evaluated resin-based dental composites in an MCF-7 culture
20 system. The response of the system was attributed to the bisphenol and its methacrylate detected
21 in the composites, but bisphenol A was not specifically tested. These papers were not reviewed
22 for this section.

23 24 2.2.3 Androgen activity

25 Transfected cell-based assays have not identified bisphenol A as having androgenic activity (165,
26 183, 188, 255). However, bisphenol A is mitogenic in cultured human prostate carcinoma cells at
27 a concentration of 1 nM (256). Based on stimulated cell growth in this system, the potency of
28 bisphenol A is about 5% that of dihydrotestosterone [estimated from a graph]. This bisphenol A
29 activity was shown to be mediated by interaction with a mutant tumor-derived androgen receptor
30 called AR-T877A. [Interaction with the mutant androgen receptor has been proposed as
31 representing a risk for men with prostate cancer but does not appear to have reproductive
32 implications and will not be further considered in this report.]

33
34 Anti-androgenic activity has been demonstrated using cells transfected with androgen receptor
35 reporting systems (Table 51). The anti-androgenic activity of bisphenol A is expressed as the
36 concentration needed to halve the androgen reporter response to a reference androgen. Studies in
37 transfected cells have shown that bisphenol A interferes with the binding of dihydrotestosterone
38 to the androgen receptor, interferes with translocation of the liganded receptor to the nucleus, and
39 prevents transactivation at the androgen-response element (257).

40

1 **Table 51. Anti-androgenicity Studies of Bisphenol A in Cells Transfected with Androgen**
 2 **Receptor Reporter**

Cell type	Reference androgen concentration (nM)	Bisphenol A median inhibitory concentration (IC ₅₀), μ M [mg/L]	Reference
Human prostate adenocarcinoma	R1881 0.1	7 [1.6]	Paris et al. (133)
Chinese hamster ovary	R1881 0.1	19.6 [4.5]	Roy et al. (258)
Yeast	Testosterone 10	1.8 [0.4]	Lee et al. (257)
Yeast	Dihydrotestosterone 1.25	2 ^a [0.5]	Sohoni and Sumpter (165)
Monkey kidney	Dihydrotestosterone 1	0.746 [0.2]	Xu et al. (255)
Mouse fibroblast	Dihydrotestosterone 0.01	4.3 [1.0]	Kitamura et al. (188)
Human hepatoma	Dihydrotestosterone 100	No anti-androgenic activity	Gaido et al. (183)

^aEstimated from a graph.

3
 4 Kim et al. (259) conducted a Hershberger assay to determine the effects of bisphenol A exposure
 5 on reproductive organs of rats. Sprague Dawley rats were fed PMI Certified Rodent LabDiet and
 6 housed in polycarbonate cages. No information was provided about bedding materials. One
 7 experiment was conducted to determine the optimum dose and age for observing testosterone
 8 exposure effects. In a second experiment, 10 rats/group rats were castrated at 5 weeks of age and
 9 7 days later gavaged with bisphenol A (99% purity) at doses of 0 (ethanol/corn oil vehicle) 10,
 10 100, or 1000 mg/kg bw/day for 7 days. A second group of castrated 6-week-old males rats was
 11 gavaged with bisphenol A at 0, 50, 100, 250, or 500 mg/kg bw/day for 7 days. In a third
 12 experiment, 10 castrated 6-week-old rats/group were treated with 0.4 mg/kg bw/day testosterone
 13 by sc injection in addition to gavaged bisphenol A at 50, 100, 250, or 500 mg/kg bw/day or
 14 flutamide at 1, 5, 10, or 25 mg/kg bw/day for 7 days. A positive control group was given 0.4
 15 mg/kg bw/day testosterone for 7 days. **[There is some confusion in the paper regarding ages at**
 16 **castration and start of treatment. For the first group of bisphenol A-treated rats, it is**
 17 **reported that rats were castrated at 5 weeks of age and treated at 6 weeks of age. For the**
 18 **other groups of bisphenol A-treated rats, the Methods section reported that treatment**
 19 **began at 6 weeks of age, but tables in the Results section indicated that rats were castrated**
 20 **at 6 weeks of age.]** During the study, clinical signs were observed and body weights were
 21 measured. Blood was collected and rats were killed ~24 hours after administration of the last
 22 dose. Accessory reproductive organs were removed and weighed. Serum luteinizing hormone
 23 (LH) and testosterone levels were measured by radioimmunoassay (RIA). Statistical analyses
 24 included Bartlett test, analysis of covariance (ANCOVA), Dunnet test, and Bonferroni test.
 25 Exposure to bisphenol A did not affect weights of the ventral prostate, seminal vesicles, glans
 26 penis, or levator ani plus bulbocavernosus muscle; or serum levels of LH or testosterone.
 27 Testosterone increased the weights of accessory reproductive organs. Flutamide increased serum
 28 LH levels and inhibited testosterone-induced increases in accessory reproductive organ weights.
 29 Study authors concluded that bisphenol A did not exhibit estrogenic or antiestrogenic effects in
 30 rats.

31
 32 Yamasaki et al. (260) conducted a Hershberger assay in rats exposed to bisphenol A or 1 of 29
 33 other chemicals. In this study, which was conducted according to GLP, animals were housed in
 34 stainless steel wire-mesh cages. Assuming these males were fed the same diets as rats used in an
 35 uterotrophic assay also described in this study, they received MF Oriental Yeast feed. Rats were
 36 randomly assigned to treatment groups. Beginning at 56 days of age and continuing for 10 days, 6
 37 castrated male Brl Han: WIST Jcl (GALAS) rats/group were administered bisphenol A by

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1 stomach tube at doses of 0 (olive oil vehicle), 50, 200, or 600 mg/kg bw/day. An additional group
 2 of rats was administered the same vehicle and doses of bisphenol A in addition to 0.2 mg/kg
 3 bw/day testosterone propionate by sc injection. Dose selection was based on results of
 4 preliminary studies. A positive control group was given 10 mg/kg bw/day flutamide in addition to
 5 0.2 mg/kg bw/day testosterone propionate. Rats were killed 24 hours after receiving the final
 6 dose. Ventral prostate with fluid, seminal vesicles with fluid, bulbocavernosus/levator ani muscle,
 7 glans penis, and Cowper gland were collected and weighed. Data were analyzed by Student *t*-test.
 8 Bisphenol A did not affect body weight and there were no clinical signs of toxicity. The only
 9 statistically significant effect on relative organ weight was a **[24%]** increase in glans penis weight
 10 in rats given 600 mg/kg bw/day bisphenol A without coadministration of testosterone. In contrast,
 11 rats treated with flutamide plus testosterone propionate experienced increases in weights of
 12 ventral prostate, seminal vesicle, bulbocavernosus/levator ani muscle, glans penis, and cowper
 13 gland. **[Absolute organ weights were not reported. It is assumed but was not stated that**
 14 **relative weights were based on body weight.]** Study authors noted that because glans penis
 15 weights were variable in control rats and weights of other accessory reproductive organs were not
 16 affected, bisphenol A could not be clearly determined to have androgen agonistic properties.

17
 18 Nishino et al. (261) performed a Hershberger assay in Wistar rats. At 2 weeks of age, rats were
 19 given ssniffR 10 diet and housed in Makrolon cages with ssniff bedding. Seven days after
 20 orchietomy, rats were placed in groups of 13 **[randomization not discussed]** and treated orally
 21 **[gavage assumed]** with bisphenol A **[purity not indicated]** in propylene glycol at 0, 3, 50, 200,
 22 or 500 mg/kg bw/day for 7 days or sc with testosterone propionate 1 mg/kg bw. Another group
 23 was given oral bisphenol A 500 mg/kg bw/day and flutamide 3 mg/kg bw/day. Rats were killed
 24 by decapitation after treatment. Seminal vesicles and prostates were weighed and fixed in 4%
 25 neutral buffered paraformaldehyde. Immunohistochemical evaluation of androgen receptor,
 26 PCNA, and MIB-5 was performed. Epithelial cell height and duct luminal area were determined
 27 morphometrically. Statistical analysis used a 2-sided *t*-test. Results are summarized in Table 52.
 28 Prostate and seminal vesicle weight were increased by testosterone propionate but not by
 29 bisphenol A. **[The study authors interpreted the changes in relative organ weights as being**
 30 **due to treatment effects on body weight.]** Study authors concluded that bisphenol A did not
 31 exert androgenic effects; they characterized the effects of 3 and 50 mg/kg bw/day bisphenol A on
 32 epithelial cell height and luminal area of the prostate and seminal vesicles as “androgen-like”
 33 effects for which the mechanism was unclear. **[Although some unusual endpoints appear to**
 34 **have been affected, they are not validated measures of hormone action. Bisphenol A had no**
 35 **androgenic or anti-androgenic effects on organ weights.]**

36
 37 **Table 52. Effects of Bisphenol A on Prostate and Seminal Vesicles in Castrated Rats**

Endpoint	Bisphenol A dose, mg/kg bw/day				
	3	50	200	500	500 + flutamide
Body weight ^a	↔	↔	↓25%	↓30%	↓32%
Absolute organ weight					
Prostate	↔	↔	↔	↔	↔
Seminal vesicles	↔	↔	↔	↔	↔
Relative organ weight ^b					
Prostate	↔	↔	↑	↑	↑
Seminal vesicles	↔	↔	↑	↑	↑
Androgen receptor staining					
Prostate					
Polyclonal antibody	↑1.5-fold	↑1.3- fold	↑1.1-fold	↓9%	↓23%
Monoclonal antibody	↑1.8-fold	↑1.8-fold	↔	↓12%	↓36%

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Endpoint	Bisphenol A dose, mg/kg bw/day				
	3	50	200	500	500 + flutamide
Seminal vesicles					
Polyclonal antibody	↔	↔	↔	↔	↔
PCNA staining (prostate)	↔	↔	↔	↔	↔
MIB-5 staining (prostate)	↔	↔	↔	↔	↔
Epithelial cell height					
Prostate	↑1.3-fold	↑1.3-fold	↔	↓18	↓27%
Seminal vesicles	↑1.2-fold	↑1.2-fold	↔	↔	↔
Luminal area					
Prostate	↑3.3-fold	↑3.6-fold	↑2.1-fold	↔	↔
Seminal vesicles	↑5-fold	↑3.9-fold	↔	↔	↔

↑,↓,↔ Statistically significant increase, decrease, or no change compared to vehicle-treated, orchietomized control.

^aEstimated from a graph.

^bData were not shown; the direction of the change or the lack of change from control was indicated in the text. From Nishino et al. (261).

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2

2.3 Genetic Toxicity

3

Assessment of mutagenicity associated with bisphenol A was based primarily on reviews by the European Union (2) and Haighton et al. (24). CERHR summarized a limited number of studies that were not included in reviews. Results of in vitro genetic toxicity testing are summarized in Table 53, and results of in vivo genetic toxicity tests are summarized in Table 54.

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The European Union (2) noted that bisphenol A demonstrated aneugenic potential and micronuclei formation in in vitro tests without metabolic activation. However, there was no evidence of micronuclei formation in an in vivo mouse study. Other studies demonstrated disruption of microtubule formation and the presence of DNA adducts. In the studies reviewed by the European Union, there was no evidence of gene mutations or structural chromosomal aberrations in in vitro tests and negative results were obtained in a dominant lethal test in rats; however, the European Union noted several limitations for those studies. Based on their review of genotoxicity data and the lack of significant tumors reported in animal studies, the European Union (2) concluded that bisphenol A does not appear to have significant mutagenicity potential in vivo. Because aneugenic potential was apparently observed only in in vitro tests, it was judged to be of no concern. The relevance of DNA adduct formation was unclear, but based on negative findings for gene mutation and clastogenicity in cultured mammalian cells, DNA adduct formation was thought unlikely to be of concern for humans.

21

22

Haighton et al. (24) concluded that results of standardized and validated genetic toxicity tests demonstrated the lack of mutagenic and genotoxic activity of bisphenol A in vivo. Studies demonstrating disrupted microtubule formation or DNA adduct formation were noted, but because the studies used high doses, they were judged to be of limited relevance. The lack of activity in an in vivo micronucleus assay in mice was said to confirm negative results observed in in vivo tests. Lastly it was concluded that bisphenol A had no structural features that suggested mutagenic activity.

29

30

Subsequent to the release of the European Union (2) and Haighton et al. (24) reviews Hunt et al. (262), published a study examining meiotic aneuploidy potential of bisphenol A in female mice. In 1998, a large increase in background rate of congression failure (from 1–2 to 40%) and in aneuploidy (from 0.7 to 5.8%) was observed in the study authors' laboratory. The increase was

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32

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found to coincide with damage to polycarbonate caging material. Removal of the most damaged cages and change to polysulfone cages resulted in decreased background rates of congression failure. Intentionally damaging polycarbonate cages and water bottles resulted in increased rates of congression failure. As noted in Table 54, congression failure rates were increased in juvenile female mice orally exposed to ≥ 20 $\mu\text{g}/\text{kg}$ bw/day bisphenol A for 6–8 days or 20 $\mu\text{g}/\text{kg}$ bw/day for 7 days. Study authors concluded that bisphenol A was a potential meiotic aneugen.

Noting the Hunt et al. (262) data, Attia et al. (263) reported on a series of studies examining possible aneugenic activity of bisphenol A. The study information is currently available only as an abstract. One in vitro study (summarized in Table 53) examined aneuploidy in mouse oocytes and reported no increase in hyperhaploidy rate, but there was an apparently non-dose-related increase in diploid metaphase II oocytes at 200 $\mu\text{g}/\text{L}$. In vivo exposure studies showed no increases in aneuploidy in mouse spermatocytes or oocytes. Attia et al. (263) concluded that the aneuploidy predicted by Hunt et al. (262) could not be demonstrated to date and that further studies were being conducted to examine this issue.

Table 53. In Vitro Genetic Toxicity Studies of Bisphenol A

Concentration	Cell	Endpoint	Results	Reference
3.3–333.3 $\mu\text{g}/\text{plate}$, with and without metabolic activation	<i>Salmonella typhimurium</i> strains TA98, TA100, TA1535, TA1537	Mutagenicity	Negative	Haworth et al. (1983) ^{a,b}
50–500 $\mu\text{g}/\text{plate}$, with and without metabolic activation	<i>Salmonella typhimurium</i> strains TA97a, TA98, TA100, TA102	Mutagenicity	Negative	Schweickl et al. (1998) ^{a,b}
≤ 5000 $\mu\text{g}/\text{plate}$ with and without metabolic activation	<i>Salmonella typhimurium</i> strains TA97, TA98, TA100, TA102	Mutagenicity	Negative	Takahata et al. (1990) ^{a,b}
≤ 1000 $\mu\text{g}/\text{mL}$, with and without metabolic activation	<i>Salmonella typhimurium</i> strain TA1538 and <i>Escherichia coli</i> strains WP2 and WP2uvrA	Mutagenicity	Negative	Dean and Brooks (1978) ^{a,b}
5–1250 $\mu\text{g}/\text{plate}$, with and without metabolic activation	<i>Salmonella typhimurium</i> strains TA98, TA100, TA1535, TA1537 and <i>Escherichia coli</i> strain WP2uvrA	Mutagenicity	Negative	JETOC (1996) ^{a,b}
1 mM [228 mg/L], with and without metabolic activation	<i>Salmonella typhimurium</i> strains TA98 and TA100	Mutagenicity	Negative	Masuda et al. (264)
0.1–0.2 mM [23–46 mg/L], without metabolic activation	Chinese hamster V79 cells, hprt locus	Mutagenicity	Negative	Schweickl et al. (1998) ^{a,b}
5–60 mg/L without metabolic activation, 25–200 mg/L with metabolic activation, or 5–60 mg/L with and without metabolic activation ^d	Mouse lymphoma L5178Y cells, tk ^{+/-} locus	Mutagenicity	Negative (results questioned due to possible inability to count small colonies)	Myhr and Caspary (1991) ^{a,b}

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Concentration	Cell	Endpoint	Results	Reference
Concentrations not specified, with and without metabolic activation	Mouse lymphoma L5178Y cells, tk ^{+/+} locus	Mutagenicity	Inconclusive without and negative with metabolic activation	Honma et al. (1999) ^{a,b} and Moore et al. (1999) ^{a,b}
25–200 µM [5.7–46 mg/L], without metabolic activation	Syrian hamster embryo cells (Na ⁺ /K ⁺ ATPase and hprt loci)	Mutagenicity	Negative	Tsutsui et al. (1998) ^{a,b}
10 ⁻⁸ –10 ⁻⁵ M [0.002–2 mg/L], without metabolic activation	Human RSa cells	Mutagenicity	↑ at all doses	Takahashi et al. (265)
≤500 mg/L, with and without metabolic activation	<i>Saccharomyces cerevisiae</i> strain JDI	Mutagenicity	Negative	Dean and Brooks (1978) ^{a,b}
10 ⁻⁸ –10 ⁻⁴ M [0.002–23 mg/L], without metabolic activation	MCF-7 cells	DNA damage (assessed by comet assay)	↑ at ≥10 ⁻⁶ M [0.2 mg/L]	Iso et al. (266)
10 ⁻⁴ M [23 mg/L], without metabolic activation	MDA-MB-231 cells	DNA damage (assessed by comet assay)	↑	
20–40 mg/L, without metabolic activation and 30–50 mg/L with metabolic activation	Chinese hamster ovary (CHO) cells	Chromosomal aberration	Negative (inconsistent ↑ at high dose with metabolic activation)	Ivett et al. (1989) ^{a,b} and Tennant et al. (1986, 1987) ^b
350–450 µM [80–103 mg/L], without metabolic activation and ≤250 µM [57 mg/L] with metabolic activation	CHO cells, clone WBL	Chromosomal aberration	Positive at ≥400 µM [91.3 mg/L] without metabolic activation ^c ; negative with metabolic activation	Hilliard et al. (1998) ^a
400 and 450 µM [91 and 103 mg/L], without metabolic activation	CHO cells, clone WBL	Chromosomal aberration	Positive ^c	Galloway et al. (1998) ^a
25–200 µM [5.7–46 mg/L], without metabolic activation	Syrian hamster embryo cells	Chromosomal aberration	Negative	Tsutsui et al. (1998) ^{a,b}
10–30 mg/L	Epithelial-type rat liver cell line (RL1)	Chromosomal aberration	Negative	Dean and Brooks (1978) ^b

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Concentration	Cell	Endpoint	Results	Reference
25–200 μM [5.7–46 mg/L], without metabolic activation	Syrian hamster embryo cells	Aneuploidy/ polyploidy	Inconclusive (non-dose-related \uparrow in aneuploidy at $\geq 50 \mu\text{M}$ [11 mg/L] ^e ; apparently positive ^f)	Tsutsui et al. (1998) ^{a,b}
0.8–25 mg/L, without metabolic activation and 30–50 $\mu\text{g/mL}$, with metabolic activation	CHO cells	Sister chromatid exchange	Negative (one small \uparrow was not reproducible)	Ivett et al. (1989) ^{a,b} and Tennant et al. (1986) ^b
0.2–0.5 mM or nM ^d [46–114 mg/L or $\mu\text{g/L}$]	Rat hepatocytes	DNA strand breaks	Negative (\uparrow noted but scored as negative by study authors due to excessive cytotoxicity)	Storer et al. (1996) ^{a,b}
10^{-9} – 10^{-5} M [0.0002–2 mg/L], without metabolic activation	Human RSa cells	Unscheduled DNA synthesis	\uparrow at 10^{-6} M [0.2 mg/L] and \downarrow at 10^{-7} [0.02 mg/L] and 10^{-5} M [2 mg/L]	Takahashi et al. (265)
Not specified, but stated to cover range of cytotoxicity	A31-1-13 clone of BALB/c-3T3 cells	Transformation	Negative	Matthews et al. (1993) ^a
25–200 μM [5.7–46 mg/L], without metabolic activation	Syrian hamster embryo cells	Transformation	Positive at $\geq 50 \mu\text{M}$ [11.4 mg/L] (non-dose-related \uparrow) ^e ; equivocal ^f	Tsutsui et al. (1998, 2000) ^{a,b}
≤ 50 mg/L for 24 hours; ≤ 30 mg/L for 7 days, without metabolic activation	Syrian hamster embryo cells	Transformation	Negative	LeBoeuf et al. (1996) ^a
2–60 mg/L	Syrian hamster embryo cells	Transformation	Negative	Jones et al. (1988) ^b
50–200 μM [5.7–46 mg/L], without metabolic activation	Syrian hamster embryo cells	DNA adduct formation	Positive at $\geq 50 \mu\text{M}$ [11 mg/L] (dose-related \uparrow)	Tsutsui et al. (1998) ^{a,b}
1000 μg presence of peroxidase and hydrogen peroxide	Purified rat DNA	DNA adduct formation	Positive	Atkinson and Roy (115)
10–100 μM [2.3–23 mg/L], metabolic activation unknown	Bovine brain microtubule protein	Inhibited microtubule polymerization	Positive	Metzler and Pfeiffer (1995) ^a
50–200 μM [11–46 mg/L], no metabolic activation	Bovine brain microtubule protein	Inhibited microtubule polymerization	Positive (dose-related)	Pfeiffer et al. (1996) ^b

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Concentration	Cell	Endpoint	Results	Reference
20–200 μM [4.6–46 mg/L], metabolic activation unknown	Bovine brain microtubule protein	Inhibited microtubule polymerization	Positive ($\text{EC}_{50} = 150 \mu\text{M}$ [34 mg/L])	Pfeiffer et al. (1997) ^{a,b}
200 μM [46 mg/L], without metabolic activation; 100 μM [23 mg/L] for metaphase arrest assay	Chinese hamster V79 cells	Aneuploidogenic potential as assessed by micronuclei formation, microtubule assay, and metaphase arrest.	Positive	Pfeiffer et al. (1997) ^{a,b}
100–200 μM [23–46 mg/L], without metabolic activation	Chinese hamster V79 cells	Aneuploidogenic potential as assessed by micronuclei formation	Positive	Ochi (1999) ^{a,b}
0.05–0.4 mg/L	Oocytes from MF_1 mice	Aneuploidy	No hyperhaploidy but \uparrow diploid metaphase II oocytes at 0.2 mg/L	Reviewed in Attia et al. (263) (abstract)

\uparrow, \downarrow increase, decrease.

^aReviewed by Haighton et al. (24).

^bReviewed by the European Union (2).

^cAccording to the Haighton et al. (24) review, positive results occurred at cytotoxic concentrations.

^dDiscrepancies noted between information presented by Haighton et al. (24) and European Union (2).

^eConclusion by Haighton et al. (24).

^fConclusion by European Union (2).

1

1 **Table 54. In Vivo Genetic Toxicity Studies of Bisphenol A**

Species and sex	Dose (route)	Cells	Endpoint	Results	Reference
Male rat	85 mg/kg bw/day for 5 days (ip)	Germ	Dominant lethality	Negative	Bond et al. (1980) ^{a,b} (abstract only)
Male rat	200 mg/kg bw (ip) and 200 mg/kg bw for 4, 8, 12, or 16 days (oral)	DNA	Adduct formation	Positive	Atkinson and Roy (116)
Male and female mouse	500–2000 mg/kg bw (oral)	Bone marrow	Micronuclei	Negative	Gudi and Krsmanovic (1999) ^a and Shell Oil Co. (1999) ^b
Male mouse	1 mmol/kg bw [228 mg/kg bw] (oral)	Peripheral blood reticulocyte	Micronuclei	Negative	Masuda et al. (264)
20–22-day-old female mouse	0.02–0.100 mg/kg bw/day (oral) for 6–8 days or 0.02 mg/kg bw for 3, 5, or 7 days	Oocyte	Congression failure	Positive at all doses; statistically significant with 7-day exposure	Hunt et al. (262)
Female mouse	0.2 or 20 mg/kg bw or 0.04 mg/kg bw/day for 7 days	Oocyte	Aneuploidy	Negative	Reviewed in Attia et al. (263) (abstract)
Male (102/E1xC3H/E1) F ₁ mouse	0.002–0.2 mg/kg bw for 6 days (oral)	Spermatocyte	Meiotic delay and aneuploidy	Negative	Reviewed in Attia et al. (263) (abstract)
<i>Drosophila melanogaster</i>	10,000 ppm (oral)	Offspring	Sex-linked recessive lethal test	Negative	Foureman et al. (1994) ^{a,b}

^aReviewed by Haighton et al. (24).

^bReviewed by the European Union (2).

2

3 **2.4. Carcinogenicity**

4 No human data examining the carcinogenicity of bisphenol A were identified.

5

6 NTP (127, 267) examined carcinogenicity of bisphenol A in F344 rats and B6C3F₁ mice.

7 Animals were randomly assigned to treatment groups. Bisphenol A (<98.2% purity) was
8 administered through feed for 103 weeks to 50 rats/sex/dose at 0, 1000, or 2000 ppm, 50 male
9 mice/group at 0, 1000, or 5000 ppm, and 50 female mice/group at 0, 5000, or 10,000 ppm. NTP
10 estimated mean bisphenol A intakes of 74 and 148 mg/kg bw/day for male rats and 74 and 135
11 mg/kg bw/day for female rats. **[Data on bisphenol A intake, food intake, and body weights
12 were not provided for mice.]** Using default values, the European Union (2) estimated bisphenol
13 A intakes of 120 and 600 mg/kg bw/day in male mice and 650 and 1300 mg/kg bw/day in female
14 mice. Concentration and stability of bisphenol A in feed were verified. Body weights and clinical
15 signs were observed during the study. Following the exposure period, animals were killed and
16 necropsied. Organs, including seminal vesicle, prostate, testis, ovary, and uterus, were preserved

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1 in 10% neutral buffered formalin and examined histologically. Statistical analyses included Cox
2 and Tarone methods, 1-tailed Fisher exact test, Bonferroni inequality criterion, Cochran-Armitage
3 test, and life table methods for linear trend.
4

5 In rats, body weights of males and females from both dose groups were lower than controls
6 throughout the study. Feed intake was decreased in females of both dose groups beginning at
7 week 12. No adverse effects on survival were observed. There were no non-neoplastic lesions
8 **[including in male and female reproductive organs]** that appeared to be treatment-related. The
9 incidence of leukemia was increased in males (13 of 50, 12 of 50, and 23 of 50 in control and
10 each respective dose group) and females (7 of 50, 13 of 50, and 12 of 50). In males the trend for
11 leukemia was significant by Cochran-Armitage test, but statistical significance was not shown by
12 life table analysis for trend or incidence in the high-dose group, according to the unpublished
13 version of the study. The published version of the study indicated statistical significance at the
14 high dose. Statistical significance was not attained for leukemia incidence in female rats. An
15 increased incidence of testicular interstitial cell tumors (35 of 49, 48 of 50, 46 of 49) was
16 statistically significant in both dose groups. An increased incidence of mammary fibroadenomas
17 in males of the high-dose group (0 of 50, 0 of 50, and 4 of 50) achieved statistical significance for
18 trend by Cochran-Armitage test but not by Fisher exact test. In bisphenol A groups, there were
19 decreased incidences of adrenal pheochromocytomas in males, adrenal cortical adenomas in
20 females, and uterine endometrial stromal polyps. The NTP concluded that none of the increases in
21 tumor incidence in rats was clearly associated with bisphenol A exposure.
22

23 In mice, body weights were lower in high-dose males and in females of both dose groups. Feed
24 intake could not be accurately determined because of spillage. Bisphenol A did not affect the
25 survival of mice. Incidence of multinucleated hepatocellular giant cells was increased in treated
26 males (1 of 49, 41 of 49, and 41 of 50). **[A review of the data indicated no increases in**
27 **incidence of non-neoplastic lesions in the reproductive organs of male or female mice.]** The
28 incidence of leukemia or lymphoma in male mice by dose group (2 of 49, 9 of 50, and 5 of 50)
29 was not statistically significant. The published version of the report indicated an increasing trend
30 for lymphoma. The linear trend for increased pituitary chromophobe carcinomas in male mice (0
31 of 37, 0 of 36, 3 of 42) was reported to be statistically significant by Cochran-Armitage test but
32 statistical significance was not shown by Fisher exact test. The study authors concluded that none
33 of the increases in tumor incidence in mice could be unequivocally associated with bisphenol A
34 exposure.
35

36 NTP concluded that under the conditions of this study, there was no convincing evidence the
37 bisphenol A was carcinogenic in F344 rats or B6C3F₁ mice. However, study authors stated that
38 there was suggestive evidence of increased cancer in the hematopoietic system based on
39 marginally significant increases in leukemia in male rats, non-statistically significant increases in
40 leukemia in female rats, and a marginally significant increase in combined incidence of
41 lymphoma and leukemia in male mice. A statistically significant increase in testicular interstitial
42 cell tumors in aging F344 rats was also considered suggestive evidence of carcinogenesis. The
43 effect was not considered conclusive evidence because of the high incidence of the testicular
44 neoplasm in aging F344 rats (88% incidence in historical controls).
45

46 The NTP study was reviewed by the European Union (2) and Haighton et al. (24). For increases
47 in leukemia, mammary gland fibroadenoma, and Leydig cell tumors in male rats, both groups
48 noted the lack of statistical significance using the appropriate analyses and the common
49 occurrence of these tumor types in F344 rats. The European Union (2) concluded, "Overall, all of
50 these [tumor] findings in rats and mice are not considered toxicologically significant.
51 Consequently, it is concluded that bisphenol A was not carcinogenic in this study in both

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1 species.” Haighton et al. (24) concluded, “Overall, the results of this bioassay did not provide any
2 compelling evidence to indicate that [bisphenol A] was carcinogenic in F344 rats or in B6C3F₁
3 mice.” Based on the experimental animal data, the European Union concluded that “. . . the
4 evidence suggests that bisphenol A does not have carcinogenic potential.” Using a weight of
5 evidence approach, Haighton et al. (24) concluded that bisphenol A was not likely to be
6 carcinogenic to humans. This conclusion was based upon NTP study results; lack of activity at
7 noncytotoxic concentrations in both in vitro genetic toxicity tests and in an in vivo mouse
8 micronucleus test; and data from metabolism studies that show rapid glucuronidation and no
9 formation of possibly reactive intermediates, with the possible exception of reactive intermediates
10 potentially generated as a result of saturated detoxification pathways at high doses.

11 2.5 Potentially Susceptible Subpopulations

12 As noted in Section 2.1.1.3, one pathway of bisphenol A metabolism in humans and experimental
13 animals is glucuronidation. Studies in experimental animals demonstrated that both the intestine
14 and liver can glucuronidate bisphenol A. UGT2B1 was identified as the isoform involved in
15 bisphenol A glucuronidation in rat liver (117). The UDPGT isoform involved in human intestinal
16 glucuronidation of bisphenol A is not known to have been identified. Despite uncertain isoform
17 identification, studies in humans and experimental animals demonstrate developmental changes
18 in expression of activities of several UDPGT isoforms that potentially affect bisphenol A
19 metabolism.

20
21 Coughtrie et al. (268) examined the ontogeny of UDPGT activity in human liver microsome
22 samples obtained postmortem from adults and premature or full-term infants. Results of this
23 analysis are listed in Table 55. Activities for isoenzymes catalyzing glucuronidation of bilirubin,
24 testosterone, and 1-naphthol were very low at birth in premature and full-term infants. Activities
25 increased with age for the isoenzymes catalyzing glucuronidation of bilirubin (~80% of adult
26 levels by 8–15 weeks of age) and 1-naphthol (~30% of adult levels at 8–15 weeks of age). During
27 the first 55 weeks of life, no consistent increase in activity was noted for the isoenzyme
28 catalyzing glucuronidation of testosterone. Using an immunoblot technique with antibodies
29 developed toward liver testosterone/4-nitrophenol and kidney naphthol/bilirubin, 1
30 immunoreactive protein was observed in microsomes of 18- and 27-week-old fetuses and 3
31 immunoreactive proteins were observed in microsomes of full-term infants. Most isoenzymes
32 present in adults were observed in infants within 3 months of age at levels ~25% those of adults.

33 **Table 55. Development of UDPGT Activity in Humans**

Age	UDPGT activity, nmol/min/mg protein		
	Bilirubin	Testosterone	1-Naphthol
30 weeks gestation	0.05	0	0.56
30 weeks gestation with 10 weeks survival	0.4; 1	0.14; 0.85	3.0; 1.8
Full-term infants surviving 1–10 days (n = 7)	0.07 ± 0.04	0.10 ± 0.06	0.75 ± 0.68
Full-term infants surviving 8–15 weeks (n = 6)	0.64 ± 0.32	0.12 ± 0.05	2.4 ± 1.1
Full-term infants surviving 22–55 weeks (n = 5)	0.99 ± 1.1	0.09 ± 0.06	3.6 ± 2.1
Adult males (n = 3)	0.76 ± 0.43	0.46 ± 0.61	7.2 ± 2.2

Data presented as individual values or mean ± SD.

From Coughtrie et al. (268).

36
37 Strassburg et al. (269) used a reverse transcript (RT)-polymerized chain reaction (PCR) technique
38 to examine developmental changes in expression for 13 *UDPGT* genes in liver samples obtained
39 from 16 pediatric patients undergoing liver transplant for extrahepatic biliary atresia (6–24
40 months old) and 12 adults undergoing liver transplant for carcinoma (25–75 years). Changes in

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1 gene expression were also assessed in hepatic RNA samples for two 20-week-old fetuses. No
2 transcripts for UDPGT were detected in samples from 20-week-old fetuses. In infant and adult
3 livers, transcripts were detected for *UGT1A1*, *UGT1A3*, *UGT1A4*, *UGT1A6*, *UGT1A9*, *UGT2B4*,
4 *UGT2B7*, *UGT2B10*, and *UGT2B15*; there were no age-related differences in expression.
5 Expression of *UGT1A9* and *UGTB4* mRNA was lower in the pediatric samples. Western blot
6 analyses of protein expression for *UGT1A1*, *UGT1A6*, and *UGT2B7* were consistent with
7 findings for mRNA expression. Activities towards 18 specific substrates were assessed in
8 microsomes. In 13–24-month-old children compared to adults, glucuronidation activity was lower
9 for ibuprofen (24-fold), amitriptyline (16-fold), 4-tert-butylphenol (40-fold), estrone (15-fold),
10 and buprenorphine (12-fold).

11
12 Cappelletto et al. (270) compared uridine 5'-diphosphoglucuronic acid levels in livers and kidneys
13 of human fetuses and adults and in placenta. In adults undergoing surgery, liver samples were
14 obtained from 1 man and 4 women (23–72 years of age) and kidney samples were obtained from
15 1 woman and 4 men (55–63 years of age). Fetal livers and kidneys were obtained from 5 fetuses
16 legally aborted between 16 and 25 weeks gestation. Five placenta samples were obtained upon
17 delivery at 17–25 weeks gestation. Compared to adults, fetal uridine 5'-diphosphoglucuronic acid
18 concentrations were 5-fold lower in liver and 1.5-fold lower in kidney. Concentrations of uridine
19 5'-diphosphoglucuronic acid in placenta were 3–4-fold lower than in fetal liver. Based on these
20 findings, study authors concluded that glucuronidation is potentially limited in the human fetus.

21
22 As noted in Sections 2.1.2.2 and 2.1.2.3, rat fetuses appear to have no or low ability to
23 glucuronidate bisphenol A (100, 121, 122). Although rats glucuronidate bisphenol A at birth,
24 glucuronidation capacity appears to increase with age (2, 92, 122).

25
26 Some possible interindividual or sex-related differences in the ability to produce the bisphenol A
27 sulfate conjugate were identified in a limited number of human studies. As discussed in more
28 detail in Section 2.1.1.3 and shown in Table 6, higher amounts of urinary bisphenol A sulfate
29 were detected in 15 adult women than in 15 adult males (70). In a study examining bisphenol A
30 metabolism by human hepatocytes, an ~10-fold higher level of a bisphenol A glucuronide/sulfate
31 conjugate was observed in the sample from 1 female than in samples from 2 other females and 2
32 males (114).

33
34 Yang et al. (61) examined the effects of polymorphisms in sulfotransferase enzymes on urinary
35 excretion of total bisphenol A (conjugated and free) in Korean volunteers. Urinary bisphenol A
36 levels were measured by HPLC and a PCR method was used to determine sulfotransferase
37 genotype. The *SULT1A1*1* allele was reported to have greater enzyme activity than the
38 *SULT1A1*2* enzyme and it was expected that individuals with the *SULT1A1*1* allele would be
39 able to rapidly eliminate bisphenol A. However, no significant differences in urinary bisphenol A
40 levels were observed between 57 individuals with the *SULT1A1*1* allele (geometric mean \pm SD
41 = 10.10 \pm 8.71 μ g/L) and 15 individuals with the *SULT1A1*2* enzyme (6.31 \pm 8.91 μ g/L).
42 Adjustment for possible bisphenol A exposure through vinyl wrap use also did not result in
43 significant differences between the 2 groups. The study authors concluded that additional
44 enzymes involved in bisphenol A metabolism should be studied to determine possible sensitivity
45 differences.

46
47 One animal study demonstrated sex-related differences in sulfation. Male versus female Sprague
48 Dawley and F344 rats were found to produce higher amounts of a bisphenol A
49 glucuronide/sulfate conjugate (114).

50

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1 As noted in Table 5, one human study reported ~2-fold higher blood bisphenol A levels in
2 Japanese men than women (64). Based on positive correlation between serum bisphenol A and
3 testosterone levels, authors speculated that sex-related differences in bisphenol A levels might be
4 due to androgen-related metabolism (65). There are no known human studies demonstrating inter-
5 individual or sex-related variations in metabolism that could lead to higher bisphenol A levels in
6 blood. Experimental animal studies have not consistently demonstrated higher levels of bisphenol
7 A or radioactive dose in one sex (93, 101). In Wistar rats orally administered 1 mg bisphenol A
8 every 2 days for 2 or 4 weeks, liver microsomal UDPGT activity towards 17 β -estradiol and
9 testosterone and expression of UGT2B1 protein and mRNA were reduced in males but not
10 females (271). One study reported an ~3-fold higher level of blood bisphenol A in male than in
11 female Wistar–Imamichi rats that were apparently not treated, but there was no sex-related
12 difference in percent glucuronidated bisphenol A in serum (272). However, in an in vitro study
13 conducted with hepatic microsomes, glucuronidation of bisphenol A and expression of *UGT2B1*
14 mRNA were higher in microsomes from female than male rats. As described in more detail in
15 Section 2.1.2.3, one study demonstrated reduced capacity to glucuronidate bisphenol A in livers
16 from pregnant than in non-pregnant rats (120).

18 2.6 Summary of General Toxicology and Biologic Effects

20 2.6.1 Toxicokinetics and metabolism

21 Human toxicokinetic data for bisphenol A are summarized in Table 56. In humans the majority of
22 ingested bisphenol A is glucuronidated and circulated as bisphenol A glucuronide (91). There is
23 no evidence of enterohepatic circulation (91). Most of the bisphenol A dose is excreted by
24 humans through urine; bisphenol A recoveries in urine were reported at $\geq 84\%$ within 5 hours of
25 dosing (62) and 100% within 42 hours of dosing (91). Human urinary profiles were reported at
26 ~33–70% bisphenol A glucuronide, ~10–33% parent compound, and ~5–34% bisphenol A sulfate
27 conjugate (69, 70). The presence of bisphenol A in human fetal tissues or fluids demonstrates that
28 bisphenol A is distributed to the human conceptus (63, 68, 83-86). Results from a limited number
29 of studies indicated that fetal bisphenol A levels are within the same order of magnitude as
30 maternal blood levels (68, 84) and amniotic fluid bisphenol A levels are ~1 order of magnitude
31 lower than maternal blood levels (85). Median or mean Bisphenol A levels in human milk were
32 reported to be ≤ 1.4 $\mu\text{g/L}$ (87, 88). One of the studies reported a milk/serum ratio of 1.3 (88).

34 **Table 56. Human Toxicokinetic Values for Total Bisphenol A Dose**

Endpoint	Value	Reference
Oral absorption, %	$\geq 84\%$	(62, 91)
Dermal absorption, in vitro, %	~10%	(2)
T _{max} , minutes	80	(91)
Elimination half-life, hours	4–5.4	(62, 91)

35
36 Animal toxicokinetic data for bisphenol A are summarized in Table 57. Following oral intake of
37 bisphenol A by rats, most of the dose ($\geq 77\%$) is glucuronidated and circulated as bisphenol A
38 glucuronide (100, 101, 121). Most bisphenol A (90–95%) circulates bound to plasma proteins
39 (110) (reviewed in (111)). In mice injected with a low dose (0.025 mg/kg bw), the most abundant
40 compound in all tissues was bisphenol A glucuronide except in placenta where bisphenol A and
41 metabolite F (most likely bisphenol A glucuronide conjugated to acetylated galactosamine or
42 glucosamine) were the major compounds detected (108). Most of a bisphenol A dose is circulated
43 as the glucuronide in monkeys (98). As noted in Table 58, free bisphenol A in blood represents
44 $\leq 8\%$ of the dose in rats and $\leq 1\%$ of the dose in monkeys following oral dosing; higher levels of
45 free bisphenol A in blood were observed following parenteral dosing ($\geq 19\%$ in rats and $\geq 5\%$ in

2.0 General Toxicology and Biological Effects

1 monkeys). The presence of 2 or more C_{max} values for radioactivity or bisphenol A, an indication
 2 of enterohepatic circulation, was noted in some rat studies (97, 100, 101). In rats, glucuronidation
 3 of bisphenol A was demonstrated to occur in intestine (118, 119) and liver (120). UGT2B1 was
 4 identified as a liver enzyme capable of glucuronidating bisphenol A, and possible involvement of
 5 other liver isoforms was noted (117). There are some data indicating that glucuronidation
 6 capacity is very limited in fetuses and lower in immature than adult animals. Little-to-no UGT2B
 7 activity towards bisphenol A was detected in microsomes of rat fetuses; activity of the enzyme
 8 increased linearly following birth (122). In an in vitro study comparing clearance of bisphenol A
 9 by hepatic microsomes from rats of different ages, activity was lower in microsomes from fetuses
 10 than in those from immature animals and adults (reviewed in (2)). As noted in Table 57,
 11 immature rats are capable of glucuronidating bisphenol A, but activity appears to increase with
 12 age. One study demonstrated that neonatal rats were able to glucuronidate a larger fraction of a
 13 lower (1 mg/kg bw) than higher (10 mg/kg bw) bisphenol A dose (92).

14
 15 The major metabolite of bisphenol A is the glucuronide conjugate. Another metabolite that has
 16 been commonly detected in urine is bisphenol A sulfate. Excretion patterns for bisphenol A are
 17 summarized in Table 59. As noted in Table 59, the major elimination routes for bisphenol A in
 18 rodents are feces and bile; a lower percentage of the dose is eliminated through urine. The major
 19 compound detected in feces is bisphenol A and the major compound detected in bile and urine is
 20 bisphenol A glucuronide. Similar excretion patterns and metabolic profiles were observed in
 21 rodents dosed orally or parenterally with low (< 1 mg/kg bw/day) or high doses (10–100 mg/kg
 22 bw/day). In contrast to rodents and similar to humans, most of the dose in orally or iv exposed
 23 monkeys was eliminated through urine.

24
 25 **Table 57. Toxicokinetic Values for Bisphenol A in Non-Pregnant Animals**

Model	Endpoint	Value	Reference
Rats orally exposed to ≤100 mg/kg bw	T_{max} , hours	0.083–0.75	(92-96)
Ovariectomized, adult rats gavaged with bisphenol A at 10 and 100 mg/kg bw	T_{max1} / T_{max2} , hours	0.5–1.5 / 3–6	(97)
Immature rats orally dosed with ≤10 mg/kg bw	T_{max} hours	0.25–3	(92)
Monkeys orally dosed with ≤100 mg/kg bw	T_{max} , hours	0.7	(96)
Chimpanzees orally dosed with 10 mg/kg bw	T_{max} , hours	0.5	(96)
Rats sc dosed with ≤100 mg/kg bw	T_{max} , hours	1	(96)
Monkeys sc dosed with ≤100 mg/kg bw	T_{max} , hours	2	(96)
Chimpanzees sc dosed with 10 mg/kg bw	T_{max} , hours	2	(96)
Ovariectomized, adult rats orally dosed with bisphenol A at 10 and 100 mg/kg bw	Bioavailability, %	16.4 and 5.6 ^a	(97)

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Model	Endpoint	Value	Reference
Rats orally dosed with 10 mg/kg bw	Bioavailability, %	5.3	(95)
Rat	Plasma protein binding, %	90–95%.	(110); reviewed in (111)
Rats orally dosed with 10 mg/kg bw	C _{max} , µg/L	14.7–63	(92, 95)
Rats orally dosed with 100 mg/kg bw	C _{max} , µg/L	580	(96).
Ovariectomized, adult rats orally dosed with (mg/kg bw):	C _{max1} /C _{max2} , µg/L		(97)
10		30/40	
100		150/134	
Oral doses (mg/kg bw) in immature rats at each age:	C _{max} (µg/L)	Range of values for males and females:	(92)
1 (PND 4)		30–60	
10 (PND 4)		10,200–48,300	
1 (PND 7)		40–80	
10 (PND 7)		1100–1400	
1 (PND 21)		5–6	
10 (PND 21)		200	
Monkeys orally dosed with 10 and 100 mg/kg bw	C _{max} , µg/L	2793 and 5732 ^a	(96)
Monkeys orally dosed with 10 mg/kg bw	C _{max} , µg/L	96–325	(96)
Rats sc dosed with 10 and 100 mg/kg bw	C _{max} , µg/L	872 and 3439 ^a	(96)
Monkeys sc dosed with 10 and 100 mg/kg bw	C _{max} , µg/L	57,934 and 10,851 ^a	(96)
Chimpanzees sc dosed with 10 mg/kg bw	C _{max} , µg/L	1026–2058	(96)
Oral doses (mg/kg bw) in immature rats at each age:	AUC, µg-hour/L	Range of values for males and females:	(92)
1 (PND 4)		100–200	
10 (PND 4)		7200–23,100	
1 (PND 7)		100	
10 (PND 7)		1700–1900	
10 (PND 21)		1000–1100	
Rats orally dosed with 10 mg/kg bw	AUC, µg-hour/L	85.6	(95)
Rats orally dosed with 100 mg/kg bw	AUC _{0–24h} , µg-hour/L	1353	(96).
Monkeys orally dosed with 10 and 100 mg/kg bw	AUC _{0–24h} , µg-hour/L	3247 and 52,595 ^a	(96).

2.0 General Toxicology and Biological Effects

Model	Endpoint	Value	Reference
Chimpanzees orally dosed with 10 mg/kg bw	AUC _{0-24h} , µg-hour/L	813-1167	(96).
Rats sc dosed with 10 and 100 mg/kg bw	AUC _{0-24h} , µg-hour/L	3377 and 23,001 ^a	(96).
Monkeys sc dosed with 10 and 100 mg/kg bw	AUC _{0-24h} , µg-hour/L	3247 and 39,040 ^a	(96).
Chimpanzees sc dosed with 10 mg/kg bw	AUC _{0-24h} , µg-hour/L	12,492-21,141	(96).
Rats orally dosed with 10 mg/kg bw	Apparent volume of distribution, L/kg	4273	(95)
Immature rats orally dosed with ≤10 mg/kg bw	Half-life, hours	4.3-21.8	(92)
Rats orally dosed with 10 mg/kg bw	Terminal elimination half-life, hours	21.3	(95)
Rats orally dosed with 10 mg/kg bw	Oral clearance, mL/minute/kg	2352.1	(95)

^aResults presented for low and high dose

1

2 **Table 58. Factors Affecting Free Bisphenol A Levels in Blood**

Model and Regimen	Findings for free bisphenol A in blood	Reference
Effects of oral dosing at:		(92)
4 days of age	10.2-48.3 mg/L	
7 days of age	1.1-1.4 mg/L	
21 days of age	0.2 mg/L	
adulthood	0.024-0.063 mg/L	
SC or gavage dosing of immature rats with 160 mg/kg bw/day	[93% lower] with oral than sc dosing	(99)
Route effects in rats administered 10 or 100 mg/kg bw:		(93)
oral	[<2-8%]	
sc	[65-76%]	
ip	[27-51%]	
Route effects in monkeys:	Percent of dose:	(98)
iv	5-29%	
oral	0-1%	

3

4 **Table 59. Summary of Elimination Information for Bisphenol A**

Model	Elimination route	Percent dose eliminated	Compound and metabolite profile	Reference
Pregnant or non-pregnant rats orally, ip, or sc dosed with <100 mg/kg bw	Feces	50-83%	Bisphenol A (83-93%); bisphenol A glucuronide (2-3%)	(93, 100, 103)
	Urine	13-42%	Bisphenol A (3-23%); bisphenol A glucuronide (57-87%); bisphenol A sulfate (2-7%)	

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Model	Elimination route	Percent dose eliminated	Compound and metabolite profile	Reference
Rats orally or iv dosed with 0.1 mg/kg bw	Feces Urine	64–88% 10–34%	Not reported	(110)
Rats orally or iv dosed with 0.1 mg/kg bw	Bile	45–66% within 5 hours	Bisphenol A glucuronide (84–88%)	(110)
Rats orally dosed with 100 mg/kg bw/day	Feces Urine Bile	Not reported	Bisphenol A (61% of dose) Bisphenol A and bisphenol A sulfate ($\leq 1.1\%$ of dose); bisphenol A glucuronide (6.5% of the dose) Bisphenol A glucuronide (41% of dose)	(110)
Pregnant mice injected with 0.025 mg/kg bw bisphenol A	Feces Urine Bile	Not reported	Bisphenol A (>95%) Major metabolites: bisphenol A glucuronide and hydroxylated bisphenol A glucuronide Bisphenol A glucuronide (>90%)	(108)
Monkeys orally or iv dosed with 0.1 mg/kg bw	Feces Urine	2–3% 79–86%	Not reported	(98)

1
2 Toxicokinetics of bisphenol A were examined in pregnant rats and are summarized in Table 60
3 for free bisphenol A and Table 61 for total dose. One study demonstrated similar disposition,
4 metabolism, and elimination of bisphenol A in pregnant and non-pregnant rats (100). A number
5 of rodent studies demonstrated distribution of bisphenol A or radioactive dose to fetuses
6 following oral dosing of the dam (94, 100-102, 105, 107). Bisphenol A distribution to fetus was
7 also demonstrated with iv dosing of rats (106) and sc dosing of mice or monkeys (108, 109). In a
8 study in which bisphenol A was orally administered to rats on GD 19, bisphenol A glucuronide
9 was not detected in fetuses (121); study authors noted the possibilities that bisphenol A
10 glucuronide was not likely transferred from dams to fetuses and that fetuses do not likely possess
11 glucuronidation ability. Some of the studies demonstrated slower elimination of bisphenol A from
12 fetuses than maternal blood following oral dosing (94, 102).

13
14 Toxicokinetics data in lactating rats are summarized in Table 62 for free bisphenol A and Table
15 63 for total dose. Distribution of bisphenol A to milk and/or nursing pups was demonstrated in
16 rodent studies with oral or iv exposures (95, 101, 103). One study reported that most of the
17 bisphenol A dose is present as bisphenol A glucuronide in milk of lactating rats (103). In a study
18 that compared bisphenol A levels in maternal serum, milk, and offspring after rat dams were
19 administered low oral doses (0.006 or 6 mg/kg bw/day), a significant increase in bisphenol A
20 level was only observed in the serum of dams from the high dose group on PND 21; no increase
21 was observed in milk or pups (104). Another study demonstrated higher levels of bisphenol A in
22 milk compared to maternal serum after iv dosing of rat dams (95).

23

2.0 General Toxicology and Biological Effects

1 **Table 60. Toxicokinetic Values for Bisphenol A in Pregnant Rats and Fetuses**

Dose	Endpoint	Maternal	Fetal	Reference
1000 mg/kg bw orally on GD 18	C _{max} , µg/L	14,700	9220	(94)
10 mg/kg bw orally on GD 19	Concentration 1 hour post dosing, µg/L	34	11	(102)
2 mg/kg bw iv on 1 day between GD 17 and 19	C _{max} , µg/L	927.3	794	(106)
1000 mg/kg bw orally on GD 18	T _{max} , minutes	20	20	(94)
2 mg/kg bw iv on 1 day between GD 17 and 19	T _{max} , hours	No data	0.6 ± 0.3	(106)
1000 mg/kg bw orally on GD 18	AUC, µg·hour/L	13,100	22,600	(94)
2 mg/kg bw iv on 1 day between GD 17 and 19	AUC, µg·hour/L	905.5	1964.7	(106)
1000 mg/kg bw orally on GD 18	Mean retention time, hours	10.6	20.0	(94)
1000 mg/kg bw orally on GD 18	Variance in retention time, hours squared	203	419	(94)
2 mg/kg bw iv on 1 day between GD 17 and 19	Mean residence time, hours	3.0	3.0	(106)
1000 mg/kg bw orally on GD 18	Half-life, hours:			(94)
	From 20 to 40 minutes	0.0952	0.55	
	From 40 minutes to 6 hours	2.58	1.60	
	From 6 to 48 hours	4.65	173	
2 mg/kg bw iv on 1 day between GD 17 and 19	Elimination half-life, hours	2.5	2.2	(106)

2

3 **Table 61. Toxicokinetic Values for Radioactive Dose in Pregnant Rats**

Endpoint	Value
C _{max1} / C _{max2} , µg eq/L	370–1006/211–336
T _{max1} / T _{max2} , hours	0.25/12–24
Time to non-quantifiable level, hours	72–96
AUC ¹⁴ C, µg·eq·hour/L	7100–12,400
AUC Bisphenol A glucuronide, µg·eq·hour/L	6800–12,300

Dams were gavaged with 10 mg/kg bw/day on GD 6–10, 14–18, or 17–21. From Dormoradzki et al. (100)

4

5 **Table 62. Toxicokinetic Values for Bisphenol A in Nursing Rats**

Endpoint	Blood Value	Milk Value
Systemic clearance, mL/minute/kg	119.2 / 142.4 / 154.1 ^a	
Steady state bisphenol A concentration, ng/mL	66.1 / 120.0 / 217.1 ^a	173.1 / 317.4 / 493.9 ^a
Milk/serum ratio		2.7 / 2.6 / 2.4 ^a

Rats were iv injected 0.47, 0.94, or 1.88 mg/kg bw and then infused over a 4 hour time period with 0.13, 0.27, 0.54 mg/hour.

^aEffect at each dose, from low to high dose. From Yoo et al. (95)

6

7 **Table 63. Toxicokinetic Values for Radioactive Dose in Nursing Rats**

Endpoint	Blood Value	Milk Value
C _{max} , µg·eq/L	27.2	4.46
T _{max} , hours	4	8
Elimination half-life, hours	31	26
AUC (0–48 hours), µg·eq·hour/L	689	156

Rats were orally dosed with 0.5 mg/kg bw on PND 11.

From Kurebayashi et al.(101)

8

2.0 General Toxicology and Biological Effects

1 A number of in vitro studies compared bisphenol A metabolic velocity rates in microsome or
2 hepatocytes from rodents and humans. Generally, faster rates were demonstrated by rodent than
3 human hepatocytes and microsomes (113, 114) and (reviewed in (2)). One of the studies noted
4 that adjustment for total hepatocyte number in vivo resulted in higher predicted rates for humans
5 than rodents (114). The European Union (2) noted that the interpretation of such studies should
6 included knowledge about in vivo conditions such as varying metabolic capacity of hepatic cells,
7 relationship of hepatic size to body size, and possibly important physiological endpoints such as
8 blood flow.

9 10 2.6.2 General Toxicity

11 Gross signs of toxicity observed in rats acutely exposed to bisphenol A included pale livers, and
12 gastrointestinal hemorrhage (reviewed by the European Union (2)). Acute effects of inhalation
13 exposure in rats included transient and slight inflammation of nasal epithelium and ulceration of
14 the oronasal duct. Based on LD₅₀s observed in animals, the European Union (2) concluded that
15 bisphenol A is of low acute toxicity through all exposure routes relevant to humans. According to
16 the European Union (2), there is evidence that bisphenol A is irritating and damaging to the eye
17 and is irritating to the respiratory tract but not the skin. Findings regarding sensitization potential
18 were not clear.

19
20 Possible target organs or systems of toxicity identified in repeat-dose animal studies with oral
21 dosing included cecum, liver, kidney, and male and female reproductive systems (reviewed in (2,
22 127, 128)). Cecal findings (effect levels) in rats included enlargement (≥ 25 mg/kg bw/day) and
23 mucosal hyperplasia (≥ 200 mg/kg bw/day). Hepatic effects included prominent hepatocyte nuclei
24 or inflammation in rats (≥ 500 mg/kg bw/day), multinucleated giant hepatocytes in mice (≥ 120
25 mg/kg bw/day), and increased weight with no evidence of histopathology in dogs (≥ 270 mg/kg
26 bw/day). Renal tubule degeneration or necrosis was observed in rats dosed with ≥ 500 mg/kg
27 bw/day. Reproductive findings in rats included seminiferous tubule degeneration and arrested
28 spermatogenesis (≥ 235 mg/kg bw/day), and disrupted estrous cycles (≥ 600 mg/kg bw/day).
29 Effects in subchronic inhalation studies in rats included cecal enlargement resulting from
30 distention by food and transient, slight hyperplasia and inflammation of epithelium in the anterior
31 nasal cavity; both effects occurred at (≥ 50 mg/m³).

32 33 2.6.3 Estrogenicity

34 Estrogenicity of bisphenol A has been evaluated using in vitro (Table 48) and in vivo (Table 49)
35 assays. In those studies estrogenic potency was compared to 17 β -estradiol, ethinyl estradiol,
36 diethylstilbestrol, and, in one study, estrone. There is considerable variability in the results of
37 these studies with the estrogenic potency of bisphenol A ranging over about 8 orders of
38 magnitude (Figure 2). Possible reasons for variability in in vitro findings include:

- 39
40
- 41 • Use of different methods for evaluating response (e.g., comparing differences in
 - 42 fractional response versus responses of equimolar test compound and reference estrogen),
 - 43 • Use of different receptor subtypes (ER α vs. ER β)
 - 44 • Interlaboratory differences (e.g., differences in methods such as counting procedures)
 - 45 • Use of different cell types

46 Most in vivo estrogenicity studies examined effects on uterine weights of immature or
47 ovariectomized rats or mice. The potency of bisphenol A in increasing uterine weight varied over
48 ~4 orders of magnitude. Uterine weight findings can be affected by the time period between
49 dosing and measurement of uterine weight (increased uterine weight 6 hours after treatment
50 represents fluid inhibition and not true tissue growth). Most but not all studies showed a greater

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1 effect on uterine weight with sc than with oral dosing. The greater activity of sc than oral
2 bisphenol A is presumably due to glucuronidation of the orally administered compound with
3 consequent loss of estrogenicity (138). Inter-strain variability in rats has been evaluated as a
4 source of variability in estrogenicity assays. Greater sensitivity of F344 than Sprague Dawley rats
5 was shown with respect to uterine weight and epithelial cell height (207), BrdU labeling of
6 vaginal epithelium (208), and increase in serum prolactin (194). In another study, bisphenol A
7 exposure increased uterine weight in DA/Han and Sprague Dawley rats but not in Wistar rats
8 (209). Inter-laboratory variability has been noted for uterotrophic effects in immature mice
9 exposed to bisphenol A (210); one factor that can result in variability is body weight of the
10 animal. Use of mice with lower body weights results in lower and less variable control uterine
11 weights and greater likelihood of bisphenol A effect (210, 211). In in vivo studies examining
12 gene expression profiles, some but not all gene expression changes were consistent between
13 bisphenol A and reference estrogens (210, 246, 247, 248); ER-independent activity was
14 suggested by 1 investigator (248). [The Expert Panel noted that oral bisphenol A does not
15 consistently produce estrogenic responses and, when seen, estrogenic effects after oral
16 treatment occur at high dose levels.]

17 2.6.4 Androgenic activity

18 In the majority of in vitro tests conducted, bisphenol A was not demonstrated to have androgenic
19 activity (165, 183, 188, 255). Anti-androgenic activity was demonstrated in in vivo systems using
20 cells transfected with androgen receptor reporting systems (Table 49). No consistent effects were
21 observed on male accessory reproductive organ weights in 3 in vivo studies in which rats were
22 dosed with bisphenol A at ≤ 600 mg/kg bw/day; the study authors concluded that bisphenol A
23 does not have anti-androgenic or androgenic activity (259-261).

24 2.6.5 Genetic Toxicity

25
26 In in vitro genetic toxicity studies reviewed by the European Union (2) and/or Haighton et al.
27 (24), evidence of aneugenic potential, chromosomal aberration, micronuclei formation, and DNA
28 adducts were observed (Table 51). Because of the lack of chromosomal effects in in vivo studies
29 (Table 52) and unknown relevance of DNA adduct formation, which only occurred at high doses,
30 both groups concluded that bisphenol A is not likely to have genotoxic activity in vivo. Subsequent
31 to the release of the European Union (2) and Haighton et al. (24) reviews, Hunt et al. (262)
32 reported that damaged polycarbonate caging was associated with increased congression failure in
33 mice. Hunt et al. (262) concluded that bisphenol A was a potential meiotic aneugen. Studies
34 examining the aneugenic potential of bisphenol A have been conducted, but results are currently
35 available only in abstract form (263).

36 2.6.6 Carcinogenicity

37
38 Carcinogenic potential of bisphenol A was evaluated in rats and mice by the NTP (127, 267).
39 NTP concluded that under the conditions of the study, there was no convincing evidence that
40 bisphenol A was carcinogenic in F344 rats or B6C3F₁ mice. However, NTP stated that there was
41 suggestive evidence of increased cancer in the hematopoietic system based on marginally
42 significant increases in leukemia in male rats, non-statistically significant increases in leukemia in
43 female rats, and a marginally significant increase in combined incidence of lymphoma and
44 leukemia in male mice. A statistically significant increase in testicular interstitial cell tumors in
45 aging F344 rats was also considered suggestive evidence of carcinogenesis. The effect was not
46 considered conclusive evidence because of the high incidence of the testicular neoplasm in aging
47 F344 rats (88% incidence in historical controls). Both the European Union (2) and Haighton et al.
48 (24) stated that the evidence does not suggest carcinogenic activity of bisphenol A in rats or mice.
49 Conclusions by the European Union and Haighton et al. were based on factors such as lack of
50
51

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1 statistical significance for leukemia, mammary gland fibroadenoma, and Leydig cell tumors, lack
2 of activity at noncytotoxic concentrations in both in vitro genetic toxicity tests and an in vivo
3 mouse micronucleus test, and unlikely formation of reactive intermediates at doses that do not
4 saturate detoxification pathways.

6 *2.6.7 Potentially Sensitive Subpopulations*

7 Studies in humans and animals demonstrated development changes in UDPGT gene expression or
8 enzyme activity that could potentially affect capacity for bisphenol A glucuronidation. In humans,
9 activities for some UDPGT isozymes were reported to be very low at birth but increased with age
10 (268). No transcripts for UDPGT were detected in samples from 20-week-old human fetuses and
11 activity for some UDPGT enzymes was lower in children than adults (269). Compared to adults,
12 human fetal uridine 5'-diphosphoglucuronic acid concentrations were 5-fold lower in liver and
13 1.5-fold lower in kidney (270). It is not clear if any of the isozymes examined are involved in
14 bisphenol A glucuronidation by humans. Human findings were consistent with rodent studies
15 that demonstrated no or limited glucuronidation capacity by fetuses (100, 121, 122) and lower
16 glucuronidation capacity in immature than adult rats (2, 92, 122).

17
18 Some studies suggested possible gender-related differences in sulfation capacity in humans (70,
19 114) and animals (114). One study in humans demonstrated no differences in urinary bisphenol A
20 levels in individuals carrying a sulfotransferase genotype associated with greater activity (61). A
21 study in humans demonstrated higher blood bisphenol A levels in males than in females (64), but
22 there are no consistent or conclusive experimental animals studies demonstrating sex-related
23 differences in bisphenol A body burden or metabolism capacity.
24

3.0 DEVELOPMENTAL TOXICITY DATA

3.1 Human

No studies were located on possible human developmental effects of bisphenol A.

3.2 Experimental animal

Studies are presented by species (rat, mouse, other), route (oral, parenteral), and by whether exposure was during pregnancy or the postnatal period. Studies in which exposures were started during pregnancy and continued after pregnancy are discussed with studies in which exposures occurred postnatally.

3.2.1 Rat—oral exposure only during pregnancy

3.2.1.1 Evaluation of pre- or perinatal growth and development

Morrissey et al. (273), supported by NTP/NCTR, examined the effects of prenatal bisphenol A exposure in rats and mice in a study conducted according to GLP. Studies are also available as NTP publications for rats (274) and mice (275). The study was conducted in two sets of rats and mice, and data were pooled for each species. **[The data for mice are discussed in Section 3.2.5.1.]** Pregnant CD rats were randomly assigned to groups of ≥ 10 animals in each set of the study, for a total of ≥ 20 animals/dose. On GD 6–15 (GD 0 = sperm or plug), rats were gavaged with bisphenol A at 0 (corn oil vehicle), 160, 320, 640, or 1280 mg/kg bw/day. Doses were based on results of preliminary studies and were expected to result in 10% maternal mortality at the high dose and no toxicity at the low dose. Purity of bisphenol A was $>95\%$ and 2,4'-bisphenol A was reported as an impurity. Dosing solution concentrations were verified. Pregnant animals were weighed during the study. Rats were killed on GD 20. Liver and uterus were weighed, and corpora lutea and implantation sites were examined. Fetuses were sexed, weighed, and examined for viability and external, visceral, and skeletal malformations. Data were analyzed by Bartlett test for homogeneity of variance, ANOVA and/or William multiple comparison, Dunnett, or Fisher exact probability tests.

An unexpectedly high number of dams (7 of 27) died in the 1280 mg/kg bw/day group, with most deaths occurring in the second set of animals. Because of the high death rate, the study authors decided not to evaluate data in the 1280 mg/kg bw/day group. Clinical signs that occurred most frequently in dams from the 640 mg/kg bw/day group included lethargy, piloerection, pica, rough coat, wet urogenital area, weight loss, and alopecia. Significant and dose-related decreases in maternal body weights were observed during the entire gestation period and thus were not confined to the GD 6–15 treatment period in rats from the 160, 320, and 640 mg/kg bw/day groups. Body weight corrected for gravid uterine weight was also decreased in all three dose groups. Effects on maternal body weight were most pronounced during the treatment period. **[During the treatment period, dam body weights were 35, 53, and 54% lower in the 160, 320, and 640 mg/kg bw/day groups than in control groups; estimated benchmark doses¹ in mg/kg bw/day were BMD₁₀ 113, BMDL₁₀ 94, BMD_{1SD} 416, BMDL_{1SD} 321].** Despite this large effect on maternal body weight, there were no effects on numbers of implantation sites or resorptions, gravid uterine weight, or liver weight. The numbers of litters available for evaluation in the control and 160, 320,

¹ Benchmark doses are used commonly in a regulatory setting; however, they are used in this report when the underlying data permit their calculation, and are only supplied to provide 1 kind of description of the dose-response relationship in the underlying study. Calculation of a benchmark dose in this report does not mean that regulation based on the underlying data is recommended, or even that the underlying data are suitable for regulatory decision-making. The BMD₁₀ is the benchmark dose associated with a 10% effect, estimated from a curve fit to the experimental data. The BMDL₁₀ represents the dose associated with the lower 95% confidence interval around this estimate. Unless otherwise indicated, BMD values in this report were calculated using a power model for continuous data and a probit model for dichotomous data using Environmental Protection Agency (EPA) Benchmark Dose Software version 1.3.2.

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1 and 640 mg/kg bw/day dose group were 23, 26, 24, or 29. There were no significant effects on fetal body
2 weight or viability, percentage males/litter, or malformed fetuses/litter. Study authors concluded that
3 bisphenol A was not teratogenic in rats at doses that cause maternal toxicity.
4

5 **Strengths/Weaknesses:** This study used adequate sample sizes to evaluate the effects of GD 6–15
6 exposure on maternal body weight during gestation and on implantation and resorption sites/dam, fetal
7 body weight, and fetal viability to GD 20. A strength is the verification of dosing solutions. Maternal
8 toxicity was observed through body weight reductions in dams receiving 160, 320, and 640 mg/kg bw/day
9 doses while 1280 mg/kg bw/day was associated with lethality. This is a “traditional” embryofetal
10 development study with all the attendant strengths (GLP, adequate n, sensitive evaluation of soft and
11 hard-tissue structures at birth) and weaknesses (no examination of any system as it matures postnatally).
12 The absence of effects on fetal endpoints despite marked reductions in maternal body weight corrected for
13 gravid uterine weight warrants the appropriate conclusion that bisphenol was not teratogenic when based
14 on GD 20 data. Weaknesses include the absence of data from the 1280 mg/kg bw/day group and the
15 absence of a no effect dose. Also, absence in all groups of information about the birth process, postnatal
16 viability, and postnatal function is a weakness. Further, a gross visceral exam is likely insensitive to
17 certain abnormalities of the reproductive tract and brain, as noted above.
18

19 **Utility (Adequacy) for CERHR Evaluation Process:** This study is adequate for the evaluation process.
20

21 **Kim et al. (276)**, support not indicated, examined the effects of prenatal bisphenol A exposure on
22 developmental toxicity in rats. Sprague Dawley rats were fed commercial rodent chow (Jeil Feed Co.,
23 Daejon, Korea) and housed in polycarbonate cages; no information was provided about bedding. Twenty
24 dams/group were gavaged with 0 (corn oil vehicle), 100, 300, or 1000 mg/kg bw/day bisphenol A on GD
25 1–20 (GD 0 = first 24 hours after detection of vaginal sperm or plug). Dose selection was based on the
26 results of a preliminary study that demonstrated maternal and developmental toxicity at doses ≥ 400 mg/kg
27 bw/day and a lack of effect at doses ≤ 200 mg/kg bw/day. Endpoints examined in dams during the study
28 were clinical signs, body weight gain, and food intake. Dams were killed on GD 21 and examined for
29 corpora lutea and implantation sites. Fetuses were sexed, weighed, and examined for viability and
30 external abnormalities. Anogenital distance was measured and alternate fetuses were examined for
31 visceral and skeletal malformations. The dam or litter was considered the statistical unit. Data were
32 analyzed by ANOVA, Scheffé multiple comparison test, Kruskal-Wallis nonparametric ANOVA, Mann-
33 Whitney *U* test, and Fisher exact probability test.
34

35 Statistically significant effects are summarized in Table 64. Dose-dependent clinical signs observed in
36 dams at the 2 highest doses included piloerection, dull fur, reduced locomotor activity, emaciation,
37 sedation, red-colored tears, soft stool, diarrhea, urination, and perineal soiling. Pregnancy failure, as
38 observed by lack of implantation sites, was increased in females from the high-dose group. Maternal body
39 weight, body weight gain, and body weight corrected for gravid uterus weight were reduced at the mid
40 and high dose. GD 4 was the only time period when food intake was significantly reduced at the mid and
41 high dose. Expansion and congestion of stomach and/or intestines were observed in dams from the high-
42 dose group. Body weights of male fetuses were decreased at the mid and high dose, and body weights of
43 female fetuses were reduced at the high dose. Increases in fetal death, early resorption, and
44 postimplantation loss, accompanied by reduced number of live fetuses, were observed at the high dose.
45 Anogenital distance was significantly reduced in males from the mid- and high-dose groups, but there
46 were no differences in anogenital distance of males or females when the values were normalized by the
47 cube root of body weight. Significantly reduced ossification was observed in the high-dose group. There
48 were no treatment-related differences in fetal sex ratio or external, visceral, or skeletal malformations.
49 Study authors concluded that exposure of rats to a maternally toxic dose of bisphenol A during the entire
50 gestation period resulted in pregnancy failure, postimplantation loss, reduced fetal body weight, and
51 retarded fetal ossification but not dysmorphogenesis.

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Table 64. Maternal and Developmental Effects in Rats Exposed to Bisphenol A

Endpoint	Dose, mg/kg bw/day						
	100	300	1000	BMD ₁₀	BMDL ₁₀	BMD _{1SD}	BMDL _{1SD}
Dams							
Number pregnant	↔	↔	↓30%				
Body weight gain	↔	↓35%	↓52%	178	152	379	304
Corrected body weight	↔	↓14%	↓15%	631	490	566	424
Food intake on GD 4	↔	↓24%	↓57%	168	147	313	257
No. fetal deaths	↔	↔	↑6.5-fold	827	13	978	585
No. early resorptions	↔	↔	↑6-fold	821	14	980	584
Postimplantation losses	↔	↔	↑11-fold	1278	394		
Fetuses							
No. live /litter	↔	↔	↓36%	929	348	982	713
Male body weight	↔	↓14%	↓20%	456	339	694	497
Female body weight	↔	↔	↓21%	439	328	682	490
Ossification	↔	↔	↓				

↑,↓ Statistically significant increase, decrease compared to controls; ↔ No statistically significant effect compared to controls.

From Kim et al. (276).

Strengths/Weaknesses: This report presents a fairly standard embryo-fetal developmental toxicity study. One strength is that the doses utilized incorporated both a no effect dose and a high maternally toxic dose, revealing fetal effects only at the high dose that showed marked maternal toxicity. Measurement of anogenital distance is another strength. Weaknesses include the absence in all groups of information about the birth process, postnatal viability, and postnatal function. Further, a gross visceral exam is likely insensitive to certain abnormalities of the reproductive tract and brain. However, this type of study does report on the ability of the exposure to cause structural malformations, which are notably absent.

Utility (Adequacy) for CERHR Evaluation Process: This study is adequate for the evaluation process.

Kim et al. (105), support not indicated, examined the effects of prenatal bisphenol A exposure on postnatal body and organ weights of Sprague Dawley rats. Rats were housed in polycarbonate cages. **[No information was provided on feed or bedding material.]** Rats were grouped according to body weight and randomly assigned to dose groups. On GD 7–17 (GD 0 = day of vaginal sperm or plug), at least 10 rats/dose group were gavaged with bisphenol A (>99.7% purity) at doses of 0 (corn oil vehicle), 0.002, 0.020, 0.200, 2, or 20 mg/kg bw/day. Dosing solution concentrations were verified. Dams were weighed and observed for clinical signs of toxicity during the study. Dams were killed on the 21st day of the postpartum period. Corpora lutea, implantation sites, resorptions, and fetal viability were assessed. Maternal liver, kidney, spleen, ovary, and gravid uterus were weighed. Live fetuses were weighed and examined for external and visceral abnormalities. Fetal liver, kidneys, spleen, and reproductive organs were weighed in half the fetuses. **[These methods are produced here as written in the original; although dams were clearly stated to have been killed on PND 21, the “fetal” examinations described appear more consistent with killing of the dams on GD 21.]** Data were analyzed by ANOVA and Student *t*-test.

A significant but non-dose-related increase in dam body weight occurred in the 0.2 mg/kg bw/day group on GD 0–15. Dam body weight was significantly increased on GD 21 in the 2 (by 53%) and 20 (by 43%) mg/kg bw/day groups. No significant differences in dam body weight were noted during the lactation period. Significant changes in dam relative organ weights (dose at which effects were observed) were: increased liver (0.002, 0.020, and 20 mg/kg bw/day); decreased right kidney (0.2 mg/kg bw/day);

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1 increased right kidney (2 mg/kg bw/day), and increased uterine (0.2 mg/kg bw/day). There was no effect
 2 on ovary weight of dams. The majority of dams were in diestrus when killed. One of 7 dams in the 0.2
 3 mg/kg bw/day group was in proestrus. One of 7 dams in the 0.2 mg/kg bw/day, 1 of 6 dams in the 2
 4 mg/kg bw/day group, and 2 of 8 dams in the 20 mg/kg bw/day group were in diestrus. Body weight
 5 effects in offspring are summarized in Table 65. **[Changes occurred at most dose levels but were not
 6 consistent over time and there was little evidence of dose-response relationships. In general, effects
 7 appeared to be most pronounced in the lowest dose group.]** Table 66 summarizes relative organ
 8 weight effects attaining statistical significance at 1 or more doses in offspring. There were no effects on
 9 ovary or uterus weights. **[In most cases, there was little evidence of a dose-response relationships for
 10 organ weights, including male reproductive organs, in offspring.]** Study authors concluded that
 11 bisphenol A had estrogenic effects on rat dams and offspring exposed during the gestation period.
 12

13 **Table 65. Postnatal Body Weight Effects in Rats Exposed to Bisphenol A During Gestation**

Dose, mg/kg bw/day	Age at evaluation, days					
	1	4	7	14	21	22
<i>Females</i>						
0.002	↑14%	↑16%	↑58%	↑43%	↑61%	↔
0.020	↔	↔	↔	↑28%	↑46%	↑65%
0.2	↔	↔	↑32%	↑34%	↑54%	↑62%
2.0	↑18%	↑16%	↑31%	↑24%	↑24%	↔
20	↑21%	↑15%	↑31%	↑12%	↔	↔
<i>Males</i>						
0.002	↔	↔	↑35%	↑26%	↑40%	↑32%
0.020	↔	↓13%	↔	↔	↑17%	↔
0.2	↔	↔	↔	↔	↔	↔
2.0	↑14%	↔	↔	↔	↔	↔
20	↑14%	↑13%	↑13%	↔	↔	↔

↑,↓ Statistically significant increase, decrease compared to controls; ↔ no statistically significant effects compared to controls. From Kim et al. (105).

14
 15 **Table 66. Relative Organ Weights in Rats Exposed to Bisphenol A During Gestation**

Organ	Dose, mg/kg bw/day				
	0.002	0.020	0.200	2	20
<i>Females</i>					
Liver	↔	↑20%	↑9%	↑9%	↑31%
Spleen	↔	↑49%	↑35%	↔	↔
Right kidney	↓5.5%	↔	↔	↔	↑10%
<i>Males</i>					
Liver	↔	↑13%	↑13%	↔	↑29%
Spleen	↔	↑37%	↑42%	↔	↔
Left kidney	↓9%	↔	↔	↔	↔
Right kidney	↔	↔	↑9%	↔	↑9%
Left testis	↔	↓18%	↔	↑10%	↔
Right testis	↔	↓20%	↔	↔	↔
Right epididymis	↔	↔	↔	↔	↑2.4-fold
Left Seminal vesicle	↔	↓36%	↔	↔	↔
Prostate gland	↔	↓24%	↔	↔	↔

↑,↓ Statistically significant increase, decrease compared to controls; ↔ no statistically significant effects compared to controls. From Kim et al. (105).

16

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1 **Strengths/Weaknesses:** While the verification of the dosing solution is a strength, this study is of unclear
2 quality, to the point that there is real confusion about what was actually done. It is indicated that 10 dams
3 were assigned to each dose group but numbers at sacrifice were 7, 7, 6, and 8 across the 4 doses. It is
4 unclear whether fetal data were appropriately analyzed with litter as the unit. It is unclear when the dams
5 were killed and analyzed. The absence of dose-related effects complicates interpretation at these low
6 doses; however, the possibility of unusual low dose effects cannot be discounted.

7
8 **Utility (Adequacy) for CERHR Evaluation Process:** This study is inadequate for inclusion into the
9 evaluation process, due to small sample size and poor documentation and communication about what was
10 done.

11 3.2.1.2 Evaluation of reproductive organ development

12 **Talsness et al. (277)**, supported by the German Federal Ministry for Environmental Protection and
13 Radiation Security, examined the effect of prenatal bisphenol A exposure on the reproductive systems of
14 male and female rats. **[No information was provided about feed, caging, and bedding materials used.]**
15 On GD 6–21, Sprague Dawley rats (n = 18–20/group) were gavaged with 2% corn starch vehicle or
16 bisphenol A at 0.1 or 50 mg/kg bw/day. A group of 11 dams was gavaged with 0.2 mg/kg bw/day ethinyl
17 estradiol. Litters were weighed during the lactation period. Pups were weaned on PND 22 (according to
18 Table 1 of the study, PND 1 was apparently the day of birth) and males and females were separated
19 around PND 30. Vaginal opening was examined in 42–91 female offspring/group, and estrous cyclicity
20 was monitored over a 3-week period in 42–53 females/group. At 4 months of age, 5–10 females/group
21 were killed during diestrus and 20 females/group were killed while in estrus. A histopathological
22 evaluation of vaginal tissue was conducted in 5 animals **[assumed 5/group]**. In 44–112 male
23 offspring/group, anogenital distance was measured on PND 3, 15, and 21 and days of testicular descent
24 and preputial separation were recorded. Males were killed on PND 70 (n = 20/group) or 170 (n = 17–
25 20/group). Blood LH and testosterone levels were measured in 14–20 animals/group/time period. Sperm
26 and spermatid numbers and sperm production and transit rates were determined in all offspring.
27 Histopathological evaluation of the testis was conducted in 2 animals **[assumed/group]**. Body,
28 reproductive organ, and liver weights were measured in all male and female offspring killed. Data from
29 female rats were analyzed by ANOVA with post hoc Dunnett test or Fisher test. Data from male rats were
30 analyzed by ANOVA and Dunnett test.

31
32
33 Pup body weights at birth were unaffected in the bisphenol A group, but on PND 22, pup body weights
34 were lower **[by 28%]** in the low-dose group than in the control group. Study authors noted that the mean
35 litter size in the low-dose group was larger by 2.6 pups than in the control group. Effects attaining
36 statistical significance in female pups are summarized in Table 67. Vaginal opening was delayed in the
37 low-dose group and accelerated in the high-dose group. When estrous cyclicity data were evaluated
38 according to total number of cycles, there was an increase in estrous phases lasting more than 1 day and
39 prolongation of the cycle length in the high-dose group. Evaluation of estrous cycles by individual rat
40 indicated a decrease in the percentage of low-dose females with 3 consecutive 1-day estrus phases. The
41 only terminal body and organ weight effects occurred in the low-dose group and included decreased
42 absolute liver weight in females killed in estrus and decreased body and uterus weights in females killed
43 in diestrus or in estrus. There were no effects on relative organ weights. Histological observations in
44 vaginal tissue of bisphenol A-exposed rats included less pronounced cornification during estrus and more
45 pronounced mucification during diestrus, with magnitude of effect greater in the low- than the high-dose
46 group. Observations in the animals exposed to ethinyl estradiol included decreased pup birth weight,
47 delayed vaginal opening, near-persistent estrus, decreased absolute and relative uterus weights, and
48 changes in vaginal histology similar to those described for the low-dose bisphenol A group.

49
50 Statistically significant effects observed in male offspring are summarized in Table 67. Decreased
51 anogenital distances was observed in the bisphenol A groups during all three time periods, but the effect

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1 remained statistically significant only in the high-dose group when normalized for body weight.
 2 Testicular descent and preputial separation were delayed in the low-dose group. Variable effects on
 3 absolute and relative organ weights at both time periods were observed, as indicated in Table 67. Organ
 4 weight effects that remained significant following adjustment for body weight included increased prostate
 5 weight in the high-dose group on PND 70 and increased testicular and epididymal weights in the low-
 6 dose group on PND 170. Variable effects on sperm endpoints such as spermatid and sperm counts, daily
 7 sperm production, and sperm transit time are summarized in Table 67. There was no effect on sperm
 8 morphology. Blood testosterone level was decreased in the high-dose group on PND 70, and blood LH
 9 level was increased in the high-dose group on PND 170. Testicular histopathology observations in the
 10 low-dose group on PND 70 included cellular debris in lumens, pyknotic nuclei in spermatids, and
 11 apoptotic debris in the region of the spermatogonia and primary spermatocyte. In testes of 70-day-old
 12 animals of the high-dose group, there were cental necrotic masses, low numbers of meiotic figures in
 13 spermatocytes, and low spermatozoa numbers. On PND 170, observations in testes from the low-dose
 14 group included low spermatozoa numbers, a thin layer of spermatocyte meiotic figures, and apoptotic
 15 debris in region of spermatids. Low spermatocyte meiotic figures were the only testicular observation in
 16 the high-group on PND 170. Effects observed in the ethinyl estradiol group included increased anogenital
 17 distance, delayed testicular descent, accelerated preputial separation, decreased testis and prostate
 18 weights, decreased sperm counts and production, increased LH levels, increased testosterone levels on
 19 PND 170, apoptotic debris, and/or low sperm numbers in testes.

20
 21 Study authors concluded that prenatal exposure to bisphenol A disrupts the reproductive systems of both
 22 male and female rats and that the effects do not occur according to a classic dose-response curve, which is
 23 generally observed in toxicology studies.
 24

25 **Table 67. Reproductive Effects in Rats Exposed to Bisphenol A During Prenatal Development**

Endpoint	Effects at each dose	
	0.1 mg/kg bw/day	50 mg/kg bw/day
Females		
Age of vaginal opening	↑5.7 days	↓1.9 days
Cycles with estrus phase > 1 day. %	↔	↑
Total 4-day cycles, %	↔	↓23%
No. with 3 consecutive 1-day estrus phase	↓	↔
No. with 3 consecutive 4-day cycles	↔	↔
Body weight of rats killed in diestrus	↓15%	↔
Absolute uterine weight of rats killed in diestrus	↓23%	↔
Body weight of rats killed in estrus	↓8%	↔
Liver weight of rats killed in estrus	↓12%	↔
Absolute uterine weights of rats killed in estrus	↓19%	↔
Males		
Anogenital distance on PND 3	↔	↓14%
Anogenital distance/body weight on PND 3	↔	↓14%
Anogenital distance on PND 15	↓12%	↓35%
Anogenital distance/body weight on PND 15	↔	↓33%
Anogenital distance on PND 21	↓8%	↓13%
Anogenital distance/body weight on PND 21	↔	↓12%
Age at testes descent	↑0.9 days	↔
Age at preputial separation	↑5.4 days	↔
Absolute organ weight, PND 70		
Paired testes	↓6%	↔
Paired epididymides	↓7%	↑7%
Prostate	↔	↑27%

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Endpoint	Effects at each dose	
	0.1 mg/kg bw/day	50 mg/kg bw/day
Seminal vesicle	↓13%	↔
Relative (to body weight) organ weight		
Prostate, PND 70	↔	↑17%
Paired testes, PND 170	↑10%	↔
Paired epididymides, PND 170	↑11%	↔
Spermatid number, PND 70	↔	↓15%
Sperm counts, PND 70	↑21%	↑16%
Daily sperm production, PND 70	↔	↓14%
Sperm transit rate, PND 70	↑23%	↑33%
Spermatid number, PND 170	↓19%	↔
Sperm count, PND 170	↑18	↑16%
Daily sperm production, PND 170	↓19%	↔
Sperm transit rate, PND 170	↑57%	↔
Blood testosterone level, PND 70	↔	↑74%
Blood LH level, PND 170	↔	↑31%

↑,↓ Statistically significant increase, decrease compared to controls; ↔ no statistically significant effect compared to controls.

Source: Talsness et al. (277).

1
2 **Strengths/Weaknesses:** Strengths are the postnatal evaluation of various endpoints to “pup” adulthood
3 and that the concentration of the dosing solutions was verified. Based on the description of numbers of
4 pups contributing to various endpoints, however, the authors do not appear to have used the litter as the
5 unit of analysis. These inflated numbers subjected to analysis complicate the interpretation of findings,
6 especially for PND 1–21 measures. A weakness also is that only 2 dose levels were examined. The
7 vaginal opening data for the controls were outside the normal range for Sprague Dawley rats, and a delay
8 in vaginal opening would be expected with an estrogen. It is unclear how the estrous cycle data were
9 analyzed. The F₁ data were not analyzed correctly. Data may be suggestive of developmental disruptions
10 at both doses, but the magnitudes are likely unreliable, and the authors’ statements about dose-response
11 peculiarities must be viewed with caution until more complete dose-response assessments are published.

12
13 **Utility (Adequacy) for CERHR Evaluation Process:** This study is not adequate for the evaluation
14 process.

15
16 **Tinwell et al. (278),** support not indicated, examined the effects of in utero exposure to bisphenol A on
17 sexual development of male rats. The study attempted to duplicate findings from Chahoud and colleagues
18 that were reported in several abstracts and as a full report (277). Sprague Dawley and Wistar-derived
19 Alderley Park rats were housed in plastic-bottomed cages containing sawdust and shredded paper
20 bedding. Rats were assigned to groups based on body weights and 7/group/strain were gavaged on GD 6–
21 21 with bisphenol A (99% purity) at 0 (arachis oil vehicle), 0.020, 0.100, or 50 mg/kg bw/day. A positive
22 control group initially received 200 µg/kg bw/day ethinyl estradiol, but the dose was reduced to 100
23 µg/kg bw/day between GD 11 and 14 due to maternal toxicity. Dosing solution concentrations and
24 stability were verified. Dams were fed RM3 breeding diet (18.5% soybean protein; Special Diet Services,
25 Ltd.) during gestation and lactation. At birth, pups were counted, sexed, and weighed. Anogenital distance
26 was measured 24 hours following birth (PND 1). On PND 5, pups were culled to 8/litter, with equal
27 numbers of males and females when possible. On PND 23, rats were weighed and housed according to
28 sex. Following weaning, pups were fed RM1 feed (6.5% soybean protein). Pups were weighed throughout
29 the post-lactation period. Ages at preputial separation, vaginal opening, and first estrus were assessed.
30 Males were killed on PND 90–91 and females on PND 98. Liver and reproductive organs were weighed.
31 Daily sperm production was determined. Data were analyzed using the litter and grouped individuals as

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1 the statistical unit. **[Litter values are discussed below.]** Data were analyzed by ANOVA, ANCOVA, and
2 Dunnett test.

3
4 The only significant effect observed in female rats exposed to bisphenol A was a 1.6-day delay in vaginal
5 opening in Alderley-Park rats of the high-dose group. The study authors stated that effect on vaginal
6 opening was correlated with body weight. **[Data were not shown by study authors.]** In Alderley Park
7 males of the high-dose group, significant reductions were observed for total sperm count/testis **[12%**
8 **lower than controls]**, sperm count/g testis **[10% reduction]**, daily sperm count/testis **[12% reduction]**,
9 and daily sperm count/g testis **[10% reduction]**. Benchmark doses for the endpoints with statistically
10 significant changes are shown in Table 68. In both strains, bisphenol A treatment had no effect on litter
11 size, sex ratio, birth weight, anogenital distance, first day of estrus, or age of preputial separation. There
12 were no significant effects on weights of liver, ovary, cervix, uterus, vagina, testis, epididymis, seminal
13 vesicle, or prostate. Rats treated with ethinyl estradiol also experienced decreased sperm counts, in
14 addition to decreased weights of male reproductive organs and advanced age of vaginal opening. Several
15 findings observed by Chahoud and colleagues (277) were not duplicated in this study including: reduced
16 anogenital distance; altered age of sexual maturation in males and females; variable changes in male
17 reproductive organ weight, including prostate weight; and reduced sperm production at low doses. Study
18 authors concluded that this study failed to confirm low-dose endocrine effects.

19
20 **Table 68. Benchmark Doses for Rat Reproductive Organ Endpoints**
21 **Affected by Prenatal Bisphenol A.**

Endpoint	Benchmark dose, mg/kg bw/day			
	BMD ₁₀	BMDL ₁₀	BMD _{1SD}	BMDL _{1SD}
Delayed vaginal opening	68	51	35	16
Sperm count/testis	55	30	57	31
Sperm count/g testis	81	41	68	34
Daily sperm count/testis	56	31	59	31
Daily sperm count/g testis	83	42	70	34

Calculated from data in Tinwell et al. (278).

22
23 **Strengths/Weaknesses:** Strengths of this study are the range and appropriateness of selected measures,
24 the utilization of 4 dose levels, the comparison between 2 strains of rat, the verification of dosing
25 solutions, and the use of ethinyl estradiol, which produced expected responses. An unfortunate weakness,
26 however, is the small sample size of 7 dams/strain/group. Nevertheless, data were appropriately analyzed
27 with the litter as the experimental unit so numbers were not inflated in the analyses, and significance
28 judgments were apparently based on 7/group. Modest effects were noted in male and female offspring in
29 the 50 mg/kg exposure group. While effects on the lowest doses in this study were not seen, it is
30 important to recognize the effects seen at 50 mg/kg bw/day (the high dose in this study) dosing on GD 6–
31 21.

32
33 **Utility (Adequacy) for CERHR Evaluation Process:** This study is adequate for the evaluation process.

34
35 **Schönfelder et al. (279)**, supported by the German Federal Ministry for Education and Research,
36 examined the effects of prenatal bisphenol A exposure on the rat vagina. Sprague Dawley rats were
37 gavaged on GD 6–21 with bisphenol A at 0 (2% corn starch vehicle), 0.1, or 50 mg/kg bw/day. A positive
38 control group was treated with 0.2 mg/kg bw/day 17 β -estradiol in a peanut oil vehicle. **[No information**
39 **was provided on the number of dams treated, the day of vaginal plug, purity of bisphenol A, or the**
40 **type of chow, bedding, and caging materials used.]** At 3 months of age, estrous cyclicity was evaluated
41 for 3 weeks in 42 female offspring of the control group, 21 offspring of the 0.1 mg/kg bw/day group, 18
42 offspring of the 50 mg/kg bw/day group, and 24 offspring of the 17 β -estradiol group. **[The number of**
43 **litters represented was not stated.]** At 4 months of age, female offspring were killed in either estrus or

3.0 Developmental Toxicity

1 diestrus. Vaginas were fixed in Bouin solution and a histopathological evaluation was conducted. Western
2 blot analyses were conducted to measure expression of ER α and ER β . **[No information was provided on
3 the number of animals examined, and it does not appear that statistical evaluations were
4 conducted.]**
5

6 Qualitative descriptions of vaginal histopathology changes and ER expression were provided by the study
7 authors. Low-dose animals killed during the estrous stage lacked keratinization of the surface epithelium
8 and demonstrated reduced thickness of the total epithelium. Similar but less pronounced effects were
9 observed in rats of the high-dose bisphenol A group. Vaginal findings were similar in the positive control
10 group, and slight desquamation of the superficial layers was also observed. There were no differences in
11 vaginal histopathology findings in rats killed during the diestrus stage. No ER β was observed in vaginas
12 of rats from any treatment group. Full-length ER α expression was not observed in either bisphenol A
13 group during estrus, but ER α in the bisphenol A-exposed groups did not differ from the control group
14 during the diestrus stage. ER α in vaginas obtained from the positive control group was either reduced or
15 was not detected. The study authors concluded that altered vaginal morphology following bisphenol A
16 treatment appears to be due to down-regulation of ER α .
17

18 Strengths/Weaknesses: Vaginal histopathology of female offspring is of interest but the quality of the
19 study cannot be judged due to unclear methodology. Uncertainty about the numbers of animals (7 or 8
20 dams may have been used in each group, but group size is uncertain) and the number of offspring
21 examined render this study of marginal value.
22

23 Utility (Adequacy) of the CERHR Evaluation Process: This study is inadequate for the evaluation
24 process.
25

26 **Schönfelder et al. (280)**, supported by the German Federal Ministry for Environmental Protection and
27 Radiation Security, examined the effects of prenatal bisphenol A exposure on the rat uterus. **[No
28 information was provided about composition of feed, caging, or bedding.]** Sprague Dawley rats
29 **[number treated not specified]** were gavaged with bisphenol A **[purity not reported]** at 0 (2% corn
30 starch vehicle), 0.1, or 50 mg/kg bw/day on GD 6–21. The high bisphenol A dose was selected because it
31 was reported to be the no observed effect level (NOEL) recommended by the Society of the Plastics
32 Industry. A positive control group was gavaged with 0.2 mg/kg bw/day ethinyl estradiol on GD 6–21.
33 Estrous cyclicity was examined for 3 weeks in 6 female offspring/group beginning at 3 months of age.
34 Six female offspring/group were killed at 4 months of age on the day of estrus. Body and reproductive
35 organ weights were measured. Uteri were fixed in methacarn solution and sectioned. Examinations of
36 uterine morphology were conducted. Immunohistochemistry techniques were used to detect ER α and ER β
37 in the uterus, and results were verified by Western blot. Data were analyzed by Mann-Whitney test.
38 Statistically significant findings are summarized in Table 69. Effects observed at both dose levels were
39 increased epithelial cell nuclei, epithelial nuclei with condensed chromatin, and epithelial cells with
40 cavities and reduced ER β -positive cells in uterine tissue. Additional effects observed only at the high dose
41 included decreased thickness of luminal epithelium and increased ER α -positive cells in the epithelium.
42 Similar findings were observed following treatment with ethinyl estradiol. The study authors concluded
43 that prenatal bisphenol A exposure causes uterine effects in rat offspring.
44

1 **Table 69. Uterine Effects in Rats Exposed to Bisphenol A During Prenatal Development**

Endpoint ^a	Dose, mg/kg bw/day	
	0.1	50
Thickness of luminal epithelium	↔	↓38%
Epithelial nuclei ^b	↑67%	↑89%
Epithelial nuclei with condensed chromatin	↑2.9-fold	↑3.2-fold
Epithelial cells with cavities	↑2.3-fold	↑87%
ER α positive cells in epithelium	↔	↑58%
ER β -positive cells in uterine tissue	↓88%	↓88%

^aValues were estimated by CERHR from a graph.

^bIt is unclear if authors were referring to numbers of nuclei.

2
3 **Strengths/Weaknesses:** A strength is the examination of effects on uterine indices in female offspring,
4 however, this strength is overwhelmed by the weakness inherent in the data being based on 6
5 females/group, which is too few animals to reach a conclusion with certainty. There are also uncertainties
6 about the number of litters examined.

7
8 **Utility (Adequacy) for CERHR Evaluation Process:** This study is inadequate for the evaluation
9 process.

10
11 **Wistuba et al. (281)**, supported by the German Federal Ministry of Education and Science, examined the
12 effects of prenatal exposure on testicular histology and sperm endpoints in rats. **[No information was**
13 **provided about chow, bedding, or caging.]** Sprague Dawley rats were gavaged with 0 (2% corn starch
14 suspension vehicle), 0.1, or 50 mg/kg bw/day bisphenol A **[purity not reported]** on GD 6–21 (GD 0 =
15 day of sperm detection). A third group was treated with 0.02 mg/kg bw/day ethinyl estradiol. The high
16 dose was said to correspond to the current accepted no observed adverse effect level (NOAEL) and the
17 lower dose was selected to determine if effects occurred at lower doses. It appears that the number of
18 dams treated was 2 in the control group, 4 in the low-dose group, 1 in the high-dose group, and 4 in the
19 ethinyl estradiol group. Litters were weighed during the lactation period. Pups were weaned on PND 22
20 **[day of birth not defined]**. Male offspring were killed between the ages of ~9 and 12 months. The
21 number of males killed was 5 from 2 litters in the control group, 15 from 4 litters in the low-dose group, 5
22 from 1 litter in the high-dose group, and 10 from 4 litters in the ethinyl estradiol group. Testes were fixed
23 in Bouin solution, and Sertoli cells were counted. Spermatogenesis was evaluated by examining germinal
24 epithelia for cell death and distribution of various cell populations. Data were analyzed by ANOVA.

25
26 Examination of tubule cross sections revealed qualitatively normal spermatogenesis in the bisphenol A
27 groups. A comparison of Sertoli cell numbers in littermates revealed high variability (20–27%) in the 0.1
28 mg/kg bw/day group. A comparison of Sertoli cell numbers in the 4 litters from the 0.1 mg/kg bw/day
29 group revealed almost identical results between litters. Sertoli cell numbers/organ were significantly
30 increased by 19.4% in the low-dose group and 19% in the high-dose group. Bisphenol A had no
31 significant effect on Sertoli cell numbers/g testis weight. The opposite situation occurred in the ethinyl
32 estradiol group, with no significant effects on Sertoli cell numbers/organ but a significant increase in
33 Sertoli cell numbers/g testis weight. Testis weight was not affected by bisphenol A treatment but was
34 decreased in the ethinyl estradiol group. The study authors concluded that the study does not support the
35 hypothesis of disruption of the male reproductive system by bisphenol A exposure.

36
37 **Strengths/Weaknesses:** The conceptual strength is the focus on the male reproductive tract/function.
38 However, a weakness is that there were too few animals to provide reliable data.

39
40 **Utility (adequacy) for the CERHR Evaluation Process:** This study is inadequate by itself based on
41 insufficient sample size. It might make a useful contribution when considered with other studies.

3.0 Developmental Toxicity

1
2 **Thuillier et al. (282)**, supported by National Institute of Environmental Health Sciences (NIEHS),
3 examined a possible role for the platelet-derived growth factor system in estrogenic effects induced by
4 bisphenol A in rats exposed during gestation. The effects of other compounds such as genistein and
5 coumestrol were also examined but will not be discussed here. Pregnant Sprague Dawley rats were
6 gavaged with bisphenol A at 0 (corn oil vehicle) or 0.1, 1, 10, or 200 mg/kg bw/day from GD 14 through
7 birth (PND 0). Additional rats were sc injected with diethylstilbestrol at 0.01–2 µg/kg bw/day during the
8 same period. **[No information was provided about number of rats treated, purity of bisphenol A,
9 feed, or materials used in bedding and caging.]** Male offspring were killed on GD 21 or PND 3 and
10 testes were collected. Expression of mRNA or protein for platelet-derived growth factor receptor- α and
11 platelet-derived growth factor receptor- β were determined in testes using RT-PCR, in situ hybridization,
12 or immunohistochemistry. Statistical analyses included unpaired *t*-test with Welch correction.
13

14 Expression of mRNA for platelet-derived growth factor receptor- α and - β was significantly increased at
15 bisphenol A doses ≥ 1 mg/kg bw/day in testes from 3-day-old rats. All other experiments with bisphenol A
16 were conducted with a single dose of 200 mg/kg bw/day. In situ hybridization examination of testes from
17 3-day-old rats from the bisphenol A group revealed an increase in expression of platelet-derived growth
18 factor receptor- α mRNA in testicular interstitium and platelet-derived growth factor receptor- β mRNA in
19 interstitium and seminiferous cords. Exposure to bisphenol A resulted in slightly increased platelet-
20 derived growth factor receptor- α protein expression and strong expression of platelet-derived growth
21 factor receptor- β in gonocytes from 3-day old rat testes. Immunolocalization studies in testes from 21-
22 day-old fetuses revealed that exposure to 200 mg/kg bw/day bisphenol A did not affect expression of
23 platelet-derived growth factor receptor- α protein in gonocytes, but platelet-derived growth factor
24 receptor- β protein appeared to be increased in gonocytes and Sertoli cells. Diethylstilbestrol tended to
25 have a biphasic effect with increased expression of platelet-derived growth factor receptor- α and - β
26 mRNA in 3-day-old rat testis at low doses and decreased expression at the high dose. Treatment with 1
27 µg/kg bw/day diethylstilbestrol decreased mRNA expression of platelet-derived growth factor receptor- α
28 in interstitium and increased platelet-derived growth factor receptor- β mRNA expression in seminiferous
29 cords. Immunoreactivity for platelet-derived growth factor receptor- α protein was maintained but there
30 was a minimal level of platelet-derived growth factor receptor- β protein expression in 3-day-old rat testes
31 following exposure to 1 µg/kg bw/day diethylstilbestrol. In testes obtained from 21-day-old fetuses,
32 expression of platelet-derived growth factor receptor- α protein was decreased in Sertoli and interstitial
33 cells and expression of platelet-derived growth factor receptor- β protein was apparently increased
34 following exposure to diethylstilbestrol. The study authors concluded that the platelet-derived growth
35 factor receptor pathway may be a target for estrogens in the testis, but the findings do not exclude the
36 possibility that effects may have occurred through an ER-independent mechanism.
37

38 **Strengths/Weaknesses:** Endpoints are a strength, but inadequate methodological detail precludes any
39 informed judgment of study quality.
40

41 **Utility (Adequacy) for CERHR Evaluation Process:** This study is inadequate for the evaluation
42 process, based on insufficient methodologic details.
43

44 **Wang et al. (283)**, supported by NIEHS, examined the effects of prenatal bisphenol A exposure on
45 expression of ER-associated proteins in rat testis. The effects of genistein and coumestrol were also
46 examined but will not be discussed here. Pregnant Sprague Dawley rats **[apparently 3/group]** were
47 gavaged with corn oil vehicle or bisphenol A at 0.1–200 mg/kg bw/day from GD 14 (14 days post-
48 coitum) through birth. Additional rats were sc injected with 0.01–2 µg/kg bw/day diethylstilbestrol during
49 the same time period. **[No information was provided about feed, caging and bedding material, or
50 compound purity.]** Male offspring from 3 independent litters were killed on GD 21, PND 3, or PND 21.
51 Western blot, RT-PCR, and immunohistochemistry techniques were used to measure expression of

3.0 Developmental Toxicity

1 protein or mRNA for *Hsp90*, *Hsp90α*, *p23*, *CYP40*, *Hsp70*, and/or *ERβ*. Spermatogonia were quantitated
2 in PND 21 rat testis. Data were analyzed by unpaired *t*-test

3
4 In testes from 3-day-old rats, RT-PCR revealed significant increases in mRNA for *hsp90* at bisphenol A
5 dose levels of 10 and 200 mg/kg bw/day, and significant decreases in expression of *CYP40* at 200 mg/kg
6 bw/day and *p23* at 1 mg/kg bw/day. In situ hybridization analyses in 3-day-old rat testes revealed that
7 bisphenol A tended to increase expression of *hsp90* throughout the testis, with patterns indicating
8 increased expression in gonocytes and interstitial Leydig cells. Examination of protein in testes from 3-
9 day old rats exposed to 200 mg/kg bw/day bisphenol A revealed significantly increased levels of hsp90
10 and hsp70, but no effect on levels of CYP40, p23, or ERβ. Immunohistochemistry revealed that hsp90
11 protein in testes from 3-day-old rats was most increased in gonocytes and less so in interstitium following
12 exposure to 200 mg/kg bw/day bisphenol A. Use of a probe specific for hsp90α protein revealed that
13 increased protein expression of hsp90 was due in a large part to the hsp90α isoform. Examination of
14 testes from GD 21 fetuses and PND 21 pups revealed that the amount of hsp90 protein in the bisphenol A
15 treatment group was similar to that observed on PND 3 but that the amount of protein did not differ from
16 controls on PND 21. In 21 day-old rats from the bisphenol A group, the number of spermatogonia/tubule
17 was significantly higher by ~2-fold compared to the control group. **[It is not clear which bisphenol A**
18 **dose induced an increase in spermatogonia, but it was most likely 200 mg/kg bw/day, because that**
19 **dose appeared to be used in all studies not examining dose-response relationships.]** Effects following
20 diethylstilbestrol exposure included increased expression of *hsp90* mRNA at 1.0 μg/kg bw/day and
21 decreased *CYP40* mRNA expression at 0.01 and 1 μg/kg bw/day, but no effect on protein levels of those
22 compounds was reported in testes from 3-day-old rats. The number of spermatogonia/tubule was also
23 increased after prenatal exposure to diethylstilbestrol. The study authors concluded that prenatal exposure
24 to bisphenol A affects *hsp90* expression in gonocytes of rats, and because hsp90 interacts with several
25 signaling molecules, changes in its expression could affect gonocyte development.

26
27 **Strengths/Weaknesses:** This study was generally well conceived, but the small sample size suggests it
28 presents pilot data only. A full study is needed to provide reliable data.

29
30 **Utility (Adequacy) for CERHR Evaluation Process:** This study is inadequate by itself based on
31 insufficient sample size. It might make a useful contribution when considered with other studies.

3.2.1.3 Neurodevelopmental endpoints

34 **Funabashi et al. (284)**, supported in part by Yokohama City University, examined the effects of
35 bisphenol A on the numbers of corticotropin-releasing hormone neurons in the preoptic area and bed
36 nucleus of the stria terminalis of rats exposed during development. **[No information was provided about**
37 **chow or composition of bedding and caging.]** Pregnant Wistar rats (n = 8–11/treatment group) were
38 given drinking water containing the 0.1% ethanol vehicle or 10 mg/L bisphenol A **[purity not reported]**
39 until their offspring were weaned at 3 weeks of age. **[It is implied but not stated that exposure**
40 **occurred during the entire gestation period.]** Bisphenol A intake was estimated by study authors at 2.5
41 mg/kg bw/day. Male and female offspring (n = 8–11/group) were killed at 4–7 months of age, and
42 immunocytochemistry techniques were used to determine the number of corticotropin-releasing hormone
43 neurons in brain. Female rats were killed during proestrus. **[Although the number of litters represented**
44 **in each group was not specified, the number of rats examined suggests that 1 rat/sex/litter was**
45 **examined.]** Histological slides of brain were evaluated by an investigator blinded to treatment conditions.
46 Two series of experiments were conducted, and data from both experiments were combined. Data were
47 analyzed by ANOVA followed by Fisher protected least significant difference post-hoc test. In the control
48 group, females had more corticotropin-releasing hormone neurons in the preoptic area and anterior and
49 posterior bed nucleus of the stria terminalis than males. Bisphenol A treatment did not change the number
50 of corticotropin-releasing hormone neurons in the preoptic areas of males. A loss in sex difference

3.0 Developmental Toxicity

1 occurred in the anterior and posterior bed nuclei of the stria terminalis following bisphenol A treatment
2 because differences in numbers of corticotropin-releasing hormone neurons between males and females
3 were no longer evident. It appears that bisphenol A treatment increased the number of corticotropin-
4 releasing hormone neurons in males and decreased the number in females. The study authors concluded
5 that exposure to bisphenol A during gestation and lactation results in a loss of sex difference in
6 corticotropin-releasing hormone neurons in the bed nucleus of the stria terminalis but not in the preoptic
7 area.

8
9 **Strengths/Weaknesses:** This study was appropriately designed to examine effects on the development of
10 brain areas known to be influenced by hormonal levels. Strengths include the relevance and subtleties of
11 the endpoints measured; weaknesses include uncertainties about the numbers of animals examined and
12 the duration of the dosing period. The results suggest a disruption of the normal pattern of sexually
13 dimorphic neurons, a result of critical importance to concerns about disruptions relevant to reproductive
14 function and sexually dimorphic behaviors. While the sample was small, the design and statistics appear
15 to be appropriate. It is a weakness that the control for litter effects was not clear.

16
17 **Utility (adequacy) for CERHR Evaluation Process:** This study is adequate for inclusion in the
18 evaluation process, although utility is decreased due to the uncertain nature of the effect.

19
20 **Fujimoto et al. (285)**, supported by the Japanese Ministry of Education, Culture, Sports, Science, and
21 Technology, examined the effect of prenatal bisphenol A exposure on sexual differentiation of
22 neurobehavioral development in rats. Wistar rats were fed CE-2 feed (CLEA, Japan). **[Caging and
23 bedding materials were not described.]** From GD 13 (day of vaginal sperm not defined) to the day of
24 birth (PND 0), 6 rats/group were given tap water containing bisphenol A **[purity not reported]** at 0 or
25 0.1 ppm. The study authors estimated the bisphenol A dose at 0.015 mg/kg bw/day. On PND 1, pups were
26 weighed and litters were culled to 4 pups/sex. Pups were weaned on PND 21. Neurobehavioral
27 evaluations conducted in 20–24 offspring/sex/group at 6–9 weeks of age included open-field, elevated
28 plus maze, passive avoidance, and forced swimming tests. Statistical analyses included ANOVA, Fisher
29 protected least significant difference test, and Mann-Whitney *U* test.

30
31 The results are summarized in Table 70. In the control group, rearing frequency and duration were
32 significantly higher in females than males, but there were no sex-related differences in rearing frequency
33 or duration in the bisphenol A group. Bisphenol A exposure caused an increase in rearing duration in
34 males when compared to males from the control group. In the forced swim test, females in the control
35 group struggled more than males but no sex-related differences in struggling were observed in the
36 bisphenol A group. The duration of immobility in the swimming test was longer in males from the
37 bisphenol A compared to males from the control group. Immobility was described as non-significantly
38 increased in females exposed to bisphenol A compared to control females. Bisphenol A exposure had no
39 effect on performance in passive avoidance and elevated plus maze test. The study authors concluded that
40 exposure of male offspring to bisphenol A during the final week of gestation resulted in impaired sexual
41 differentiation in rearing and struggling behaviors and facilitated depression-like behavior.

1 **Table 70. Sexually-Dimorphic Behaviors in Rats Exposed Prenatally to Bisphenol A**

Endpoint	Controls	Bisphenol A-exposed
Open-field		
Rearing frequency	Females 40% higher than males ^a	No sex difference
Rearing duration	Females 68% higher than males ^a	No sex difference; treated males reared ~50% longer than control males ^a
Time in center area	Females 55% higher than males	Females 60% higher than males
Total distance moved	Females 15% higher than males	Females 12% higher than males
Rapid movements	Females 21% higher than males	Females 21% higher than males
Elevated plus maze		
Time in open arms	No sex difference	No sex difference
Open arm entries	No sex difference	No sex difference
Total entries	No sex difference	No sex difference
Passive avoidance latency	Males 32% longer than females ^a	Males 46% longer than females ^a
Forced swim, time		
Struggling	34% longer in females than males ^a	No sex difference
Immobile	No sex difference	No sex difference; treated males were immobile 75% longer than control males
Shaking head	Males 31% longer than females	Males 31% longer than females
Diving	No sex difference	Females 96% longer than males
Moving limbs	No sex difference	No sex difference

^aEstimated from a graph.

Data from Fujimoto et al. (285).

2
3 **Strengths/Weaknesses:** This study utilized a good choice of methods to examine functional disruptions
4 in sexually dimorphic behaviors. Weaknesses include a lack of clarity about the nature of disruption of
5 sexually dimorphic behavior patterns that was indicated in the authors' conclusions, the somewhat small
6 sample size, the use of a single dose level, which was not confirmed, and the lack of clarity of the
7 statistical methods regarding litter. Nevertheless, the strengths (subtlety and appropriateness of the
8 behavioral measures examined) may outweigh the weaknesses.

9
10 **Utility (Adequacy) for CERHR Process:** This paper is adequate for the evaluation process.

11 3.2.2 Rat—parenteral exposure only during pregnancy

12 **Ramos et al. (286)**, supported by the Argentine National Council for Science and Technology, the
13 Argentine National Agency for the Promotion of Science and Technology, and the Ministry of Health,
14 examined the effects of bisphenol A exposure on the rat prostate. Wistar rats were housed in stainless
15 steel cages. [No information was provided about chow or bedding material.] Four dams/group were
16 exposed to bisphenol A [purity not reported] at 0 (DMSO vehicle), 0.025, or 0.250 mg/kg bw/day by sc
17 pump on GD 8–23 (GD 1 = day of vaginal sperm). Pups were weighed and sexed at birth. Litters were
18 culled to 8 pups, with 4/sex when possible. Pups were weaned on PND 22 [day of birth not defined]. On
19 PND 30, pups were injected with bromodeoxyuridine and killed 2 hours later. Ventral prostates were
20 dissected and fixed in 10% neutral buffered formalin. Immunohistochemical techniques were used to
21 measure proteins associated with cell proliferation and cell phenotypes. Morphometric measurements
22 were taken. [It was not clear how many rats/treatment group were examined for each endpoint.
23 Although a statement was made that males from a single dam were evaluated, it was later stated
24 that siblings were excluded from the same experimental group. Therefore it appears that different
25 litters were represented.] Data were analyzed by Kruskal-Wallis ANOVA and Mann-Whitney *U* test.
26

3.0 Developmental Toxicity

1
2 Statistically significant effects observed in the ventral prostates of rats treated with both doses of
3 bisphenol A are summarized in Table 71. In the periductal stroma, the fibroblastic layer was increased,
4 the smooth muscle layer was reduced, and androgen receptor-positive cells were decreased. Prostatic acid
5 phosphatase-positive cells were reduced in epithelial cells. There were no effects on cell proliferation and
6 ER α was not detected. No changes were observed in interductal stromal cells.

7
8 **Table 71. Effects of Bisphenol A on Proliferation and Differentiation Markers in Rat Prostates**

Endpoint	Doses, mg/kg bw/day	
	0.025	0.250
Relative area of vimentin-positive cells (marker of fibroblast cells) in periductal stroma	↑4.2-fold	↑3.7-fold
Relative area of α -smooth muscle actin positive cells (marker of smooth muscle) in periductal stroma	↓39%	↓45%
Area of androgen receptor-positive cells in periductal stroma	↓48%	↓45%
Area of prostatic acid phosphate-positive cells in epithelial cells ^a	↓45%	↓47%

9
10 ↑, ↓ Statistically significant increase, decrease.

11 ^aThe text indicated that the effect was statistically significant, but Table 2 of the study did not identify the effect as being statistically significant.

12 From Ramos et al. (286).

13
14 **Strengths/Weaknesses:** This study has an interesting design with respect to choice of endpoints. Certain design aspects are unclear, but statistical approaches appear “conservative.” The sample size was small and there was considerable uncertainty about numbers and litter effects. The use of neat DMSO is of concern, as this can modify the effects of the solute.

15
16 **Utility (Adequacy) for CERHR Evaluation Process:** This study is considered marginally adequate, with guarded interpretation because of uncertainties about numbers and origins of the examined males and the concerns about the solution vehicle.

17
18
19 **Ramos et al. (287)**, supported by the Argentine Ministry of Health, Argentine National Agency for the Promotion of Science and Technology, and the National University of Litoral, examined the effects of bisphenol A exposure on the prostate and the hypothalamic-pituitary-gonadal axis in Wistar rats. Rats were housed in stainless steel cages and 7–9/group were administered DMSO vehicle or bisphenol A at 0.025 or 0.250 mg/kg bw/day by sc pump on GD 8–23 (GD 1 = day of vaginal sperm). **[No information was provided on purity of bisphenol A, the type of feed used, or composition of bedding.]** After birth, pups were weighed and sexed. Litters were culled to 8 pups with equal numbers of male and female pups when possible. Pups were weaned on PND 22 **[day of birth not defined]**. During prepuberty (PND 15), peripuberty (PND 30), and adulthood (PND 120), 6–8 males/group were injected with bromodeoxyuridine and killed 2 hours later. **[Although a statement was made that males from a single dam were evaluated, it was later stated that siblings were excluded from the same experimental group. Therefore, it appears that different litters were represented.]** Serum was collected for measurement of LH and prolactin by RIA. Immunohistochemistry techniques were used to evaluate markers of cell proliferation, estrogen/androgen receptors, and prostatic cells. Expression of mRNA for ER α and ER β in the preoptic area and medial basal hypothalamus was determined by RT-PCR. Data were analyzed by Kruskal-Wallis 1-way ANOVA using Dunn post-test.

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36 Statistically significant effects are summarized in Table 72. No significant effects were observed for ventral prostate weight. Numerous transient effects were observed in both bisphenol A dose groups. On

3.0 Developmental Toxicity

PND 15, cellular proliferation was increased in the periductal stroma of the prostate, and serum testosterone levels were elevated. On PND 30, the fibroblasts (vimentin-positive cells) in the prostatic periductal stroma was increased and the area of smooth muscle cells (α -smooth muscle actin) was decreased. Also observed on PND 30 was a reduction in androgen-receptor positive stromal cells, a decrease in epithelial cells positive for prostatic acid phosphatase, and an increase in serum prolactin levels. Expression of *ER β* mRNA was increased in the preoptic areas on PND 30 and 120, and the study authors considered the effect to be permanent because it occurred on both days. The study authors concluded that prenatal exposure to environmental concentrations of bisphenol A during gestation results in transient and permanent changes in the male reproductive axis.

Table 72. Effects of Bisphenol A on Proliferation and Differentiation Markers in the Rat Prostate

Endpoint	Dose, mg/kg bw/day	
	0.025	0.250
Incorporation of bromodeoxyuridine by periductal cells, PND 15	↑88%	↑155%
Periductal cells occupied by vimentin, PND 30	↑205%	↑164%
Periductal cells occupied by α -smooth muscle actin, PND 30	↓49%	↓41%
Periductal androgen receptor-positive cells	↓48%	↓45%
Prostatic acid phosphate-positive cells	↓45%	↓45%
Serum prolactin levels, PND 30 ^a	↑300%	↑367%
Serum testosterone level, PND 15 ^a	↑33%	↑25%
Expression of <i>ERβ</i> mRNA in preoptic area, PND 30 and 120 ^a	↑3.5–4-fold	↑3.5–4-fold

↑,↓ Statistically significant increase, decrease.

^aValues estimated by CERHR from a graph.

From Ramos et al. (287).

Strengths/Weaknesses: The design seems reasonable as a means to address the study questions. Weaknesses include a very small n for most of these measures and uncertainty about the litter origin and representation in each necropsy group. Like many of these studies, altered values are given without addressing the normal range of variation or the likely functional significance of the changes.

Utility (Adequacy) for CERHR Evaluation Process: This study is inadequate for the evaluation process, based on the small n for these measures and the uncertainty this creates.

Naciff et al. (246), from The Procter and Gamble Company, examined the effects of prenatal bisphenol A exposure on gene expression and, to a limited extent, development in female rat reproductive organs. Pregnant Sprague Dawley rats were fed Purina 5K96, a casein-based soy- and alfalfa-free diet.

[Composition of caging and bedding materials was not reported.] The rats were randomly assigned to groups (≥ 7 rats/group) sc injected with bisphenol A (~99% purity) in DMSO vehicle at 0, 5, 50, or 400 mg/kg bw/day on GD 11–20 (day of sperm detection = GD 0). Dams were killed on GD 20, and ovaries and uteri were removed from fetuses. In 4 litters/group, 1 female fetus/litter was examined for ovarian and uterine histopathology. In 5 litters/group, ovaries and uteri from at least 5 littermates were pooled for a microarray analysis of gene expression. Changes in gene expression were further quantified using RT-PCR. Data were analyzed by *t*-test, ANOVA, and Jonkheere-Terpstra test. Comparisons of gene expression among estrogenic compounds were made by Wilcoxon-Mann-Whitney and Jonkheere-Terpstra tests. Results of gene expression assays are discussed in Section 2. Vaginal bleeding and early parturition occurred in 1 of 8 dams in the high-dose group. Bisphenol A treatment had no effect on maternal body weight or number of live fetuses/litter, and there were no gross or histopathological effects on ovary or uterus. Prominent nipples and areolas were observed in males and females in the high-dose bisphenol A group **[number of fetuses and litters affected were not reported]**.

3.0 Developmental Toxicity

1 **Strengths/Weaknesses:** The sample sizes are small but adequate for these sorts of analyses. Strengths are
2 that these endpoints appear appropriate; weaknesses are the limited nature of the endpoints and the use of
3 neat DMSO as vehicle. While endocrine disruption may certainly affect reproductive tissue development,
4 of greater concern are potential disruptions in the neural control centers that are programmed in early
5 development for performance at puberty and beyond.

6
7 **Utility (Adequacy) for CERHR Evaluation Process:** This study is adequate but ancillary for the
8 evaluation process.

9
10 **Naciff et al. (288)**, from The Procter and Gamble Company, examined the effect of prenatal bisphenol A
11 exposure on male rat reproductive organ histology and gene expression.
12 Pregnant Sprague Dawley rats were fed Purina 5K96, a casein-based soy- and alfalfa-free diet. Rats were
13 housed in stainless steel cages prior to mating. Rats were randomly assigned to groups (≥ 8 rats/group) and
14 sc injected with bisphenol A (~99% purity) in DMSO at 0, 0.002, 0.02, 0.5, 50, or 400 mg/kg bw/day on
15 GD 11–20 (day of sperm detection = GD 0). Dams were killed on GD 20, and testes and epididymides
16 were removed from fetuses. In 4 litters/dose group, 1 male fetus/litter was examined for testicular
17 histopathology. In 5 litters/group, testes and epididymides from 5 littermates were pooled for a microarray
18 analysis of gene expression. Changes in gene expression were further quantified using RT-PCR. Data
19 were analyzed by *t*-test, ANOVA, and Jonkheere-Terpstra test. Comparisons of gene expression among
20 estrogenic compounds were analyzed by Wilcoxon-Mann-Whitney and Jonkheere-Terpstra tests.
21 Bisphenol A treatment had no effect on maternal body weight or number of live fetuses/litter, and there
22 were no gross or histopathological effects on the testis or epididymis. Prominent nipples/areolas were
23 observed in male and female fetuses from the high-dose group [**numbers of fetuses and litters affected**
24 **were not reported**]. In pooled testis and epididymis samples from the high-dose bisphenol A group,
25 expression of 15 genes was significantly altered in a dose-related manner. When bisphenol A data were
26 pooled with data obtained from ethinyl estradiol and genistein and globally analyzed, there were 50 genes
27 that were significantly altered in the same direction by all 3 compounds. The study authors concluded that
28 transplacental exposure to high doses of bisphenol A alters the expression of certain genes in the testis
29 and epididymis of fetal rats without causing malformations in those organs. The study authors noted that
30 the dose response to bisphenol A was monotonic with no evidence of robust quantifiable responses at low
31 doses.

32
33 **Strengths/Weaknesses:** Strengths of this study are the relevance of the endpoints, the strategy used, and
34 the adequate numbers of animals for gene expression. Weaknesses include the small number of animals
35 used to evaluate histopathology and the use of neat DMSO. The study shows the commonality of gene
36 response to several compounds thought to impact the estrogen response system.

37
38 **Utility (Adequacy) for CERHR Evaluation Process:** This study is adequate for the evaluation process.

39
40 **Saito et al. (289)**, support not indicated, examined the effect of prenatal bisphenol A exposure on
41 testosterone production during adulthood in rats. On GD 12–19 (day of vaginal plug not reported), 2
42 Wistar rats were sc injected with the corn oil vehicle, 4 rats were sc injected with 0.005 mg/day bisphenol
43 A [**purity not indicated**], and 2 rats were injected with 5 μ g/day 17 β -estradiol. [**Assuming a pregnant**
44 **Wistar rat weights ~0.33 kg, 0.005 mg/day would be equivalent to 0.015 mg/kg bw/day bisphenol A.**]
45 Other materials found in dental composites were also evaluated but will not be discussed. During the
46 lactation period, rats were housed in polypropylene cages with synthetic bedding. [**No information was**
47 **provided on feed.**] Offspring were housed separately at 3 weeks of age and killed at 13 weeks of age.
48 Body and testis weights were measured in all male offspring (22 in the bisphenol A group, 11 in the
49 vehicle control group, and 5 in the 17 β -estradiol group). Plasma testosterone level was measured by RIA,
50 and plasma cholesterol level was measured using a kit. Activities of testicular enzymes involved in the
51 production of testosterone from progesterone were also examined in an in vitro assay in which testicular

3.0 Developmental Toxicity

1 microsomes were incubated with ^{14}C -progesterone and ^{14}C - Δ^4 -androstendione for 20 minutes. Data were
2 analyzed using unspecified post hoc tests.

3
4 Bisphenol A exposure had no effect on pup sex ratio. No effects on body weight or absolute testicular
5 weight were observed in the bisphenol A group at 13 weeks of age. However, relative (to body weight)
6 testicular weight was lower [**by 6%**] in rats of the bisphenol A compared to the control group. Also
7 observed in the bisphenol A group was a reduction in plasma testosterone level [**by ~28%**]. No effect was
8 observed on cholesterol level. In the ex vivo study, prenatal bisphenol A exposure increased activities of
9 17α -hydroxysteroid dehydrogenase [**by ~140%**] and 17β -hydroxysteroid dehydrogenase [**by ~70%**].
10 Observations in the 17β -estradiol compared to the control group included decreased numbers of offspring
11 delivered, higher body weight of male offspring at 13 weeks of age, reduced plasma testosterone level,
12 and increased testicular 17α -hydroxysteroid dehydrogenase activity. The study authors concluded that
13 bisphenol A had an estrogenic effect on the testis but did not decrease activities of enzymes involved in
14 testosterone production.

15
16 **Strengths/Weaknesses:** A strength of this study is the examination of testosterone levels at 13 weeks of
17 age. This strength is negated by the sample size, which is too small to draw any firm conclusions.

18
19 **Utility (Adequacy) for CERHR Evaluation Process:** This study is inadequate by itself based on
20 insufficient sample size. It might make a useful contribution when considered with other studies.

21
22 **Murray et al. (290)**, supported by NIH, examined the effect of prenatal bisphenol A exposure on in situ
23 induction of mammary tumors. Wistar-Furth rats were fed Harlan Teklad 2018, which was reported to
24 contain 20 fmol/g estrogen equivalents. Water was supplied in glass bottles. Caging and bedding
25 materials were not reported, but they were stated that to test negative in the E-SCREEN. From GD 9 (GD
26 1 = day of vaginal sperm) through PND 1 [**assumed to be day of birth**], rats received the 50% DMSO
27 vehicle or bisphenol A [**purity not reported**] at 0.0025, 0.025, 0.250, or 1 mg/kg bw/day. Dosing
28 solutions were delivered by implanted [**assumed SC**] osmotic pumps. [**Number of dams treated was not**
29 **reported. Based on a limited amount of information provided on the number of offspring examined,**
30 **it appears that ≤ 6 dams/group were treated.**] Pup viability was assessed on PND 1. On PND 2 pups
31 were sexed and litters were culled to 8 pups. Anogenital distance was measured on PND 4. Litters were
32 weighed during the lactation period. Female offspring were monitored for body weight and vaginal
33 opening in the post weaning period. Female offspring were killed on PND 50 or 95. Mammary glands
34 were collected and whole-mounted or sectioned for histopathological examination. Morphometric
35 analyses were conducted to examine possible presence of preneoplastic lesions. Mammary glands were
36 examined for ER α and Ki-67 protein by an immunohistochemistry technique. Maximal numbers of
37 maternal units were represented in each dose group. One female/litter was included in histological
38 examinations. [**Apparently ≤ 6 offspring/group were examined in histopathological examinations.**
39 **Number of offspring examined for other endpoints was not reported.**] Statistical analyses included
40 ANOVA followed by post hoc tests (Bonferroni or *t*-test) when significant effects were observed by
41 ANOVA.

42
43 Significant findings are summarized in Table 73. Bisphenol A exposure did not affect offspring viability,
44 sex ratio, age at vaginal opening, or female anogenital distance. Anogenital distance was reduced on PND
45 4 in males from the 0.250 mg/kg bw/day group. Percent hyperplastic ducts was increased in all dose
46 groups on PND 50 and in the 0.0025 mg/kg bw/day group on PND 95; the study authors noted that the
47 effect on PND 50 was quantitatively similar in all dose groups (i.e. 3–4-fold increase). Cribriform
48 structures were observed in the 0.25 and 1 mg/kg bw/day groups (Table 73). [**Incidence was not**
49 **reported for the control and lower dose groups.**] The structures were classified as carcinomas-in-situ
50 and were characterized by increased ductal size resulting from luminal epithelium proliferation, enlarged
51 luminal epithelial cells, presence of a nucleolus, variable chromatin pattern, and rounded luminal spaces

3.0 Developmental Toxicity

1 consisting of trabecular rods of cells perpendicularly aligned to the longer duct axis. Numbers of Ki-67-
 2 and ER- α positive cells were increased in aberrant compared to normal tissues, regardless of dose.
 3 **[Results in treated compared to control groups were not reported.]** The study authors concluded that
 4 fetal bisphenol A exposure in rats is sufficient to induce development of preneoplastic and neoplastic
 5 mammary lesions.

6
 7 Table 73. Effects in Offspring of Rats Exposed to Bisphenol A by Osmotic Pump During Gestation

Endpoint	Bisphenol A dose in mg/kg bw/day			
	0.0025	0.025	0.250	1
Male anogenital distance, PND 4	↔	↔	↓12%	↔
% Hyperplastic ducts, PND 50 (~7% incidence in control) ^a	↑ (~24%)	↑ (~20%)	↑ (~25%)	↑ (~20%)
% Hyperplastic ducts, PND 95 (~4% incidence in controls) ^a	↑ (~13%)	↔	↔	↔
Cribriform structures, PND 50 [incidence not reported for controls]	Not reported	Not reported	1 of 4	1 of 4
Cribriform structures, PND 95 [incidence not reported for controls]	Not reported	Not reported	2 of 6	2 of 6

From Murray et al. (290)

↑,↓ Statistically significant increase, decrease compared to controls; ↔ no statistically significant effects compared to controls

^aValues estimated from a graph by CERHR

8
 9 **Strengths/Weaknesses:** To be added

10
 11 **Utility/Adequacy for CERHR Evaluation:** To be added

12
 13 **Durando et al. (291)**, supported by Universidad Nacional del Litoral, Argentine National Agency for the
 14 Promotion of Science and technology, and NIH, examined the effects of prenatal bisphenol A exposure
 15 on susceptibility to mammary tumors in rats. Wistar rats were fed Cooperación (Buenos Aires, Argentina)
 16 and housed in stainless steel cages. **[It was not clear if bedding was used.]** On GD 8–23 (GD 1 = day of
 17 vaginal sperm), 11–14 dams/group were sc dosed by osmotic pump with the DMSO vehicle or 0.025
 18 mg/kg bw/day bisphenol A **[purity not indicated]**. Pups were delivered on GD 23 and weaned on PND
 19 21. It was not indicated if day of birth was considered PND 0 or 1. During the study, body weights and
 20 day of vaginal opening were monitored. Offspring were killed before puberty (PND 30), after puberty
 21 (PND 50), or in adulthood (PND 110 and 180). In mammary gland stroma and epithelium, proliferation
 22 was assessed by BrdU incorporation and apoptotic cells were identified by TUNEL method.
 23 Morphometric analyses were conducted in sectioned mammary glands. Mast cells were identified by
 24 immunostaining for proteinase I. **[The number of offspring killed and evaluated on PND 30, 50, 110,
 25 and 180 was not reported, and it was not indicated if all litters were represented.]** Additional
 26 offspring were examined for responsiveness to chemically-induced mammary preneoplastic or neoplastic
 27 lesions. On PND 50, N-nitroso-N-methylurea was administered to 10–16 offspring from the vehicle
 28 control group at 25 or 50 mg/kg bw and 21 offspring from the bisphenol A group at 25 mg/kg bw. Based
 29 on findings from a pilot study, 25 mg/kg bw was considered a subcarcinogenic dose and 50 mg/kg bw
 30 was considered a positive control. During the study, rats were palpated for tumors. Rats that received 50
 31 mg/kg bw N-nitroso-N-methylurea were killed on PND 180 and rats that received 25 mg/kg bw N-

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1 nitroso-N-methylurea were killed on PND 110 or 180. Whole-mounted mammary glands were examined
2 for tumors. Immunostaining was conducted to identify cytokeratin 8 (an epithelial marker) and p63 (a
3 myoepithelial marker). Data were statistically analyzed using the Mann-Whitney *U* test.

4
5 Bisphenol A exposure did not affect successful pregnancies, dam weight gain, pregnancy duration,
6 number of pups/litter, or percent females/litter. Anogenital distance on PND 1 or 5 and postnatal body
7 weights were unaffected in pups exposed to bisphenol A. Vaginal opening was accelerated in pups from
8 the bisphenol A group (mean 34 days of age compared to 39 days of age in controls). On PND 50, the
9 BrdU/apoptosis ratio was significantly increased and apoptosis was significantly decreased in
10 parenchyma and stroma of bisphenol A-exposed animals; the effects were not observed on PND 30 or
11 110. Significantly increased percentages of hyperplastic ducts, density of stromal nuclei, and numbers of
12 mast cells were observed in the bisphenol A group on PND 110 and 180. Exposure to bisphenol A
13 resulted in formation of a dense stroma layer around mammary epithelial structures and replacement of
14 normal adipose tissue with a fibroblastic stroma. In rats exposed to 25 mg/kg bw N-nitroso-N-methylurea
15 on PND 50, incidence of hyperplastic lesions on PND 180 was significantly higher in the group with
16 prenatal bisphenol A compared to DMSO exposure (mean incidence of 35.5% compared to 15.7% in
17 controls). Although statistical significance was not achieved, exposure to 25 mg/kg bw N-nitroso-N-
18 methylurea resulted in tumors in 2 of 15 mice in the prenatal bisphenol A group and 0 of 10 mice in the
19 prenatal vehicle control group on PND 180. Cytokeratin 8 immunostaining revealed no invasion by
20 stromal epithelial cells. The study authors concluded that rats prenatally exposed to environmentally
21 relevant doses of bisphenol A may have an increased risk of developing mammary tumors.

22
23 **Strengths/Weaknesses:** To be added

24
25 **Utility (Adequacy) for CERHR Evaluation Process:** To be added

26 27 3.2.3 Rat—oral exposure postnatally with or without prenatal exposure

28 29 3.2.3.1 Multigeneration studies

30 **Emm et al. (292)**, supported by the Japanese Ministry of Health and Welfare, examined developmental
31 toxicity endpoints, in a 2-generation rats study described in detail in Section 4.2.3.1. Two generations of
32 rats were gavaged with 0, 0.0002, 0.002, 0.020, or 0.200 mg/kg bw/day bisphenol A prior to and during
33 mating and throughout the gestation and lactation period. These doses were based on previous studies
34 which found effects at 0.002 and 0.020 mg/kg bw/day. There were some non-dose-related and sporadic
35 effects, but the study authors concluded that none of the effects were related to bisphenol A treatment.
36 Bisphenol A exposure did not adversely affect prenatal or postnatal growth or survival, developmental
37 landmarks, anogenital distance, or age of puberty. In adult animals exposed to bisphenol A during
38 development, there was no evidence of adverse effects on reproductive endpoints such as fertility, estrous
39 cyclicity, or sperm counts. Prostate and other male reproductive organ weights were unaffected.

40
41 **Strengths/Weaknesses:** Strengths of this study were the thoroughness of the evaluation, the size of the
42 dose range, the large number of animals, the litter-based analysis, and the verification of the dosing
43 solution. A weakness is the failure to replicate previous findings at 0.002 and 0.020 mg/kg bw/day and
44 the lack of a positive control group, which leaves a persistent questions about the ability of this group of
45 rats to respond.

46
47 **Utility (Adequacy) for CERHR Evaluation Process:** This study is adequate for the evaluation process.

48
49 **Tyl et al. (293)**, supported by The Society of the Plastics Industry, Inc., reported some developmental
50 toxicity effects in a multigeneration bisphenol A study in Sprague Dawley rats that is reported in detail in
51 Section 4.2.3.1. In that study, F₁, F₂, and F₃ rats were exposed to bisphenol A indirectly during gestation

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1 and lactation and directly through feed after weaning. Dietary doses were 0, 0.015, 0.3, 4.5, 75, 750, or
2 7500 ppm, and target intakes were ~0.001, 0.02, 0.30, 5, 50, and 500 mg/kg bw/day. At the 7500 ppm
3 dose there were fewer pups and live pups/litter and body weight gain of pups was lower during the
4 lactation period. Delayed puberty in both males and females of the 7500 ppm group was most likely
5 related to reduced body weights according to the study authors. Bisphenol A exposure during
6 development did not increase the weight of the prostate in adult rats. Although some decreases in
7 epididymal sperm concentration and daily sperm endpoints were each observed in 1 generation of males
8 from the high-dose group, the study authors concluded there were no treatment-related effects on sperm
9 endpoints or reproductive function. The study authors identified an offspring and reproductive NOAEL of
10 750 ppm (~50 mg/kg bw/day). A systemic NOAEL for adult rats was identified at 75 ppm (~5 mg/kg
11 bw/day) by the study authors; therefore, bisphenol A was not considered a selective developmental
12 toxicant.

13
14 **Strengths/Weaknesses:** This study has numerous strengths, including the quality and number of the
15 endpoints evaluated, the number of dose groups and generations examined, and the confirmation of
16 dosing solutions. This study incorporated screening-level endpoints within the context of a
17 multigeneration study. As such, it addresses gross issues but does provide helpful data regarding the
18 NOAEL.

19
20 **Utility (Adequacy) for CERHR Evaluation Process:** This study is adequate for the evaluation process.

21 22 *3.2.3.2 Development of the reproductive or endocrine systems*

23 **Cagen et al. (294)**, support not indicated (but all authors affiliated with industry), conducted a study to
24 examine the effects of prenatal and lactational bisphenol A exposure on reproductive development of rats.
25 The study attempted to replicate findings by Sharpe et al. that appeared in an unpublished meeting
26 abstract. The protocol used by Cagen et al. was the same as that used by Sharpe et al., with the exception
27 that more dose levels were included, group sizes were larger, and a greater number of reproductive
28 endpoints were examined. Animals were fed Certified Rodent Chow #5002. Music was played at a low
29 volume to provide background noise. Female Han-Wistar rats were randomly assigned to groups. For 2
30 weeks prior to mating, during a 2-week mating period, and during the gestation and lactation periods, 28
31 rats/group were given drinking water containing bisphenol A (>99% purity) at 0.01, 0.1, 1.0, or 10 ppm
32 (0.001–0.004, 0.008–0.038, 0.100–0.391, or 0.775–4.022 mg/kg bw/day). Two negative control groups of
33 28 rats each were given undosed drinking water. Because the two control groups were determined to be
34 statistically equivalent, data from the two groups were pooled. A positive control group of 28 rats was
35 given drinking water with diethylstilbestrol at 0.1 ppm (0.006–0.036 mg/kg bw/day). Dosing solutions
36 were prepared weekly, and concentrations were verified. Dams were evaluated for food and water intake,
37 weight gain, and fertility endpoints. Pups were sexed, weighed, and counted at birth. During the postnatal
38 period, pups were evaluated for growth and survival. On PND 4, litters were culled to 8 pups with as
39 many male pups retained as possible. At weaning on PND 22, up to 4 males/litter (86–109 pups/group)
40 were randomly selected to continue in the study until 90 days of age and were individually housed. At
41 necropsy, brain, liver, kidneys, and reproductive organs were weighed, daily sperm production was
42 determined, and testes were examined histologically. Technicians were blinded to treatment group. The
43 litter was considered the experimental unit in statistical analyses. Data were analyzed by Levene test,
44 ANOVA, Dunnett test, rank transformation, and Wilcoxon rank sum test with Bonferroni correction.

45
46 In the bisphenol A groups, there were no significant effects on dam body weight gain or food or water
47 intake. **[Food and water intake data were not shown.]** There were also no effects on dam fertility,
48 mating, gestation index and duration, live litter size, or pup survival and body weight gain during the
49 postnatal period. Male sex ratio was increased in the 0.1 ppm bisphenol A group (56.7% males versus
50 48.4% in control), but the study authors did not consider the effect to be treatment-related due to the lack
51 of a dose response relationship. Dams in the diethylstilbestrol group experienced decreased body weight

3.0 Developmental Toxicity

1 gain and food intake, increased duration of gestation, smaller litter size at birth, and decreased pup
2 survival in the postnatal period.

3
4 In adult offspring from the bisphenol A groups, there were no significant effects on terminal body weight
5 or organ weights including prostate, epididymis, preputial gland, prostate, seminal vesicle, or testis. There
6 were also no significant effects on epididymal sperm concentration, efficiency of sperm production, or
7 daily sperm production. No histopathological alterations were observed in the testis. Reproductive
8 development in male offspring was also unaffected by prenatal exposure to diethylstilbestrol. The study
9 authors noted that the reduced testis weight and sperm production reported by Sharpe et al. was not
10 confirmed in this study and that bisphenol A should not be considered a selective reproductive or
11 developmental toxicant.

12
13 **[The NTP Statistics Subpanel (295) concluded that the statistical methods used by Cagen et al. were**
14 **appropriate. Although the Subpanel agreed with the study author conclusions, 2 matters were**
15 **noted. The first was that a significant ANOVA is not a requirement for Dunnett test. The second**
16 **was that a Bonferroni correction of Wilcoxon-rank sum test was not needed because the authors**
17 **already required significance by ANOVA, which was sufficient.]**

18
19 **Strengths/Weaknesses:** Significant strengths of this study include the large number of dose levels and
20 animals per dose level and the technical care with which the study was performed, as well as the inclusion
21 of a positive control group and two negative controls. The lack of much effect with diethylstilbestrol
22 treatment is a weakness.

23
24 **Utility (Adequacy) for CERHR Evaluation Process:** This study is adequate for the evaluation process,
25 although the lack of much effect with the positive control raises uncertainties about interpretation of the
26 results..

27
28 **Elswick et al. (296)**, from the Chemical Industry Institute of Toxicology [CIIT], examined the effects of
29 sampling design on conclusions made about bisphenol A effects on prostate weight. Statistical
30 evaluations were conducted using data generated in 3 studies in Sprague Dawwley rats performed at CIIT
31 between 1997 and 1999. In those studies, the litter was considered the experimental unit in statistical
32 analyses. Organ weights were analyzed using a nested ANOVA with litter within dose as the random
33 effect. Post hoc tests were conducted when appropriate.

34
35 A manuscript was reported to be in preparation for a low-dose drinking water study but no additional
36 reference information was provided. Dams were given drinking water containing 0, 0.005, 0.05, 0.5, 5, or
37 50 mg/L bisphenol A from GD 2 to PND 21. The study authors estimated bisphenol A intakes at ~0.001–
38 10 mg/kg bw/day. The lowest doses were reported to be similar to human exposure levels. The study was
39 conducted in 2 blocks separated by 4 months. A total of 16 dams/group were exposed, and the overall
40 sample size was ultimately 13–16/group. In the first block, 2 males/litter were most often retained and in
41 the second block, 1 male/litter was retained until 6 months of age. Fresh ventral prostate weights were
42 recorded. Analysis of data from the first study block revealed no treatment-related effects on ventral
43 prostate weight. Within litters, ventral prostate weights were observed to be very variable, with weights
44 sometimes differing by values of 2-fold or more. In the second study block, mean weights in the 0.05, 5,
45 and 50 mg/kg bw/day groups were significantly higher than those of the control group. It was noted that
46 mean prostate weight in the control group from block 2 (0.387 g) was much lower than the mean weight
47 observed in block 1 (0.517 g) and that the standard error in block 2 (0.174 g) was almost two times higher
48 than the standard error in block 1 (0.092 g). When data from the 2 blocks were combined, statistical
49 significance remained. The study authors noted that no historical control database was available at CIIT at
50 the time of the analysis.

3.0 Developmental Toxicity

1 Data from a high-dose bisphenol A gavage study were next examined. Details are available in the study
2 by Kwon et al. (297) [discussed in Section 3.2.3.3], which was in press at the time this report was
3 published. Eight dams/group were gavaged with bisphenol A in corn oil at 0, 3.2, 32, or 320 mg/kg
4 bw/day on GD 11 through PND 20, excluding the day of parturition. All males in each litter were retained
5 until necropsy at 6 months of age. Tissues of interest were collected and weighed. Prostate glands were
6 dissected by 1 experienced prosector who was blinded to treatment conditions. In addition, simulations
7 were conducted to determine conclusions after random selection and analysis of 1, 2, or 3 pups/litter.
8 Simulated data were analyzed by one-way nested ANOVA with litter within dose as the random effect.
9 Simulated results were compared to results where all males within a litter were retained and analyzed. In
10 the analyses of all male pups/litter, large inter-litter variability (14%) was observed for ventral prostate
11 weights and ranges of litter means were reported at 0.505–0.727 g. Statistical significance was not
12 attained for a 23% increase in ventral prostate weight in the 320 mg/kg bw/day group. In simulation
13 analyses for the high-dose group, false positive conclusions were 19% when 1 pup/litter was analyzed and
14 20% when either 2 or 3 pups/litter were analyzed.

15
16 Simulation analyses were also conducted for data on ventral prostate weights obtained from offspring of
17 rats gavaged with 0, 0.5, 5, 50, 100, or 500 mg/kg bw/day dibutyl phthalate on GD 12–21. The study, by
18 Mylchreest et al. (298), was conducted in 2 blocks, with total number of dams targeted at 10 in the 500
19 mg/kg bw/day group and 20 in the other dose groups. In contrast to the original study, which did not
20 include data from animals with missing or incompletely formed reproductive organs, data from all
21 animals were included in the Elswick et al. analyses. Male offspring were necropsied at ~100 days of age.
22 The median intra-litter coefficient of variation for ventral prostate weight was reported at 18% and ranged
23 from 7 to 44%. Ventral prostate weights that differed by more than 2-fold from 1 pup to another were
24 reported in 6 of 19 litters. In the 500 mg/kg bw/day group, ventral prostate weights were significantly
25 lower than in controls in block 1 and in both blocks combined, but not in block 2. In the original study,
26 only marginal significance was obtained when ventral prostate weights were not included for animals
27 with reproductive tract malformations. In simulation analyses, the mean incorrect conclusions in block 1
28 were 44% when 1 pup was analyzed, 8% when 2 pups were analyzed, and <1% when 3 pups were
29 analyzed. In block 2, incorrect conclusions were <1% or 0, regardless of the number of pups analyzed.
30 When blocks were combined, false results following analysis of 1, 2, or 3 pups/litter were 91, 56, and
31 16%. The study authors noted that incorrect results were more likely with *P* values closest to 0.05. The *P*
32 values were 0.0005 for block 1, 0.6685 for block 2, and 0.0038 for block 3. The study authors concluded
33 that sampling strategies can lead to incorrect conclusions.

34
35 **[The NTP Statistics Subpanel (295) concluded that the results and conclusions of their analyses of**
36 **Elswick et al. (296) showed a consistent increase in ventral prostate weight in the 2 replicates.]**

37
38 **Strengths/Weaknesses:** This paper presents an interesting approach to the examination of background
39 variance and litter effects as related to the endpoints under study. These data argue for multiple pup/litter
40 sampling, a characteristics that has been uncommon in this literature. The significant effects noted only in
41 1 block raise the question of a lack of experience or training among the technicians.

42
43 **Utility (Adequacy) for CERHR Evaluation Process:** This study is adequate for the evaluation process,
44 although it does not provide original data.

45
46 **Rubin et al. (217)**, supported by the Tufts Institute of the Environment and NIH, examined the effects of
47 perinatal bisphenol A exposure on estrous cyclicity and LH levels in rats. Uterotropic responses were
48 examined in a second group of rats, and those results are listed in Table 49. Sprague Dawley rats were fed
49 Purina Rodent Chow and provided drinking water in glass bottles. The rats were housed in plastic cages;
50 estrogenicity testing of ethanol extracts indicated that estrogenic compounds did not leach from cages at
51 detectable levels. **[No information was provided about bedding.]** Dams were weighed and randomly

3.0 Developmental Toxicity

1 assigned to treatment groups of 6 animals given drinking water containing bisphenol A [**purity not**
2 **reported**] at 0 (1% ethanol vehicle), 1, or 10 mg/L from GD 6 (plug day not indicated) through the
3 lactation period. Mean bisphenol A doses were estimated by study authors at 0.1 and 1.2 mg/kg bw/day.
4 At weaning, pups were given untreated water. Dams were examined and weighed during the studies.
5 Offspring were sexed on PND 2 and weighed beginning in the postnatal period and continuing through
6 adulthood (n = 40–53/group during the neonatal period and 19–27/sex/group during adulthood).
7 Anogenital distance was examined during the neonatal period. [**It was not clear how many time points**
8 **and animals were examined.**] Genital tracts were examined for gross abnormalities in males killed
9 during the neonatal period, at 3 months, and at 5 months of age and in females killed during the neonatal
10 period, at 8 months, and at 12–16 months of age. [**The total number of animals examined at each time**
11 **period was reported as 12–34, but it is not known how many/dose group were examined.**] Animals
12 were selected from as many different litters as possible at each time point. Day of vaginal opening was
13 monitored. Estrous cyclicity was evaluated daily for 18 days at 4 and 6 months of age in 18–28
14 rats/group. Eight female offspring/group were killed 3 months later to measure serum LH levels using an
15 LH assay kit; a total of 6–8 values/group were obtained. Body and uterine weights and LH levels were
16 analyzed by ANOVA followed by *t*-test, Tukey test, or least significant difference test. Mammary tumors
17 were analyzed by chi-squared test, and estrous cyclicity data were analyzed by Kruskal-Wallis test and
18 Mann-Whitney *U* test.

19
20 On PND 4, 7, and 11, body weights were significantly higher in pups from the bisphenol A groups than in
21 the control group; body weights were higher in animals of the low compared to the high dose group. Body
22 weights of low-dose females were higher than body weights of control and high-dose females at PND 28
23 and beyond. While the percentage of control females with regular estrous cycles was 83% at 4 months of
24 age and 60% at 6 months of age, the values were significantly reduced in the high dose group to 21% at 4
25 months of age and 23% at 6 months of age. There were no clear patterns of estrous cycle changes. Periods
26 of diestrus were extended in some animals and other animals had extended periods of proestrus and/or
27 estrus. The mean number of 4–5-day estrous cycles was significantly reduced in rats of the high-dose
28 group at 6 months of age. Serum LH levels in the high-dose group were significantly reduced by ~19%
29 compared to the control group [**BMD₁₀ = 0.94, BMDL₁₀ = 0.48, BMD_{1SD} = 1.6, and BMDL_{1SD} = 0.78**
30 **mg/kg bw/day**]. The treatment group incidences of females with mammary tumors (10% in controls,
31 20% in the low-dose group, and 28% in the high-dose group) were not statistically different. The study
32 authors noted that the study was not designed to detect mammary tumors and that the tumors were
33 detected during routine handling. No effects were reported for mean number of pups/litter, sex ratio, day
34 of vaginal opening, or anogenital distance in the neonatal period. [**Data were not shown for anogenital**
35 **distance.**] In comparing the effects on estrous cycles and LH levels in animals exposed in the perinatal
36 period to the lack of uterotrophic effects in animals exposed in the postpubertal period, the study authors
37 concluded that there was evidence of increased sensitivity to bisphenol A during the perinatal period.

38
39 **Strengths/Weaknesses:** This study incorporates a range of basic developmental and gross functional
40 reproductive endpoints, but the sample sizes are small and the statistical approach does not appear to use
41 litter as the unit. Actual exposures are poorly defined, particularly postnatally. The plausibility of the
42 estrous cycle changes is a strength.

43
44 **Utility (Adequacy) for CERHR Evaluation Process:** This study is barely adequate for the evaluation
45 process, with guarded weighting based on sample sizes and statistical approaches.

46
47 **Takashima et al. (299)**, supported by a Grant-in-Aid for Health Sciences Research [**sponsor not**
48 **indicated**], examined the effect of bisphenol A exposure during development on carcinogenicity induced
49 by N-nitrosobis (2-hydroxypropyl)amine. [**No information was provided about caging and bedding**
50 **materials used in this study.**] Female Wistar rats were fed either MF diet or soybean-devoid powder diet
51 (Oriental Yeast Co.). In each dietary group, 10–11 rats/group received bisphenol A at 0 or 1.0% diet.

3.0 Developmental Toxicity

1 Bisphenol A exposure commenced 10 weeks prior to mating and was continued through the mating,
2 gestation, and lactation periods. Total intakes of bisphenol A were reported at 21–22 g/rat. **[Assuming an**
3 **exposure period of ~16 weeks, mean bisphenol A intake over the course of the study was estimated**
4 **at ~200 mg/day. Based on reported body weights, bisphenol A intake was ~1600 mg/kg bw/day**
5 **during the prebreeding stage and 1000 mg/kg bw/day during gestation and at weaning.]** The rats
6 were mated to males fed CE-2 basal pellet diet (Clea, Inc.), and GD 0 was defined as the day of the
7 vaginal plug. Endpoints associated with pregnancy, delivery, and nursing were evaluated. Dam body
8 weight and food intake were measured. Offspring were not culled and were weaned at 3 weeks of age.
9 Dams were killed following weaning of offspring. Serum levels of thyroid hormones were measured in 2–
10 4 dams/group. Implantation sites were evaluated. Weights of several organs, including ovary, were
11 measured. The organs were fixed in 10% buffered formalin and processed for histopathological
12 evaluation. Offspring (n = 32–50/group) were evaluated for body weight gain, preputial separation, and
13 vaginal opening. Beginning at 5 weeks of age and continuing for 12 weeks, offspring in each group were
14 subdivided into 2 groups (n = 17–21/group/sex) that received either undosed tap water or tap water
15 containing 2000 ppm N-nitrosobis (2-hydroxypropyl)amine. Offspring were killed at 25 weeks of age.
16 Serum thyroid hormone levels were measured. Organs, including testis, ovary, and uterus were weighed.
17 In 5–19 offspring/sex/group, histopathological examinations were conducted in organs targeted by N-
18 nitrosobis (2-hydroxypropyl)amine (lungs, thyroid, esophagus, liver, and thymus). Data were analyzed by
19 Dunnett and chi-squared tests.

20
21 Dam body weight was lower in the 1.0% bisphenol A group fed MF diet compared to the MF diet control
22 during the gestation period and at weaning. Food intake and maternal serum levels of triiodothyronine,
23 thyroxine, and thyroid-stimulating hormone were unaffected by bisphenol A exposure. Changes in
24 weights or histopathological alterations of maternal organs, including uterus and ovary, were not observed
25 in the bisphenol A groups. **[Data were not shown by the study authors.]** Bisphenol A had no significant
26 effect on mating, fertility, duration of gestation, live-born pups, implantation loss, or offspring viability
27 through PND 21. In pups from dams exposed to 1.0% bisphenol A fed MF diet compared to pups from
28 MF controls, body weights were higher **[by 11%]** in females at 3 days of age and lower in males and
29 females at 10 days and 2 weeks of age **[16–22% decreases in males and 12–19% decreases in**
30 **females]**. In pups from dams exposed to 1.0% bisphenol A and fed soybean-free diet compared to pups
31 from the soybean-free controls, body weights of pups were increased in males at 3 weeks of age **[13%**
32 **increase]** and in females at 10 days and 3 weeks of age **[13–19% increase]**. Prenatal exposure to
33 bisphenol A did not affect preputial separation or vaginal opening. In 25-week-old rats that were not
34 exposed to N-nitrosobis (2-hydroxypropyl)amine, prenatal bisphenol A exposure was associated with
35 some thyroid-stimulating hormone elevations in males and females from the MF and soybean-free diet
36 groups. According to a statement in the study abstract, the study authors did not consider the effect on
37 thyroid-stimulating hormone to be related to bisphenol A exposure. There were no effects of N-nitrosobis
38 (2-hydroxypropyl)amine exposure on serum thyroid-stimulating hormone, triiodothyronine, or thyroxine
39 levels or on thyroid histopathology. No effects were observed on offspring organ weights. **[With the**
40 **exception of uterus and ovary, no organ weight data were shown.]** Prenatal bisphenol A exposure was
41 not associated with significant differences in the development of N-nitrosobis (2-hydroxypropyl)amine-
42 induced neoplasms in the offspring. The study authors concluded that bisphenol A exposure did not
43 induce tissue injury in rat dams or their offspring or affect the development of tumors in offspring
44 exposed to N-nitrosobis (2-hydroxypropyl)amine.

45
46 **Strengths/Weaknesses:** Strengths include the size and duration of the study. Surprisingly small effects
47 were found given the high exposure levels. This study seems to discount the importance of certain effects
48 on body weight and thyroid-stimulating hormone levels that might have received more attention in a
49 study with a non-tumor focus. Sample size is inadequate to address neoplasm endpoints. Information is
50 insufficient to judge the appropriateness of the statistical analyses and hence the reliability of findings.

51

3.0 Developmental Toxicity

1 **Utility (Adequacy) for CERHR Evaluation Process:** While the study is adequate for the evaluation
2 process, it uses insufficient animals for a full-strength evaluation of many of the main endpoints, and so
3 the weight and application of the conclusions must be limited.
4

5 **Kobayashi et al. (300)**, supported by the Japanese Ministry of the Environment, examined the effect of
6 prenatal and lactational bisphenol A exposure on somatic growth and anogenital distance in Sprague
7 Dawley rats. The same rats were used to measure plasma hormone levels and testicular testosterone
8 content in a study by Watanabe et al. (301) and apparently thyroid function in a study by Koybayashi et
9 al. (302). Rats were fed standard laboratory feed (CE-2, CLEA Japan, Inc.). **[No information was**
10 **provided about caging or bedding materials.]** Rats were randomly assigned to groups and 6 rats/group
11 were gavaged with bisphenol A (99.8% purity) at 0 (corn oil vehicle), 4, 40, or 400 mg/kg bw/day from
12 GD 6 through PND 20. GD 0 was defined as the day a vaginal plug was observed, but the day of birth
13 was not defined. Doses were based on the study by Kwon et al. (297) **[discussed in Section 3.2.3.3]**. On
14 PND 7, litters were culled to 10 pups, with equal numbers of males and females when possible. Offspring
15 were weaned on PND 21. Dams were weighed during the study. Body weight and anogenital distance
16 were measured in offspring at 1, 3, and 9 weeks of age. Plasma and testicular testosterone levels were
17 measured at 9 and 36 weeks of age, and plasma LH and FSH concentration were measured at 9 weeks of
18 age. Weights of liver, kidney, and testis were examined in offspring at 3 and 9 weeks of age. Ont to 10
19 (most often 6–10) offspring/group/sex were examined for body weight and anogenital distance at 1 week
20 of age and 4–6/sex/group at 3 and 9 weeks of age. A pair of male and female offspring/litter **[assuming**
21 **authors meant 1/sex/litter]** was examined for organ weights, and 4–6 males/group were used in hormone
22 analyses at 3 and 9 weeks of age. Statistical analyses included ANOVA followed by Dunnett test.
23

24 In the 40 mg/kg bw/day group, all pups from 1 dam were found dead on PND 2. Four of 6 dams of the
25 400 mg/kg bw/day group died on GD 21, and all pups born to 1 dam in that group died by PND 2.
26 Maternal body weight gain during pregnancy was reduced in the 400 mg/kg bw/day group. A transient
27 decrease in body weight gain was observed early in the lactation period in dams of the 40 mg/kg bw/day
28 group. In offspring from the 4 and 40 mg/kg bw/day group, no statistically significant effects were
29 observed for body or organ weights, anogenital distance, anogenital distance/g body weight, or anogenital
30 distance/body weight cubed at any time point in the study. At 9 weeks of age, plasma testosterone levels
31 were significantly increased by 88% in the 4 mg/kg bw/day group and by 123% in the 40 mg/kg bw/day
32 group. **[Benchmark dose was not calculated because the SD was provided only graphically.]** The
33 study authors stated that there was a tendency for plasma testosterone to increase with dose at 36 weeks,
34 but neither of the values were significantly increased compared to control. Testis testosterone was not
35 statistically different from control at either dose at 9 or 36 weeks of age. There were no significant effects
36 on plasma LH and FSH levels at 9 weeks of age. Plasma levels of 17 β -estradiol were also unaffected by
37 bisphenol A exposure. **[Data were not shown.]** The study authors concluded that gestational and
38 lactational exposure to bisphenol A did not affect somatic growth or anogenital distance but did have a
39 significant effect on testosterone homeostasis in rat offspring.
40

41 **Strengths/Weaknesses:** The study appears better able to address maternal toxicity than offspring
42 outcomes, for which it appears to be best considered a screening study. Sample sizes are too small to
43 reliably judge postnatal endpoints.
44

45 **Utility (Adequacy) for CERHR Evaluation Process:** This study is inadequate for the evaluation of
46 bisphenol A effects on postnatal outcome.
47

48 **Yoshino et al. (303)**, supported by the Japanese Ministry of Health, Labor, and Welfare, examined the
49 effects of prenatal and lactational bisphenol A exposure in the prostate and testis of rats. In this study,
50 pregnant and lactating dams were fed NMF feed and offspring were fed MF feed (Oriental Yeast Co.,
51 Tokyo). The animals were housed in an unspecified type of cage containing wood chip bedding. F344 rat

3.0 Developmental Toxicity

1 dams (n = 19–22/group) were gavaged with bisphenol A (99.9% purity) at 0 (0.5% sodium
2 carboxymethylcellulose vehicle), 7.5, or 120 mg/kg bw/day during mating, gestation, and lactation
3 periods. Doses were based on the result of an NTP study (127). Clinical signs, food intake, and body
4 weight were monitored in dams during the study. After birth, pups were counted and weighed. Pups were
5 randomly culled to 8/litter on PND 4 (day of birth not defined). On PND 21, weaning occurred and
6 female pups were killed and discarded. Dams were killed at weaning for examination of implantation
7 sites. Male pups were weighed during the post-weaning period. On PND 23, 28, and 91, five male
8 offspring/group were killed. Ventral prostate weights were measured during each evaluation period, and
9 anterior and dorsolateral prostate, testis, and epididymis weight were also measured on PND 91.
10 Reproductive organs were preserved in 10% buffered formalin and examined histologically. Sperm count,
11 motility, and morphology were examined on PND 91. The study was repeated with evaluation of 10 male
12 offspring/group. **[The number of dams treated/group was not reported. Based on body weights
13 reported for rats in experiment 2, it appears they were evaluated at adulthood, but it was not
14 specified if they were evaluated on PND 91.]** Data were analyzed by Student *t*-test.

15
16 In the first experiment, bisphenol A exposure had no effect on dam body weights during gestation or
17 lactation, duration of the gestation period, or number of implantation sites. There were no effects on litter
18 size, pup viability, or sex ratio. On PND 21, relative dorsolateral prostate weight was significantly higher
19 **[by 23%]** in the low-dose group than in controls. **[It was not stated if organ weights were relative to
20 body weight.]** There were no effects on final body weight or weights of anterior and ventral prostate,
21 testis, or epididymis. There were no increases in malformations of reproductive organs. **[Data were not
22 shown by study authors.]** Testicular sperm counts were significantly lower **[by 22%]** in males of the
23 high-dose group, but there were no effects on epididymal sperm counts. There were also no effects on
24 sperm motility or abnormalities. **[Data were not shown by authors.]** In the second experiment
25 examining 10 offspring/group, exposure to bisphenol A had no effects on final body weights or relative
26 weights of testis, epididymis, or ventral, anterior, or dorsolateral prostate. There were no adverse effects
27 on testicular or epididymal sperm count, motility, or morphology. Morphologically abnormal sperm were
28 reduced in rats of the low-dose group. Study authors concluded that under the conditions of their study,
29 exposure of dams to bisphenol A during the gestation and lactation periods did not result in adverse effect
30 on the reproductive system of male offspring.

31
32 **Strengths/Weaknesses:** The number of dams used in Experiment 1 appears adequate; however, small
33 numbers of male offspring were examined at multiple time points to determine various organ endpoints. It
34 is unfortunate that these data were then analyzed by many *t*-tests rather than multivariate analyses. This
35 report seems to more closely resemble a screening study than a definitive study.

36
37 **Utility (Adequacy) for CERHR Evaluation Process:** This study is considered inadequate because of
38 small sample size and statistical insufficiencies.

39
40 **Ichihara et al. (304)**, supported by the Japanese Ministry of Health, Labor, and Welfare, examined the
41 effects of prenatal and lactational exposure to bisphenol A on the development of prostate cancer in rats.
42 F344 rat dams were fed NMF feed during pregnancy and lactation and their offspring were fed MF
43 (Oriental Yeast Co.) following weaning. Rats were housed in cages containing wood chip bedding. **[No
44 information was provided about caging materials.]** During pregnancy and lactation, ~8–15 dams/group
45 were gavaged with bisphenol A (99.9% purity) at 0 (0.5% carboxymethyl cellulose sodium salt vehicle),
46 0.05, 7.5, 30, or 120 mg/kg bw/day. Doses were based on findings from an NTP study **[citation not
47 provided]**. Dam body weight and food intake were monitored during the study. Gestation period duration
48 and implantation sites were evaluated. Pups were counted and sexed at birth. Litters were randomly culled
49 to 8 pups on PND 4, and pups were weaned on PND 21 **[day of birth not defined]**. At 5 weeks of age, 21
50 male rats/group were injected sc with 50 mg/kg bw 3,2-dimethyl-4-aminobiphenyl 10 times at 2-week
51 intervals. An additional 12 rats/group in the 0, 0.05, 7.5, and 120 mg/kg bw/day bisphenol A groups were

3.0 Developmental Toxicity

1 injected with corn oil during the same time period. Surviving male offspring were killed and necropsied at
2 65 weeks of age. Blood was collected for analysis of serum testosterone levels in 5 rats/group.
3 Reproductive organs were examined for gross abnormalities, weighed, and fixed in 10% buffered
4 formalin. A histopathological examination of the prostate was conducted. Body and organ weight data
5 were analyzed by Student *t*-test. The incidence of histopathological lesions was evaluated by Fisher exact
6 probability test.

7
8 Body weights of dams in the 120 mg/kg bw/day group were significantly lower than control values from
9 GD 14 to 20. There were no consistent or dose-related effects on dam body weights during lactation,
10 although a significant increase in body weight was observed in dams of the 0.05 mg/kg bw/day group on
11 PND 14. Exposure to bisphenol A had no effect on gestation period duration or number of implantation
12 sites. In pups exposed to bisphenol A, there were no differences in number of live births, sex ratio,
13 external anomalies, or body weights during the lactation period. **[Data for pup body weights were not
14 shown by study authors.]** Terminal body weight of pups exposed to 0.05 mg/kg bw/day bisphenol A
15 prior to treatment with 3,2-dimethyl-4-aminobiphenyl were significantly higher than controls **[by 12%]**.
16 Exposure to bisphenol A had no effect on weights of prostate, testis, or epididymis. Incidences of
17 prostatic intraepithelial neoplasia, carcinoma, and atypical hyperplasia were not increased by bisphenol A
18 treatment, and there were no increases in tumors found in non-reproductive organs. No effect was
19 observed on serum testosterone levels. The study authors concluded that exposure of rat dams to
20 bisphenol A during the gestation and lactation periods does not predispose their offspring to prostate
21 cancer development.

22
23 **Strengths/Weaknesses:** Strengths are the range of dose levels and the endpoints evaluated. The design is
24 reasonable for the endpoints measured, but sample sizes are inadequate for the prostate cancer endpoint
25 and hormonal endpoints in particular. This experiment represents a good screening study.

26
27 **Utility (Adequacy) for CERHR Evaluation Process:** This study is inadequate by itself based on
28 insufficient sample size. It might make a useful contribution when considered with other studies.

29
30 **Yoshida et al. (104)**, supported by the Japanese Ministry of Health, Labor, and Welfare, examined the
31 effects of bisphenol A exposure on development of the rat female reproductive tract. Donryu rats (12–
32 19/group) were gavaged with bisphenol A **[purity not reported]** at 0 (carboxymethylcellulose solution),
33 0.006, or 6 mg/kg bw/day from GD 2 to the day before weaning of pups at 21 days post delivery. The low
34 dose was said to represent average daily intake from canned foods and the high dose was reported to
35 represent the maximum dose level detected in plastic plates for children. **[It is assumed the authors
36 meant estimated exposure levels for children eating off plastic plates.]** Bisphenol A levels were
37 measured in maternal and pup tissues, and those values are reported in Section 2.1.2.2.1. After delivery,
38 dams and litters were housed in plastic cages with wood chip bedding. Tap water was stored in plastic
39 containers. The only information provided about feed was that it was a commercial pellet diet. Samples of
40 tap water, drinking water from plastic containers, and feed were measured for bisphenol A content by
41 HPLC. Offspring were sexed, weighed, and examined for external abnormalities on PND 1 and then
42 weighed weekly through PND 21. Litters were adjusted to 8–10 pups at PND 4 or 6 (day of birth = PND
43 0). Dams were weighed, and observed during the study and killed following weaning of litters on PND
44 21. Implantation sites were examined and organs including uterus, vagina, and ovaries were fixed in 10%
45 neutral buffered formalin and examined histologically. **[It does not appear that results of
46 histopathological testing in dams was reported.]** All female offspring were examined for vaginal
47 opening, and following vaginal opening, vaginal smears were taken for the remainder of the study. Three
48 to 5 offspring/group from different litters were killed on PND 10, 14, 21, or 28 and at 8 weeks of age. At
49 most time periods, uteri were weighed, preserved in 10% neutral buffered formalin, and examined
50 histopathologically to determine development of uterine glands. Ovaries and vagina were also examined
51 histologically. ER α was determined using an immunohistochemical method. Serum was collected for

3.0 Developmental Toxicity

1 measurement of FSH and LH by RIA. Four offspring/group from different litters were killed at 8 weeks
2 of age on the morning of estrus to examine ovulation by counting ova in oviducts. Initiation of
3 carcinogenesis following injection of the uterine horn with N-ethyl-N'-nitro-nitrosoguanidine was
4 examined at 11 weeks of age in 35 or 36 animals/group. At ~24 weeks following cancer initiation, the 24–
5 30 surviving animals/group were killed and uteri were examined histologically to determine the presence
6 of tumors and other lesions. Statistical analyses included ANOVA and Dunnett test.

7
8 Bisphenol A was not detected in fresh tap water but was detected at ~3 ng/mL following storage in plastic
9 containers. The bisphenol A concentration in feed was ~40 ng/g. In dams exposed to bisphenol A, there
10 were no clinical signs of toxicity or effects on body weight, implantation sites, or gestation length.
11 Bisphenol A exposure had no effect on litter size, pup body weight at birth and through PND 21, external
12 abnormalities in pups or age of vaginal opening. In uteri of bisphenol A-exposed offspring, there were no
13 effects on weight, gland development, ER α , or cell proliferation. No increase in lesions was reported in
14 organs of the alimentary, urinary, respiratory, or nervous system. **[Data were not shown by study**
15 **authors.]** Bisphenol A exposure had no effect on ovulation, estrous cyclicity, or serum FSH or LH levels.
16 There were no effects on uterine preneoplastic or neoplastic lesions or ovarian histopathology following
17 bisphenol A treatment. The study authors concluded that perinatal exposure to bisphenol A at levels
18 comparable to human exposure did not affect the reproductive system of female rats.

19
20 **Strengths/Weaknesses:** Strengths of this study were the bisphenol A determinations that were made and
21 the anchoring of animal exposure levels to human exposures. The design appears sound with a good range
22 of endpoints measured. Small numbers of animals were sacrificed at several time points and cellular
23 analyses were performed; these numbers were too small for a definitive cancer evaluation and were, in
24 fact, too small for definitive conclusions to be reached for most of the adult reproductive endpoints
25 Statistics are not described in enough detail to determine how data from multiple sampling points were
26 evaluated. This experiment represents a thorough screening study.

27
28 **Utility (Adequacy) for CERHR Evaluation Process:** This study is inadequate by itself based on
29 insufficient sample size. It might make a useful contribution when considered with other studies.

30
31 **Takagi et al. (305)**, supported by the Japanese Ministry of Health, Labor, and Welfare, examined the
32 effect of perinatal bisphenol A exposure on the reproductive and endocrine systems of rats. Nonylphenol
33 was also examined but will not be discussed. Sprague Dawley rat dams were fed a soy-free diet (Oriental
34 Yeast Co., Tokyo) prepared according to the formula for NIH-07. At weaning, the offspring were fed
35 CRF-1 diet (Oriental Yeast Co., Tokyo), which contains soybean and alfalfa-derived proteins. Rats were
36 housed in polycarbonate cages containing wood chip bedding. Dams were randomly assigned to groups,
37 and 5–6 dams/group were fed diets containing bisphenol A (96.5% purity) at 0, 60, 600, or 3000 ppm
38 from GD 15 (GD 0 = day of vaginal plug) to PND10 (PND 1 = day of birth). The study authors estimated
39 bisphenol A intake at ~5, 49, and 232 mg/kg bw/day during the gestation period and ~9, 80, and 384
40 mg/kg bw/day during the lactation period. Dose levels were based on results of preliminary studies, and
41 selected with a goal of achieving weak to moderate toxicity in dams at the highest dose. In a separate
42 study, rats were fed diets containing ethinyl estradiol at 0 or 0.5 ppm from GD 15 to PND 10. On PND 2,
43 offspring were counted, sexed, and weighed and anogenital distance was measured. Litters were culled to
44 6 pups on PND 10, and pups were weaned on PND 21. Five pups/sex/group (1/sex/litter) were selected
45 for necropsy on PND 21 and brain, adrenals, testis, ovary, and uterus were weighed. Eight
46 offspring/sex/group (at least 1/sex/litter) were selected for evaluation in adulthood, and these rats were
47 observed for age and body weight at puberty. Estrous cyclicity was observed from 8 to 11 weeks of age.
48 Offspring were killed at 11 weeks of age, on the day of diestrus for cycling female rats. Brain, pituitary,
49 thyroid, adrenal mammary gland, epididymis, prostate, seminal vesicles, ovary, uterus, and vagina were
50 weighed and examined histologically. The testis was fixed in Bouin solution, and other organs were fixed
51 in 10% neutral buffered formalin. The volume of the sexually dimorphic nucleus of the preoptic area

3.0 Developmental Toxicity

(SDN-POA) was measured. It appears that endpoints were assessed in 8 adult rats/sex/group, with the exception of histopathological evaluations, which were conducted in 5 rats/sex/group. The litter was considered the experimental unit in statistical analyses of data from PND 21 offspring, and the individual animal was considered the statistical unit for data obtained from adult offspring. Homogenous numerical data were analyzed by ANOVA and Dunnett test, and heterogenous numerical data were analyzed by Kruskal-Wallis *H* test and Dunnett-type rank sum test. Data for histopathological lesions and vaginal cyclicity were analyzed by Fisher exact probability test or Mann-Whitney *U* test.

Maternal body weight gain was significantly decreased the high-dose bisphenol A group during gestation, but there were no effects on body weight gain during lactation or food intake. In offspring evaluated on PND 2, there were significant decreases in body weight in low- and high-dose males [13 and 22%] and in high-dose females [20%], but there were no effects on number of live offspring or anogenital distance. Body weight gain was lower in high-dose males [21%] and females [29%] from PND 2 to 10. Benchmark doses calculated for dams and developing offspring are presented in Table 74. Increased relative brain weight as a result of growth retardation was reported in high-dose offspring evaluated on PND 21. [Data were not shown by study authors.] Exposure to bisphenol A did not affect onset of vaginal opening, preputial separation, or estrous cyclicity. Body weight of males was significantly lower [by 9.3%] at adult necropsy. Weights and histopathology of brain, pituitary, thyroid, adrenal mammary gland, epididymis, prostate, seminal vesicles, ovary uterus, and vagina in adulthood were unaffected in rats from the bisphenol A group. [Organ weight data were not shown by study authors.] Bisphenol A did not affect SDN-POA volume. Effects observed in offspring from the ethinyl estradiol study included reduced numbers of live offspring, increased male:female ratio, decreased body weight and body weight gain, accelerated vaginal opening, delayed preputial separation, increased estrous cycle irregularities, and histopathological alterations in pituitary, ovary, uterus, vagina, and mammary gland. The study authors concluded that bisphenol A did not affect endocrine or reproductive system development of rats at doses that induced maternal toxicity.

Table 74. Summary of Benchmark Doses for Body Weight Effects in Rats Exposed to Bisphenol A.

Affected population	Benchmark dose, ppm in diet [estimated mg/kg bw/day ^a]			
	BMD ₁₀	BMDL ₁₀	BMD _{1SD}	BMDL _{1SD}
Dam, during gestation	734 [60]	566 [46]	1208 [99]	850 [69]
Male pup, PND 2	2835 [232–378]	1294 [106–173]	2770 [226–369]	975 [80–130]
Female pup, PND 2	2860 [234–381]	1550 [127–207]	2808 [229–374]	1271 [104–169]
Male pup, PND 2–10	1595 [130–213]	890 [73–119]	1906 [156–254]	1060 [87–141]
Female pup, PND 2–10	1814 [148–242]	1080 [88–144]	2795 [228–373]	984 [80–131]

^aEstimates are based on author calculation of dam intake during gestation for dam weight and during gestation and lactation for pup weights. From Takagi et al. (305).

Strengths/Weaknesses: Strengths include the range of endpoints measured and the use of the 17 β -estradiol comparator group, as well as the complete statistical evaluation. The study used small sample sizes of dams as well as offspring for selected endpoints, though slightly larger than many other screening studies. The range of endpoints was better than average and included gross assessment of the volume of the highly relevant SDN-POA.

Utility (Adequacy) for CERHR Evaluation Process: This study is considered barely adequate for the evaluative process, based on sample size.

Akingbemi et al. (306), supported by NIEHS, US EPA, NICHHD, and NIH, conducted a series of studies in Long Evans rats to determine the effects of postweaning and perinatal exposure to bisphenol A on testicular steroidogenesis. In vitro studies were also conducted and are described in Section 4 because

3.0 Developmental Toxicity

1 cells used in the studies were obtained from adult animals. Rats were fed Purina chow, which contains
2 soybean meal, and given drinking water in polycarbonate bottles. Pregnant and nursing dams were housed
3 in polycarbonate cages lined with wood bedding, but no information was provided on caging used at the
4 other life stages. To reduce leaching of bisphenol A, the cages were washed, rinsed, and dried at least
5 twice/week and were discarded once they began getting cloudy; water bottles were cleaned daily. Corn oil
6 vehicle was used for bisphenol A and was administered to control animals. Rats were stratified according
7 to body weight and randomly assigned to treatment groups. RIA methods were used to measure steroid
8 hormone concentrations in serum or testicular fluid. RT/PCR methods were used to examine changes in
9 mRNA expression. Statistical analyses included ANOVA with multiple comparisons conducted by the
10 Duncan multiple range test.

11
12 In the first study, rats were gavaged with bisphenol A [**purity not given**] at 0, 0.0024, 0.010, 100, or 200
13 mg/kg bw/day from PND 21 through 35. The two lowest doses were selected to represent environmental
14 exposures, and the higher doses were selected to compare the effects between low and high doses. Rats
15 were killed at the end of treatment and blood was collected for measurement of serum LH, testosterone,
16 and 17 β -estradiol levels. Leydig cell cultures were prepared for measurement of ex vivo testosterone
17 production with and without the addition of LH, testosterone precursors, or metabolizing enzymes.
18 Additional weanling rats were exposed to bisphenol A at 0 or 0.0024 mg/kg bw/day on PND 21–35. At
19 the end of treatment, mRNA for *LH β* , *ER β* , and *ER α* was measured in pituitary using an RT-PCR
20 technique. All endpoints were reported for 7–12 rats/group. Compared to rats in the control group, rats
21 exposed to bisphenol A at 0.0024 mg/kg bw/day had significantly lower levels of serum LH [**by 62%**]
22 and testosterone [**by 40%**]. Serum 17 β -estradiol levels were decreased in rats exposed to 0.0024, 0.010,
23 and 100 mg/kg bw/day bisphenol A [**by ~30, 40, and 25% in each respective dose group**]. There were
24 no effects on basal ex vivo testosterone production by Leydig cells. In Leydig cells obtained from rats
25 exposed to 0.0024 mg/kg bw/day bisphenol A, testosterone production was significantly reduced when
26 cells were exposed to LH or CYP450 17 α -hydroxylase/17–20 lyase. In Leydig cells obtained from rats
27 exposed to 0.0024 or 0.010 mg/kg bw/day bisphenol A, testosterone production was significantly reduced
28 following exposure of the cells to pregnenolone or progesterone. No effects on blood hormone levels or
29 ex vivo testosterone production were observed at higher doses. Significant effects observed in pituitaries
30 obtained from rats exposed to 0.0024 mg/kg bw/day bisphenol A were decreased *LH β* mRNA and
31 increased *ER β* mRNA. The study authors concluded that the decreased serum LH level resulted from
32 bisphenol A effects on the pituitary and that decreased LH stimulation of Leydig cells was the cause of
33 reduced serum testosterone levels.

34
35 In the second experiment, 7 dams/group were gavaged with bisphenol A at 0 or 0.0024 mg/kg bw/day
36 from GD 12 through PND 21. Male offspring received no further treatment following weaning. Males
37 were randomly selected from each dam and killed on PND 90. Endpoints evaluated in 10–12 male
38 offspring/group included serum LH and testosterone levels, ex vivo testosterone production by Leydig
39 cells, testosterone levels in testicular interstitial fluid, and seminal vesicle and prostate weight. Significant
40 ($P < 0.01$ or 0.05) effects observed in 90-day-old males that had been perinatally exposed to bisphenol A
41 compared to the control group included increased body weight [**10%**], decreased relative weight (to body
42 weight) of paired testes [**17%**] and seminal vesicles [**17%**], reduced testicular testosterone level [**~39%**],
43 and reduced basal and LH-induced ex vivo testosterone production.

44
45 In the third experiment, 10–12 rats/group were gavaged with bisphenol A at 0 or 0.0024 mg/kg bw/day
46 from PND 21 through 90. Within 24 hours following treatment, rats were killed and examined for the
47 same endpoints described for the second experiment. Significant ($P < 0.01$ or 0.05) effects compared to
48 the control group included increased serum LH level [**117%**], decreased seminal vesicles weight
49 [**absolute: 15%, relative: 16%**], reduced testicular testosterone level [**~24%**], and decreased basal and
50 LH-induced ex vivo testosterone production. For the second and third experiments, the study authors
51 concluded that bisphenol A exposure inhibits androgen production by Leydig cells.

3.0 Developmental Toxicity

1
2 **Strengths/Weaknesses:** Significant strengths of this report were the sequential nature of the work, in
3 which later studies built on the previous data, and the clear expertise that the authors brought to this
4 endeavor. Experiment 1 provided a helpful examination of postnatal effects following adolescent
5 exposure and examined hormonal levels under stimulated and unstimulated conditions, thus separating
6 pituitary from target organ contributions to serum levels. In Experiment 2, the sample size of 7
7 dams/prenatal treatment group and the examination of 10–12 offspring/group raise questions about the
8 adequacy of the sample size with respect to the number of litters represented and the number of offspring
9 used to represent each litter. In Experiment 3, 10–12 rats/group were treated through adolescence and into
10 early adulthood and then examined according to endpoints common to Experiments 1 and 2. Weaknesses
11 include an inadequate number of animals to obtain confidence about the hormonal changes (indeed, LH
12 was decreased in the first experiment and increased in the third), the lack of histopathology evaluation,
13 and lack of an estrogenic positive control..

14 **Utility (Adequacy) for CERHR Evaluation Process:** While individual studies are borderline adequate
15 based on insufficient sample size for determining in vivo hormone levels for these episodically-released
16 hormones, the data collectively appear adequate to indicate that these low exposures are interfering with
17 androgen production. In aggregate, the paper is considered adequate.
18

19 **Tan et al. (307)**, supported by the University of Malaya and the Ministry of Science, Technology, and
20 Environment, examined the effects of bisphenol A exposure on pubertal development of male rats.
21 Sprague Dawley rats were fed soy-free feed (Gold Coin Feedmills) and housed in aluminum cages
22 containing shredded recycled paper as bedding. On PND 23–53, twelve rats/group were gavaged with 100
23 mg/kg bw/day bisphenol A [**purity not reported**] in a Tween-80/water solution (1:9 v/v), 100 mg/kg
24 bw/day nonylphenol in corn oil, or a mixture of 100 mg/kg bw/day bisphenol A and nonylphenol. A
25 control group of 12 rats was gavaged with Tween 80 in corn oil. Dosage selection was based upon
26 published studies reporting NOAELs of 50 mg/kg bw/day for both compounds. Rats were examined for
27 preputial separation during the study. Six rats/group were killed on PND 52, and the other 6/group were
28 killed on PND 53. Testes, epididymides, liver, kidney, adrenal, seminal vesicles plus coagulation gland,
29 and thyroid were weighed. [**The Expert Panel assumes that by coagulation gland, the authors mean**
30 **the anterior prostate or coagulating gland.**] Thyroid, testis, kidney, and liver were fixed in 10%
31 formalin and examined histologically. Statistical analyses included ANOVA and Fisher protected least
32 significant difference test.
33

34 There was no significant effect on weight gain in rats treated with bisphenol A. In the bisphenol A group,
35 preputial separation occurred by PND 53 in 66.7% of rats compared to 100% of rats in the control group.
36 In the bisphenol A group, significant increases were observed in absolute and relative (to body weight)
37 kidney and thyroid weights and significant decreases were observed for absolute and relative liver weight.
38 Cortical thickness of the kidney was significantly decreased [**by ~13% compared to controls according**
39 **to CERHR calculations and ~30% according to study authors**]. There was no effect on testicular
40 weight or tubule diameter. Normal patterns of spermatogenesis were observed in rats from the control
41 group. Multinucleated giant cells were observed in seminiferous tubules and there was no indication of
42 spermatogenesis in 4 of 12 rats of the bisphenol A group. Giant cells were observed and spermatogenesis
43 was found to occur in only some seminiferous tubules of the remaining rats treated with bisphenol A.
44 Moderate-to-severe hydronephrosis was observed in 50% of rats and mild hydronephrosis was observed
45 in the other 50% of rats from the bisphenol A group.
46

47 Preputial separation occurred by PND 53 in 33.3% of rats in the nonylphenol group and 58.3% of rats
48 exposed to the bisphenol A/nonylphenol mixture. In animals treated with nonylphenol, relative liver
49 weight was increased, absolute and relative seminal vesicle weights were decreased, and the diameter of
50 testicular tubules was reduced. A decrease in relative seminal vesicle weight was the only significant
51 organ weight effect observed in the group treated with both bisphenol A and nonylphenol. Moderate

3.0 Developmental Toxicity

1 hydronephrosis was observed in 25% of rats exposed to the bisphenol A/nonylphenol mixture and mild
2 hydronephrosis was observed in the other rats from that exposure group. No spermatogenesis was
3 observed in 3–5 of 12 rats/group treated with nonylphenol or the mixture of bisphenol A/nonylphenol.
4 The study authors concluded that both bisphenol A and nonylphenol affected the reproductive system of
5 rats, while only bisphenol A affected the kidneys. They also noted a less-than-additive effect with
6 administration of the bisphenol A/nonylphenol mixture.

7
8 **Strengths/Weaknesses:** This study was apparently well performed and documents the endpoints tested.
9 A weakness is the use of a single high dose level of bisphenol A administered by a single route. Other
10 minor weaknesses include the exposure period, which specifically avoids early development thus only
11 providing information on exposure in an important but possibly less biologically critical window. No
12 attempt was made to measure the tissue levels of bisphenol A achieved by the dosing regimen. It is also
13 unfortunate that no histology was performed on the seminal vesicles and coagulating gland tissue as
14 histology would have provided additional information relating to testicular function, specifically,
15 providing an integration of serum testosterone levels.

16
17 **Utility (Adequacy) for CERHR Process:** This study has moderate utility for the evaluation process and
18 raises concern about the effects of bisphenol A on testicular development and function, consistent with
19 observations in other studies. The utility is limited by the high dose level used.

20
21 **Kobayashi et al. (302)**, supported by the Japanese Ministry of Health, Labor, and Welfare, examined the
22 effects of developmental exposure to bisphenol A on thyroid status in rats. Rats used in this study were
23 fed standard laboratory chow (CE-2, Clea Japan). **[No information was provided about caging or**
24 **bedding materials.]** From GD 6 (day of copulatory plug = GD 0) through PND 20 (day of birth not
25 defined), 6 maternal CD rats/group were gavaged with bisphenol A (>99.8% purity) at 0 (corn oil
26 vehicle), 4, 40, or 400 mg/kg bw/day. The 400 mg/kg bw/day maternal group was excluded from analysis
27 because of excessive maternal toxicity. Details about maternal toxicity and additional aspects of this study
28 are available in the summary for the study by Kobayashi et al. (300). On PND 7, litters were culled to 5
29 pups/sex when possible. It appears that culled pups may have been used in analyses conducted at 1 week
30 of age. Pups were weaned on PND 21. Approximately 1 male and female pup/litter were killed at 3 and 9
31 weeks of age. Plasma thyroxine levels were measured by chemiluminescence immunoassay in 1–9
32 offspring/group/sex at 1 week of age and 3–6 offspring/sex/group at 3 and 9 weeks of age. At 9 weeks of
33 age, thyroid stimulating hormone test was conducted in 2–7 rats/sex/group by measuring thyroxine levels
34 after injection with bovine thyroid stimulating hormone. Statistical analyses included ANOVA followed
35 by Dunnett test or Student or Welch *t*-test. In the 4 and 40 mg/kg bw/day groups, there were no significant
36 differences in thyroxine levels at 1, 3, or 9 weeks of age or in thyroid stimulating hormone-induced
37 increases in thyroxine levels at 9 weeks of age. Based on the findings of this study, the study authors
38 concluded that prenatal and lactational exposure of rats to bisphenol A does not appear to affect thyroid
39 function.

40
41 **Strengths/Weaknesses:** Strengths of this study include the use of a range of dose levels of bisphenol A,
42 with a proviso that these dose levels were extremely high. Weaknesses include the limited endpoints
43 addressed (thyroid function) and the small numbers of animals used, which limits our confidence in the
44 conclusions.

45
46 **Utility (Adequacy) for CERHR Evaluation Process:** As presented, this publication is suitable but has
47 minimal utility for this evaluation.

48
49 **Zoeller et al. (308)**, supported in part by NIH, examined the effect of bisphenol A exposure on the
50 thyroid of developing rats. Sprague Dawley rats were housed in plastic cages. **[No information was**
51 **provided about composition of feed or bedding materials.]** Prior to initiation of dosing, rats were

3.0 Developmental Toxicity

1 trained to eat an untreated wafer each day. On GD 6 (day of vaginal plug not defined) through the
2 remainder of the experiment (the remainder of the gestation and lactation periods, according to the study
3 abstract), 9 rats/group were given a wafer dosed with bisphenol A [**purity not reported**] at levels
4 resulting in exposure to 0 (methanol vehicle), 1, 10, or 50 mg/kg bw/day. Doses were based on those used
5 in the study by Tyl et al. (293). Pups (n = 7–9/group/sex/time period) were weighed and killed on PND 4,
6 8, 15, or 35 (day of birth not defined). During each of those time periods, serum thyroxine was measured
7 by RIA. On PND 15, brains of male pups were sectioned and examined for presence of RC3/neurogranin
8 mRNA, a thyroid hormone-responsive gene, using an in situ hybridization and autoradiography
9 technique. Serum thyroid-stimulating hormone was measured using an unspecified method in 6–8 male
10 pups/group on PND 15. Statistical analyses included ANOVA and Bonferroni *t*-test.

11
12 The text of the study indicated a significant reduction in maternal body weight gain during pregnancy in
13 the high dose group, while Figure 1 of the study indicated a significant reduction in maternal body weight
14 gain during pregnancy at all dose levels. Maternal body weight gain during the lactation period was
15 unaffected by bisphenol A treatment. Bisphenol A exposure had no effect on litter size at birth. [**Data**
16 **were not shown by study authors.**] Bisphenol A had no effect on pup body weights on PND 4, 8, or 15.
17 On PND 15, but at no other time period, there was a significant increase in serum thyroxine levels in all
18 dose groups of male and female pups [**percent increases compared to controls were ~11, 35, and 37%**
19 **in each respective dose group.**] Significant increases in expression of RC3/neurogranin mRNA were
20 observed in the upper and lower dentate gyrus in males from each treatment group [**with no apparent**
21 **dose-response relationship**]. Expression of RC3/neurogranin mRNA in cortex was unaffected by
22 bisphenol A treatment. No significant effects were observed for thyroid-stimulating hormone levels in
23 males on PND 15. The study authors concluded that bisphenol A acts as a thyroid antagonist at these
24 concentrations.

25
26 **Strengths/Weaknesses:** Strengths of the study include use of a range of doses and examination of a role
27 of bisphenol A as a thyroid hormone antagonist. Weaknesses include the extremely high doses of
28 bisphenol A required to elicit these effects, the lack of litter-based analysis, and the lack of a positive
29 control.

30
31 **Utility (Adequacy) for CERHR Evaluation Process:** This can be considered in the evaluation process
32 but it is of limited utility based on the inadequate statistical analysis.

3.2.3.3 Studies with neurobehavioral endpoints

35 **Kwon et al. (297)**, from CIIT, examined the effects of bisphenol A exposure during pre- and postnatal
36 development on reproductive endpoints and the SDN-POA of rats. Sprague Dawley rats were fed NIH-07
37 feed from Zeigler Brothers and housed in polycarbonate cages with cellulose fiber-chip bedding. Water
38 was provided in glass bottles with Teflon-lined caps. Pregnant rats were randomly assigned to groups
39 according to body weight. Rats (n = 8/group) were gavaged with bisphenol A (~99% purity) at 0 (corn oil
40 vehicle), 3.2, 32, or 320 mg/kg bw/day from GD 11 (GD 0 = day of sperm detection) through PND 20,
41 excluding the day of parturition. It was not stated if the day of parturition was considered PND 0 or 1. A
42 positive control group was treated with 15 µg/kg bw/day diethylstilbestrol. Rats were examined for
43 clinical signs of toxicity and weighed during the study. Pups were weighed on PND 1 and 7. Pups were
44 weaned on PND 21. After pups were weaned, dams were killed for assessment of body and organ
45 weights. On PND 10, brains were collected from 1–3 female offspring/litter from 6–8 litters/group for
46 measurement of SDN-POA volume. All remaining female pups were examined for age of vaginal
47 opening and day of first estrus, and estrous cyclicity was monitored for 22 days, beginning at ~4 months
48 of age. Lordosis behavior was examined at 6 months of age in 1–2 female offspring/litter from 7–9
49 litters/group that had been ovariectomized 2 weeks prior to reproductive behavior testing and primed with
50 estradiol benzoate and progesterone. Male offspring were killed on PND 180 for measurement of body
51 and reproductive organ weights and histopathological evaluation of ventral prostates fixed in 10% neutral

3.0 Developmental Toxicity

1 buffered formalin. **[Based on information presented in the results section, it also appears that ovaries**
2 **and uteri were examined in an unspecified number of female offspring at 6 months of age.]**

3 Statistical analyses included ANOVA, Dunnet test, ANCOVA, and pair-wise comparison of least square
4 means.

5
6 Bisphenol A treatment had no significant effect on maternal body weight during pregnancy or lactation or
7 on maternal liver, kidney, adrenal, ovary, or uterus weights. There was no effect on number of live
8 pups/litter at birth or pup weight on PND 1 or 7. In female offspring, bisphenol A exposure had no
9 significant effect on volume of SDN-POA, age or weight at vaginal opening or first estrous, estrous
10 cyclicity, or mean lordosis intensity. In male offspring, there were no significant effects on body weight
11 or weights of testis, epididymis, seminal vesicle, or prostate. The study authors noted that a 23% increase
12 in ventral prostate weight in the high-dose group did not reach statistical significance. No treatment-
13 related histopathological alterations were reported for ventral prostate, ovary, or uterus. Effects observed
14 in the diethylstilbestrol group included increased maternal liver weight, increased SDN-POA volume in
15 female offspring, and disrupted estrous cycles. The study authors concluded that indirect exposure of
16 offspring to bisphenol A at these levels during gestation and lactation did not affect estrogen-mediated
17 reproductive endpoints. A similar study with comparable findings in females was reported in abstract
18 form (309).

19
20 **Strengths/Weaknesses:** This study was well performed and presented. The wide coverage of the dose
21 range (across a three log range) is a major strength. The use of diethylstilbestrol as a positive control is a
22 strength, as is the number of reproductive organs and endpoints evaluated. A weakness was the limited
23 analysis of those reproductive organs (wet weight only; histopathology was only performed on the
24 prostate) and a lack of determination of pup exposure during lactation.

25
26 **Utility (Adequacy) for CERHR Evaluation Process:** This study is adequate for the evaluation process.

27
28 **Kubo et al. (310)**, supported by the Japanese Ministry of Education, Science, and Culture, examined the
29 effect of prenatal bisphenol A exposure on sexually dimorphic behavior and brain development in rats.
30 **[No information was provided about the type of chow, bedding, or caging materials used.]**
31 Throughout the gestation (from the day that sperm were detected in the vagina) and lactation periods, 5
32 Wistar rats/group were administered bisphenol A through drinking water at 0 or 5 mg/L. **[No**
33 **information was provided on bisphenol A purity or use of a vehicle.]** The study authors estimated the
34 bisphenol A dose at 1.5 mg/kg bw/day and stated that the dose was below the NOAEL of 50 mg/kg
35 bw/day. **[It is not clear whether this is an estimate based upon water consumption or dosing by a**
36 **separate, unspecified route. If based upon drinking, this estimate is suspect because it implies a**
37 **daily consumption of approximately 70 mL water (because the weight of the rats is not supplied this**
38 **must of necessity be a guess), which is well in excess of published intakes for post partum rats**
39 **(generally noted as around 20 mL/day). It is also noted that water consumption varies widely in**
40 **non-lactating rats and throughout the period of lactation in rats, reflecting milk production, so any**
41 **such estimate would of necessity be suspect, and doses will vary with time post partum.]** Litters were
42 adjusted to 5 pups/sex on the day following birth. Pups were weaned on PND 21 **[day of birth not**
43 **defined]** and housed according to sex and litter. Behavior was tested for 10 minutes in an open-field
44 apparatus at 6 weeks of age (n = 11–14) **[It was not clear if the number of animals examined included**
45 **total animals, total/group, or total/sex/group. Litter distribution was not indicated.]** A passive
46 avoidance test was conducted at 7 weeks of age (n = 11–14); the test included a habituation period and
47 testing of retention 24 hours later. An unspecified number of rats were killed and necropsied at 12 weeks
48 of age, with females killed in diestrus. Reproductive organs were weighed (n = 12–14) and sperm
49 endpoints were evaluated in an unspecified number of rats. Serum hormone levels were measured by RIA
50 (n = 5–10/group). At 20 weeks of age, 6 rats/sex from the control group and 7 rats/sex from the treated
51 group were killed to measure the volume of the SDN-POA and the locus ceruleus. Behavioral data were

3.0 Developmental Toxicity

1 analyzed by Student *t*-test or Mann-Whitney *U* test, and brain morphology data were analyzed by Student
2 *t*-test. **[No information was provided on data analyses for reproductive organ weight, serum**
3 **hormone levels, or sperm endpoints.]**
4

5 In open-field testing of controls, females moved significantly greater distances, reared more times, and
6 stayed in the center of the apparatus for a longer period of time than males. In passive avoidance testing
7 of controls, latency to enter the dark chamber following electric shock was significantly longer in male
8 than female rats. In rats exposed to bisphenol A, there were no significant differences in the behaviors of
9 males compared to females. Study authors attributed the loss of sexually dimorphic behaviors to
10 demasculinization of male behavior and defeminization of female behavior. Bisphenol A treatment did
11 not affect brain weight, which was higher in male than female controls. The larger size of SDN-POA in
12 male compared to female controls was retained following bisphenol A treatment. The volume of the locus
13 ceruleus was significantly larger in females than males of the control group. In the bisphenol A group, the
14 volume of the locus ceruleus was described as larger in males than females, but the stated increase was
15 not statistically significant ($P = 0.12$). **[Graphically, there is an estimated 14% difference between**
16 **male and female locus ceruleus volume in controls and in bisphenol A-exposed animals, with the**
17 **direction of the difference apparently reversed by treatment.]** Bisphenol A treatment had no effect on
18 absolute weight of the testis or epididymis or relative weights of the ventral prostate, ovaries, or uterus.
19 There were no significant effects on serum levels of LH, FSH, testosterone, or 17 β -estradiol. Sperm count
20 and motility were also unaffected by bisphenol A exposure. The study authors concluded that current
21 methods for establishing NOAELs may not be sufficient to detect disrupted sexual dimorphism in the
22 brain.
23

24 **Strengths/Weaknesses:** A strength is the variety of biological and behavioral endpoints assessed. The
25 major weakness of the study is the lack of experimental detail, which makes it difficult to determine how
26 much bisphenol A was received by the animals.
27

28 **Utility (Adequacy) for CERHR Evaluation Process:** Given the lack of methodologic data provided in
29 the paper, this communication has limited utility for the evaluation process.
30

31 **Kubo et al. (311)** examined the effect of prenatal bisphenol A exposure on sexually dimorphic behavior
32 and brain structure of rats. **[No information was provided on the type of feed or materials used in**
33 **bedding or caging.]** Wistar rats were dosed with the 0.1% ethanol in distilled water vehicle ($n = 5$
34 dams/group) or bisphenol A **[purity not reported]** at 0.1 or 1 mg/L ($n = 6$ dams/group). The study
35 authors estimated bisphenol A intake at 0.030 and 0.3 mg/kg bw/day and noted that the levels were below
36 the tolerated daily intake. **[Though not clearly stated, it appears that as in the previous study by**
37 **Kubo et al. (310), exposures occurred through drinking water during the entire gestation and**
38 **lactation period.]** Five dams/group were exposed to *trans*-resveratrol, an estrogenic compound found in
39 grapes, at 5 mg/L or diethylstilbestrol at 50 μ g/L. Body weight and anogenital distance were measured in
40 pups on PND 1 (the day following birth). **[All litters were examined and although the number of pups**
41 **examined in each litter was not clearly stated, it was implied that all pups were analyzed.]**
42 Anogenital distance was adjusted by the cube root of body weight. Following the evaluations on PND 1,
43 litters were standardized to 5 pups/sex. Pups were weaned on PND 21 and housed according to sex and
44 litter. Day of testicular descent or vaginal opening was monitored in all remaining offspring ($n = 25$ /sex in
45 the control group and 30–31/sex in the treated group). Open-field testing was conducted in 20–24
46 animals/group at 6 weeks of age. **[It is not clear if the authors meant 20–24 animals/group or**
47 **animals/group/sex].** Sexual behavior of 7–13 male and female rats/sex/group was tested at 11–12 weeks
48 of age. Males and females ($n = 11$ –15/sex/group) were killed at 12 weeks of age, females during
49 proestrus. Reproductive organs were weighed. Serum hormone levels were measured by RIA. Sperm
50 from one testis and cauda epididymis were counted. Histological examinations were conducted on testis

3.0 Developmental Toxicity

1 fixed in Bouin solution and ovary fixed in 10% neutral buffered formalin. Rats were killed at 14 weeks of
2 age for measurement of SDN-POA and locus ceruleus volume in 7–8 males and females/group.

3
4 Because of the large number of animals used, the experiment was conducted in 3 blocks representing
5 identical experiments. All data were collected and analyzed following completion of the third block of the
6 study. The litter was considered the statistical unit in analyses of data collected prior to weaning of
7 animals. Individual animals were considered the statistical unit in analyses of data collected subsequent to
8 weaning. Behavior and brain structure data were analyzed by ANOVA and differences between sexes
9 were analyzed by Student *t*-test. Reproductive data were analyzed by ANOVA followed by Fisher
10 protected least significant difference test for each sex.

11
12 Bisphenol A exposure had no significant effect on body weight on PND 1, anogenital distance in males
13 and females, day of testicular descent, or day of vaginal opening. Body weight at vaginal opening was
14 significantly higher [by 7%] in the high-dose bisphenol A group. In sexual behavior testing of males, a
15 non-dose-related decrease in the intromission rate observed in the low-dose group was the only significant
16 effect reported following bisphenol A exposure. There were no effects on mounting or ejaculation.
17 Bisphenol A exposure had no significant effects on female sexual behavior as measured by ear wiggle,
18 lordosis quotient, and rejection of males. The study authors concluded that bisphenol A exposure had no
19 remarkable effects on male or female sexual behavior. The only significant effect on organ weights was
20 an [9%] increase in testis weight in the high-dose bisphenol group. There were no significant effects on
21 absolute weight or relative (to body weight) weights of ventral prostate, seminal vesicle, uterus, or ovary.
22 Bisphenol A treatment did not affect sperm count or motility or estrous cycles. Serum levels of LH, FSH,
23 prolactin, testosterone, and 17 β -estradiol were also unaffected by treatment. No histopathological findings
24 were observed in testis or ovary. [Data were not shown.]

25
26 In open-field testing of control rats, females moved greater distances, reared more often, and spent more
27 time in the center of the testing apparatus. Following treatment with the low or high dose of bisphenol A,
28 there were no longer significant differences between males and females in frequency of rearing and or
29 duration of time spent in the center of the apparatus. Differences in distances moved by males versus
30 females were no longer significant following exposure to the high bisphenol A dose. Males in the low
31 bisphenol A group reared significantly more times than males in the control group. Bisphenol A treatment
32 had no significant effect on the sex-related difference in size of the SDN-POA, which was significantly
33 larger in males than females in the control and treatment groups. Although the volume of the locus
34 ceruleus was significantly greater in females than males of the control group, locus ceruleus volume was
35 significantly larger in males than females of both bisphenol A groups. The change was due to a significant
36 increase in volume in males at the low dose and significant decrease in volume in females at both dose
37 levels of bisphenol A. [Magnitude of locus ceruleus volume changes in males and females was ~12–
38 17% compared to controls, as estimated from a graph.] The numbers of neurons in the locus ceruleus
39 was affected in the same manner as volume by bisphenol A treatment, except that increases in neuron
40 numbers following bisphenol A treatment were also significant in males of the high-dose group.

41
42 Diethylstilbestrol mainly affected open-field behavior, locus ceruleus volume, and the reproductive
43 system. *Trans*-resveratrol mainly affected locus ceruleus volume and the reproductive system. The study
44 authors concluded that the brain is highly sensitive to bisphenol A at levels below the tolerable daily
45 intake and disruptions in sexual differentiation may differ from effects observed with diethylstilbestrol
46 and *trans*-resveratrol.

47
48 **Strengths/Weaknesses:** As with the previous study by this group (310) the main weakness of the paper
49 lies in the failure to accurately describe the methods to allow a reader to determine how much bisphenol
50 A the dams received during the experiment. The subtlety and relevance of the neurologic endpoints
51 assessed was a major strength of this paper. The most significant finding related to brain development and

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1 the size of the locus ceruleus along with possibly related behavioral changes. No effects on the
2 reproductive tract were noted. A major strength is that this work apparently explored low dose exposures
3 to bisphenol A.
4

5 **Utility (Adequacy) for CERHR Evaluation Process:** This paper is adequate for the evaluation process
6 and gives cause for concern that even low levels of bisphenol A exposure in utero and/or during lactation
7 could result in gender-specific changes in brain maturation and resulting changes in sexual behavior.
8

9 **Facciolo et al. (312)**, supported in part by the Italian Ministry of University Education and Research,
10 examined the effects of developmental exposure to bisphenol A on the somatostatin receptor subtype sst₂
11 in the limbic circuit of rats. Sprague Dawley dams were exposed orally to bisphenol A at 0 (arachis oil
12 vehicle), 0.040, or 0.400 mg/kg bw/day. **[No information was provided on the specific method of oral
13 dosing, the purity of bisphenol A, or the number of dams treated/group. There was no information
14 on the type of chow used or composition of cage and bedding materials.]** The authors stated that the
15 doses selected were relevant to human exposures from can linings and dental sealants and had been
16 reported to induce morphometric changes in offspring. The rats were mated for 5 days during the
17 treatment period, and treatment was continued through gestation and lactation. Litters were culled to 8
18 pups at birth and it was 1 pup/litter was randomly assigned to a dam in the same treatment group. Pups
19 were weaned on PND 23 (day of birth not defined). On PND 10 and 23, 4–7 rats/group **[10–11/group
20 according to figures in the study]** were killed and their brains were removed to examine effects on sst₂
21 receptors in the limbic region. Receptor binding was assessed using ¹²⁵I-Tyr⁰-somatostatin-14 as a ligand.
22 At the same ages, interactions of sst₂ with α -containing γ -aminobutyric acid (GABA) receptors, using the
23 agonists zolpidem and Ro 15-4513, were examined in 12–13 rats/group. Results were reported for only
24 the high dose of bisphenol A (0.400 mg/kg bw/day) because higher affinity was obtained for receptor
25 ligand binding. Statistical analyses included Student *t*-test, ANOVA, and Newman-Keuls multiple range
26 test. Statistically significant results are summarized in Table 75, which shows variable results depending
27 on age and brain region. The study authors concluded, “These results support the contention that an sst₂
28 subtype α -containing GABA type A receptor system might represent an important neuromediating station
29 capable of promoting estrogenlike mechanism of [bisphenol A], especially during the early
30 developmental phases.”
31

1 **Table 75. Effects on sst₂ Receptors in Rats Exposed to 0.4 mg/kg bw/day Bisphenol A During**
 2 **Prenatal and Postnatal Development.**

Receptors	Binding levels	
	PND 10	PND 23
High affinity		
Periventricular nucleus	↑	↔
Ventromedial hypothalamic nucleus	↔	↓
in presence of Ro 15-4513	↓	↓
in presence of zolpidem	↓	↔
Dentate gyrus	↔	↑
in presence of Ro 15-4513	↓	↑
in presence of zolpidem	↑	↔
Basomedial nucleus of the amygdala	↔	↓
in presence of Ro 15-4513	↔	↑
in presence of zolpidem	↔	↑
Low affinity		
Dentate gyrus	↓	↔
in presence of Ro 15-4513	↑	↑
in presence of zolpidem	↑	↔
CA1 layer of the hippocampus	↑	↓
in presence of Ro 15-4513	↔	↓
Basomedial nucleus of the amygdala	↓	↔
in presence of Ro 15-4513	↑	↑
Cortico-medial nucleus of the amygdala	↔	↓
Ventromedial hypothalamic nucleus in presence of Ro 15-4513	↑	↔
Periventricular nucleus in presence of zolpidem or Ro 15-4513	↑	↑

↑,↓ Statistically significant increase, decrease compared to controls; ↔ no statistically significant effects compared to controls.

From Facciolo et al. (312).

3
 4 **Strengths/Weaknesses:** Strengths of this study are the fact that it appears to have been carefully
 5 performed and used biologically-relevant concentrations with specific neuropeptide receptors. These data
 6 suggest that the GABA system could mediate some of the xenoestrogenic effects of bisphenol A. Minor
 7 weaknesses include lack of some specific experimental details as noted above. The random assignment of
 8 1 pup/litter within treatment groups is a weakness.

9 **Utility (Adequacy) for CERHR Evaluation Process:** This paper represents well performed and
 10 appropriately controlled work, and is thus useful for inclusion in the evaluation process.

11
 12 **Facciolo et al. (313)**, supported by the Italian Ministry of University Education and Research, examined
 13 the effects of bisphenol A on expression of somatostatin subtype 3 (*sst₃*) receptor mRNA in brains of
 14 female rats exposed during development and investigated whether the α GABA_A receptor is also involved
 15 in this effect. Sprague Dawley rats were housed in stainless steel cages. **[No information was provided**
 16 **about the type of feed or bedding used.]** Beginning 8 days before mating and continuing through the
 17 mating period (5 or 8 days) and during pregnancy and lactation (42 days), 8 rats received the arachis oil
 18 vehicle and 12 rats/group received bisphenol A **[purity not reported]** at 0.040 or 0.400 mg/kg bw/day.
 19 Vehicle or bisphenol A were orally administered by pipette. To minimize litter effects, 1 female pup from
 20 each litter was fostered to a dam from the same treatment group (8 pups/dam). Pups were weaned on PND
 21 23. On PND 7 and at 55 days of age, 4 rats/group/time period were killed. Brains were sectioned and a
 22 ³²S-labeled probe was used in an in situ hybridization method to measure *sst₃* mRNA expression. The
 23 effects of α GABA_A receptor subunits on expression of *sst₃* mRNA was examined by incubating the brain
 24 sections in 1 nM–100 μ M of α GABA_A receptor agonists (zolpidem, flunitrazepam, RY 080, and RO 15-

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4513). Additional brain sections from high-dose rats were used to determine interactions between *sst*₃ with α_1 and α_5 subunits with or without addition of 5–500 nM zolpidem or RY 080. Statistical analyses included ANOVA followed by Dunnett *t*-test or Neuman-Keuls multiple range post hoc test, when analysis by ANOVA indicated statistical significance. The effects of bisphenol A on *sst*₃ mRNA expression are summarized in Table 76. Changes in *sst*₃ expression varied with dose and age. Expression patterns were changed in the presence of α GABA_A receptor agonists. Based on their findings, the study authors concluded that bisphenol A exposure can affect cross-talking mechanisms involved in the plasticity of neural circuits with resulting influences on neuroendocrine/sociosexual behaviors.

Table 76. Effects of Bisphenol A on Expression of *sst*₃ mRNA in Rat Brain

Brain region	Age in days and bisphenol A dose as mg/kg bw/day			
	7		55	
	0.040	0.400	0.040	0.400
<i>Incubated without αGABA_A receptor agonists</i>				
Layer III of frontoparietal cortex	↔	↑	↔	↑
Layer V of frontoparietal cortex	↔	↔	↔	↓
Radiatum hippocampal lacunosum molecular CA1 fields	↓	↓	↓	↓
Hypothalamic arcuate nucleus	Not reported	↔		↑
Ventromedial hypothalamic nuclei	↔	↑	↑	↑
Periventricular nucleus	↔	↓	Not reported	
<i>Incubated with the αGABA_A receptor agonist zolpidem</i>				
Layer III of frontoparietal cortex	Not examined	Not reported	Not examined	↑
Radiatum hippocampal lacunosum molecular CA1 fields	examined	↔	examined	↑
Stratum oriens and pyramidale of the CA1 hippocampus field		↓		Not reported
<i>Incubated with the αGABA_A receptor agonist RY080</i>				
Layer V of frontoparietal cortex	Not examined	Not reported	Not examined	↓
Radiatum hippocampal lacunosum molecular CA1 fields	examined	↓	examined	↓
Ventromedial hypothalamic nuclei		↓		↓
Stratum oriens and pyramidale of the CA1 hippocampus field		↓		Not reported
Hypothalamic arcuate nucleus		↑		
Periventricular nucleus		↓		

↑,↓ Statistically significant increase, decrease compared to controls; no statistically significant effect.

From Facciollo et al. (313).

Strengths/Weaknesses: This was well performed and appropriately controlled work examining the effects of antenatal and lactational exposure to bisphenol A provided orally to the dam on the expression profile of somatostatin receptor subtype 3 and the role of GABA in its expression profile. The strengths of the paper were the rigor with which the study was performed and the nature of the endpoints (receptor binding assays and in situ hybridization to assess localization of receptors). No major weaknesses were noted.

Utility (Adequacy) for CERHR Evaluation Process: This work was well performed and is valuable for the evaluation process.

Aloisi et al. (314), supported in part by the Italian Ministry for Universities and Scientific and Technological Research (MURST), examined the effects of prenatal or postnatal bisphenol A exposure on

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1 the pain response of rats. **[No information was provided on chow or composition of caging and**
2 **bedding.]** Sprague Dawley rats were fed peanut oil vehicle (n = 13) or 0.040 mg/kg bw/day bisphenol A
3 **[purity not given]** (n = 7/group) by pipette during pregnancy and lactation. Within 48 hours after birth,
4 the offspring were sexed and cross-fostered to form the following groups:

- 5
- 6 • Prenatal exposure group—born to dams receiving bisphenol A and nursed by dams receiving the
- 7 peanut oil vehicle (n = 11 males; 9 females)
- 8 • Postnatal exposure group—born to vehicle control dams but fostered to bisphenol A treated dams
- 9 (n = 11 males; 9 females)
- 10 • Vehicle control group—born to and nursed by dams exposed to the vehicle control (n=16 males
- 11 and 11 females)
- 12

13 At 22 weeks of age, the rats were randomly assigned to sham or formalin treatment groups, but the sham
14 group was not analysed. The formalin group was sc injected with 10% formalin on the dorsal surface of
15 the right hind paw. Pain behaviors, such as licking, flexing, and jerking of the paw were recorded for 60
16 minutes. Following testing, the phase of the estrous cycle was determined and blood was drawn to
17 measure plasma levels of testosterone in males and corticosterone and 17 β -estradiol in both sexes by RIA.
18 Data were analyzed by ANOVA followed by post hoc least significant difference test.

19
20 The frequency of paw jerking was decreased at 30–60 minutes following formalin injection in postnatally
21 exposed rats. **[The study abstract and results section indicate that the effect occurred in males and**
22 **females, but according to data presented in figures of the study, the effect only appeared to have**
23 **occurred in males.]** Duration of flexion was increased 0–30 minutes following formalin injection in both
24 sexes exposed prenatally to bisphenol A. Although statistical significance was not attained, the study
25 authors noted an increase in licking duration at 0–30 minutes following formalin injection in females
26 exposed to bisphenol A during prenatal development. No effects were observed on open-field behaviors
27 or plasma levels of testosterone, 17 β -estradiol, or corticosterone. The study authors concluded that their
28 findings indicated sex- and exposure-related modifications of neural pathway activity or nociception
29 centers following exposure to bisphenol A.

30
31 **Strengths/Weaknesses:** A strength of this study is the added dimension being investigated (pain
32 response). The lack of some methodologic details are annoying minor weaknesses, and confidence is
33 eroded by inconsistencies in the data presentation. This study separates out antenatal and postnatal
34 exposure but unfortunately does not include a group exposed at both times, another mild weakness.
35 Another weakness is the use of a single dose level.

36
37 **Utility (Adequacy) for CERHR Evaluation Process:** The data presented are weak, and are considered
38 barely adequate or useful for the evaluation process. Thus, limited weight is given to this report.

39
40 **Negishi et al. (315)**, support not indicated, examined the effect of perinatal bisphenol A exposure on
41 behavior of rats. F344/N rats (n = 8–9/group) were orally exposed to bisphenol A at 0 (olive oil vehicle),
42 4, 40, or 400 mg/kg bw/day from GD 10 through PND 20. GD 0 was defined as the day that vaginal
43 sperm were detected and PND 0 was defined as the day of parturition. **[No information was provided on**
44 **purity of bisphenol A, the specific method of oral dosing, type of chow used, or composition of**
45 **bedding or caging materials.]** Dams were observed and weighed throughout the study. On PND 0, pups
46 were counted, weighed, and culled to 8/litter with equal numbers/sex when possible. Pups were weighed
47 periodically from PND 7 through 84. Pups were housed as same-sex littermates following weaning on
48 PND 21. Upon weaning of pups, dams were killed and body and organ weights were recorded. Behavioral
49 testing of offspring consisted of spontaneous motor activity measured at 28–34 days of age (n = 12–
50 27/group), active avoidance testing conducted at 28–34 and 56–62 days of age (n = 8–9/group), and open-

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field behavior evaluations at 56–62 days of age (n= 9–18/group). **[It was not indicated if group sizes were for total numbers of animals/group or numbers/sex/group.]** On PND 62, offspring were randomly selected (8/sex/group) and killed for evaluation of body and organ weights. Statistical analyses included ANOVA, nested ANCOVA, and post hoc Fisher protected least significant difference test.

Statistically significant results are summarized in Table 77. Maternal body weight gain was reduced during the gestation and lactation period in dams exposed to the mid or high dose. The only organ weight effects in dams were reduced absolute and relative (to body weight) thymus weight. There were no effects on weights of liver, kidney, or spleen in dams. Bisphenol A treatment did not affect the number of pups/litter or sex ratio. In male offspring, body weights were lower than control values on PND 7 and 28 at the mid dose, and PND 7, 21, 28, and 56 at the high dose. Body weights of female offspring were lower than controls at PND 7 and 28 at the low and mid dose and PND 7, 21, and 28 at the high dose. On PND 62, there were no effects on body weight or liver, kidney, spleen, thymus, brain, or testis weights. There were no effects on spontaneous activity, but total immobile time was increased in females of the mid-dose group. Performance of males in avoidance testing improved in the mid- and high-dose group at 4 weeks of age but decreased in the low-dose group at 8 weeks of age. Increased grooming by males of the low-dose group was observed in open-field testing. The study authors concluded that perinatal bisphenol A exposure caused behavioral alterations that differed by sex.

Table 77. Effects in Rats Exposed Perinatally to Bisphenol A

Endpoint	Dose (mg/kg bw/day)		
	4	40	400
Maternal body weight during gestation and lactation	↔	↓	↓
Maternal thymus weight relative to body weight	↔	↔	↓35%
Body weight of male pups on PND 7, 21 and/or 28 ^a	↔	↓4%	↓7–10%
Body weight of female pups on PND 7, 21, and/or 28 ^a	↓3–4%	↓4–5%	↓4–8%
Body weight of male offspring on PND 56	↔	↔	↓6%
Immobile time by females in activity testing ^b	↔	↑33%	↔
Percent responses in avoidance testing by 4-week-old males ^b	↔	↑ 13–43%	↑ 13–71%
Percent responses in avoidance testing by 8-week-old males ^b	↓15%	↔	↔
Grooming by males at 8 weeks of age	↑77%	↔	↔

↑,↓ Statistically significant increase, decrease compared to controls; ↔no statistically significant effects compared to controls.

^aSee text for specific days pup body weight was affected.

^bValues estimated from a graph by CERHR.

From Negishi et al. (315).

Strengths/Weaknesses: Doses were sufficiently high to produce gross body weight changes, and 3 different measures of behavior were collected, as well as organ weights at necropsy from the same animals. The lack of an evaluation of hormone-dependent behaviors is a weakness, as is the lack of assessment of more hormone-dependent tissues (prostate, levator ani muscle, etc.) or processes (age at balanopreputial separation, natal anogenital distance). Analysis was apparently not litter-based and there was no positive control.

Utility (Adequacy) for CERHR Evaluation Process: This paper shows that behavioral changes can occur as a result of early bisphenol A exposure and is adequate for the evaluation process. The utility of this paper is decreased by the weaknesses noted above.

Negishi et al. (316), support not indicated, examined the effect of perinatal bisphenol A exposure on the behavior of rats. The effects of nonylphenol were also examined but will not be discussed. F344/N rats

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1 (10 or 11/group) were gavaged with bisphenol A at 0 (corn oil vehicle) or 0.1 mg/kg bw/day from GD 3
2 to PND 20. GD 0 was defined as the day that vaginal sperm was detected, and PND 0 was the day of
3 parturition. At birth, pups were counted and weighed. Litters were culled to 6 pups, with equal numbers
4 of each sex when possible. Pups were weighed throughout the postnatal period. At weaning, dams were
5 killed and organ weights were measured. One male pup/litter (n = 8–10/group) was subjected to a series
6 of behavioral tests. The remaining male pups were killed for measurement of organ weights at 21 days or
7 8 weeks of age. Neurobehavioral endpoints evaluated included open-field behavior at 8 weeks of age,
8 spontaneous motor activity at 12 weeks of age, passive avoidance at 13 weeks of age, performance in the
9 elevated-plus maze at 14 weeks of age, and active avoidance at 15 weeks of age. At 22–24 weeks of age,
10 a monoamine reduction test was performed: rats were injected with the monoamine oxidase inhibitor
11 trans-2-phenylcyclopropylamine hydrochloride or with saline, and behavior was then evaluated. Data
12 were analyzed by ANOVA, and if statistical significance was obtained, Fisher protected least significant
13 difference test was conducted.

14
15 Bisphenol A exposure did not affect dam body weights during gestation or lactation, gestation duration,
16 litter size, number of male and female pups, or final dam body and organ weights. **[Data were not
17 shown.]** Body and organ weights of male offspring at 21 days and 8 weeks of age, behavior in open-field
18 testing, spontaneous motor activity, and performance in the elevated-plus maze were also unaffected by
19 bisphenol A exposure. **[Data were not shown by study authors.]** Bisphenol A had no significant effect
20 on performance in the passive avoidance test, although tendencies for increased latency were observed. In
21 active avoidance testing, rats from the bisphenol A group had significantly ($P < 0.01$) fewer correct
22 avoidance responses during the first, second, and third of 5 sessions, and failure of avoidance was
23 significantly increased [**~2.5% in the bisphenol A group compared to 0.2% in controls**]. In contrast to
24 control rats, bisphenol A-treated rats did not show an increase in locomotion following a challenge with
25 trans-2-phenylcyclopropylamine hydrochloride. The number of rearings following 2-
26 phenylcyclopropylamine hydrochloride exposure did not differ significantly between rats from the
27 bisphenol A and control groups. The study authors concluded that perinatal exposure of rat dams to
28 bisphenol A at concentrations slightly higher than environmental exposures irreversibly affected
29 perception of fear-provoking stimuli and monoaminergic neural pathways in male offspring.

30
31 **Strengths/Weaknesses:** The use of a single dose level is a weakness. Strengths include the variety of
32 endpoints used to provide data, which point to effects that are not gross structural changes but relatively
33 subtle behavioral effects.

34
35 **Utility (Adequacy) for CERHR Evaluation Process:** These data are adequate for the evaluation process
36 and raise concern that the primary effects of low dose bisphenol A could be exhibited in terms of
37 behavioral modifications.

38
39 **Farabollini et al. (317)**, supported by the University of Siena, University of Firenze, MURST, and the
40 Italian National Research Council, examined the effects of perinatal bisphenol A exposure on behavior in
41 male and female rats. **[No information was provided about chow or composition of cage and bedding
42 materials.]** Three groups of Sprague Dawley rats were orally dosed with the arachis oil vehicle or
43 bisphenol A **[purity not reported]** by micropipette. One group of 11 rats was administered 0.040 mg/kg
44 bw/day bisphenol A from 10 days prior to conception until weaning of pups at 21 days of age. A second
45 group of 11 rats was given arachis oil from 10 days prior to conception through GD 13, 0.400 mg/kg
46 bw/day bisphenol A from GD 14 **[day of vaginal plug not stated]** through 6 days following delivery of
47 pups, and arachis oil until weaning of pups. A control group of 9 rats was given arachis oil from 10 days
48 prior to conception until weaning of pups. Beginning at 85 days of age and continuing for 3 days,
49 behavioral testing was conducted using a holeboard and elevated-plus maze in 15 offspring/sex from the
50 low-dose group, 11–12 offspring/sex from the high-dose group, and 14 pups/sex from the control group.
51 **[Litter distribution was not reported.]** Separate sessions were conducted for each sex and treatment

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group. Data were analyzed by ANOVA and Fisher least significant difference test. A factor analysis was conducted using the principal components method with an orthogonal rotation of the factor matrix.

Results of detailed analyses involving individual doses and sexes are summarized in Table 78. In general, head dipping (extending head over edge of apparatus) and arm entries were reduced and self-grooming was increased in exposed females. Head dipping and stretched-attend posture (moving body forward without moving paws and then returning to original position) were inhibited and arm entries were increased in exposed males. A factor analysis indicated reduced anxiety and motivation to explore in treated males and reduced activity and motivation to explore in treated females. The study authors concluded that although sex-related differences in behavior were noted following bisphenol A treatment, there was no clear masculinization of behavior in females. The authors also noted the lack of substantial differences in results between the two exposure protocols.

Table 78. Behavioral Testing in Rats Exposed Perinatally to Bisphenol A

Endpoint	Females, doses in mg/kg bw/day		Males, doses in mg/kg bw/day	
	0.040 ^a	0.400 ^b	0.040 ^a	0.400 ^b
<i>Holeboard test</i>				
Frequency of head dipping	↔	↓59%	↔	↓52%
Duration of head dipping	↓45%	↓74%	↔	↔
No. crosses	↓31%	↔	↔	↔
<i>Elevated plus maze test</i>				
Open-arm entries	↔	↔	↑178%	↔
Percent time in open arms	↔	↔	↑158%	↔
Closed-arm entries	↔	↓40%	↔	↔
Percent time in center	↓59%	↓71%	↔	↔
Total entries	↔	↓42%	↔	↔
Percent open/total entries	↔	↔	↑236%	↑206%
Frequency of self grooming	↑47%	↔	↔	↔
Frequency of stretched-attend posture	↔	↔	↓63%	↓43%

↑,↓ Statistically significant increase, decrease compared to control; ↔ no statistically significant change.

^aIn the low-dose group, dams were exposed to bisphenol A from 10 days prior to conception through gestation and lactation.

^bIn the high-dose group, dams were exposed to bisphenol A from GD 14 to PND 6.

From Farabollini et al. (317).

Strengths/Weaknesses: The unusual exposure scenario in this paper is both a strength and a weakness. The study appears to have been adequately and rigorously performed. The lack of the two obvious controls (prolonged high dose exposure and short low dose exposure) is a weakness. The behavioral effects induced by the two protocols were similar.

Utility (adequacy) for CERHR Evaluation Process: This study is adequate and useful for the evaluation process and raises concern that early exposure, even to low doses of bisphenol A, can give rise to permanent behavioral changes.

Farabollini et al. (318), supported by the University of Siena, University of Firenze, and MURST, examined the effects of perinatal bisphenol A exposure on sociosexual behavior in rats. Sprague Dawley rats were housed in polysulfone cages. [No information was provided on type of feed or composition of bedding materials.] Dams received arachis oil vehicle (n = 13) or 0.040 mg/kg bw/day bisphenol A (n = 7) through a micropipette from mating through weaning of pups. On the 2nd day following delivery, litters were culled to 4 pups/sex and cross-fostered to obtain the following exposure groups of 12 animals/sex:

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- 1
- 2 • Prenatal exposure group: born to bisphenol A-treated dams and nursed by vehicle-treated dams
- 3 • Postnatal group: born to vehicle-treated dams and nursed by bisphenol A-treated dams
- 4 • Control group: born to and nursed by vehicle-treated dams
- 5

6 Litters were weaned on PND 21 (day of birth not defined). On day 45 [**assumed to be PND 45**], animals
7 of the same sex were randomly chosen and housed 4/cage, with no siblings in any cage. At 100 days of
8 age, behavior in the presence of an intruder rat was observed. In female rats, vaginal smears were taken at
9 the end of intruder testing and only females in diestrus were considered (n = 8 – 9/group). One week later,
10 sexual orientation was tested in 12 rats/sex/group by placing a rat between cages containing a sexually
11 receptive female and sexually mature male and recording the number of visits to each rat. Sexual
12 performance was tested next in males; evaluation was restricted to only males that ejaculated (n = 10–12
13 group). One week later, sexual behavior was tested in females during the diestrus or proestrus phase.
14 **[It is not clear how many females were evaluated for sexual behavior.]** Behavior testing sessions were
15 videorecorded and later evaluated by a blinded observer. Data were analyzed by ANOVA followed by
16 post hoc Fisher least significant difference test.

17
18 In intruder testing, statistically significant effects observed in males exposed prenatally to bisphenol A
19 included an increased number showing defensive behavior (9 of 10 versus 4 of 10 in the control group), a
20 decreased number showing ambivalent behavior (3/10 versus 8/10 in the control group), and increased
21 ratio of defensive/agonistic behaviors [**by 280% compared to controls**]. No significant effects were
22 observed in intruder testing of female rats. There was no effect on sexual preference of males or females.
23 For sexual behavior testing of females, data from the pre- and postnatal exposure groups were pooled
24 because there were no significant differences between groups. Bisphenol A exposure significantly
25 decreased exit latency in females in diestrus [**by ~66%**] and proestrus [**by ~83%**] and significantly ($P <$
26 0.05) increased lordosis frequency in females in proestrus [**~11.75 versus 3.75 times in controls**].
27 Statistically significant effects on sexual performance of treated males included an increased number of
28 intromissions [**~15 compared to 11 in controls**] in the postnatal exposure group and increased duration
29 of intromission latency [**~115 versus 40 seconds in controls**] and genital sniffing [**~40 versus 16**
30 **seconds in controls**] in the prenatal exposure group. The study authors stated that the results suggested a
31 slight intensification of sexual behavior in females, slightly reduced performance in a limited number of
32 endpoints in males, but no effect on other important sexual endpoints in males (e.g., latency of ejaculation
33 and refractory period). It was concluded that pre- or postnatal exposure to bisphenol A potentiated female
34 behavior and depotentiated male behavior.

35
36 **Strengths/Weaknesses:** The work was carefully performed. The use of a single dose level of bisphenol A
37 is a weakness; however, this dosing paradigm is consistent with many other papers by this group making
38 comparisons between the papers relevant. Addressing aggressive/defensive behavior as well as sexual
39 performance and interest in both male and female offspring is a strength. The failure to address
40 underlying biological mechanisms is a weakness.

41
42 **Utility (Adequacy) for CERHR Evaluation Process:** This study is a serious contribution to the
43 bisphenol A literature and is suitable for the evaluation process. The observations of this study suggest a
44 potentiation of female behavior and a decrease in masculinity in adults resulting from perinatal exposure
45 to low doses of bisphenol A.

46
47 **Dessi-Fulgheri et al. (319)**, supported by the University of Firenze, University of Siena, and MURST,
48 examined the effect of perinatal bisphenol A exposure on play behavior in rats. Sprague Dawley rats were
49 housed in polysulfone cages. **[No information was provided on chow or bedding material.]** Using a
50 pipette, rats were fed solutions containing the arachis oil vehicle and/or bisphenol A according to 1 of 3
51 exposure scenarios. A control group of 9 rats was given arachis oil from 10 days prior to mating until

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1 weaning of pups on PND 21 [**day of birth not defined**]. Eleven rats in the low-dose group were given
2 0.040 mg/kg bw/day bisphenol from 10 days prior to mating until weaning of pups. Eleven rats in the
3 high-dose group received arachis oil vehicle from 10 days prior to mating until GD 13 [**day of vaginal**
4 **plug not defined**], 0.400 mg/kg bw/day bisphenol A from GD 14 to PND 6, and arachis oil from PND 7
5 until weaning. Both doses were considered to be within the range of human exposure. The low dose was
6 said to represent exposures through food occurring over a long period of time. The high dose was said to
7 represent exposures occurring through dental procedures occurring over a short period of time. Litters
8 were culled to 8 pups at birth. [**No information was provided on the sex distribution of the retained**
9 **pups.**] After pups were weaned, 3 male and 3 female pups were randomly caged together, with no
10 siblings co-housed in any cage. Behavioral testing was conducted on PND 35, 45, and 55. For the
11 behavioral testing, rats from the same cage were individually identified by marking them with dye. On
12 each day of testing, the 6 cage mates were transferred to a neutral arena that was covered in clean sawdust
13 and videorecorded for 6 minutes. Behaviors recorded during the 2nd and 3rd minute of each testing session
14 were evaluated. There were 12–15rats/sex/group. [**The methods section indicates that 15 rats/sex were**
15 **tested at the high dose, 12 rats/sex at the low dose, and 15 rats/sex in the control group. According**
16 **to Table 4 of the study, which gives the pooled number of rats tested for 3 age periods, it appears**
17 **that 12/sex were tested in the high-dose group, 15/sex in the low-dose group, and 15/sex in the**
18 **control group.**] For statistical analyses, individual factor scores were used as independent variables in a
19 3-way ANOVA that considered treatment, sex, and age. Fisher least significant difference test was used
20 when appropriate.

21
22 Behavioral elements were categorized under 8 general factors. The authors first presented results that
23 were pooled for the 3 different age groups. In females of the low-dose group, bisphenol A treatment was
24 found to significantly increase factors addressing play directed towards females. Factors affecting low-
25 intensity mating elements (e.g. crawling-under behavior) were significantly reduced in high-dose males
26 and females. Factors of sociosexual exploration (e.g., genital and body sniffing) were significantly
27 reduced in high-dose females and in males from both dose groups. Factors of social interest (e.g.,
28 approaching) were significantly reduced in both sexes at the high dose but increased in low-dose males.
29 The authors next discussed results for PND 35, because it is the approximate time period of vaginal
30 opening in females. Factors that were significantly affected at PND 35 included increased social interest
31 by males and females of the low-dose group, decreased low-intensity mating elements by females of both
32 dose groups, and decreased sociosexual exploration by males of both dose groups. The study authors
33 concluded that 2 factors of female behavior were masculinized by treatment: play with females and
34 sociosexual exploration.

35
36 **Strengths/Weaknesses:** A strength of this work is that it evaluated the socio-sexual consequences of
37 exposure, and specifically at a young age. Weaknesses included the fact that the two obvious controls
38 (prolonged high-dose exposure and short low-dose exposure) were apparently not performed, which
39 limits the degree to which the data can be interpreted.

40 **Utility (Adequacy) for CERHR Evaluation Process:** This paper is marginally useful for CERHR
41 evaluation process. That being said, the data are broadly consistent with other, perhaps more directly
42 relevant, papers. For this reason this paper raises only mild concern, although it is consistent with a larger
43 and perhaps more ominous overall image.

44
45 **Adriani et al. (320)**, supported by the Nervous and Mental Disorders Research Area, Istituto Superiore di
46 Sanità, Italy, and by MURST, examined the effects of perinatal exposure to bisphenol A on behavior in
47 rats. Sprague Dawley rats were housed in Plexiglass cages with sawdust bedding. [**No information was**
48 **provided about feed.**] Nine dams/group were dosed with bisphenol A [**purity not reported**] orally by
49 micropipette at doses of 0 (arachis oil vehicle) or 0.040 mg/kg bw/day from the day of mating to the day
50 pups were weaned. Pups were weaned on PND 25 (PND 0 = day of birth) and housed in groups of 3
51 according to sex. One male and 1 female/litter were observed in testing that included novelty-seeking

3.0 Developmental Toxicity

1 behavior during adolescence (PND 30–45), impulsivity during adulthood (PND 70), and open-field
2 behavior following injection with 1 mg/kg bw *d*-amphetamine during adulthood. It appears that the same
3 animals were tested at each time period. Data were analyzed by Tukey HSD test and ANOVA.
4

5 In novelty testing, the time spent in a new area of the testing apparatus was lower in females exposed to
6 bisphenol A [**~45–55% compared to vehicle control, $P < 0.05$**]. Males and females of the bisphenol A
7 group exhibited increased activity in the novel area [**increases of ~75% in males and 35–55% in**
8 **females, $P < 0.05$**]. The study authors interpreted the effects of novelty testing as suggesting a less
9 pronounced habituation profile and increased stress in a novel situation. In the impulsivity testing, food-
10 restricted animals were placed in an apparatus that involved nose poking in a small hole to immediately
11 deliver 1 pellet of feed or a larger hole to deliver 5 pellets of feed following a delay that was increased
12 over the time of the study. Lights were turned on during the delay periods following nose poking and
13 for 25 seconds after delivery of feed, time periods in which no feed could be delivered. Both groups of rats
14 preferred the larger hole with delayed delivery, but treatment with bisphenol A resulted in a more marked
15 preference for the larger hole ($P < 0.05$), thus indicating reduced impulsivity. When the length of the
16 delay was increased for the large hole, the frequency of inadequate responding (i.e., nose poking during
17 the delay) was decreased in males from the bisphenol A group; the study authors interpreted the effect as
18 indicating a demasculinization of the restlessness profile. [**Figures 2 and 3 of the study suggest that**
19 **there was decreased preference of the larger reinforcer and increased inadequate hole poking by**
20 **the bisphenol A group. It is not clear if an error may have been made in the text or illustration of**
21 **data. Because the statement, made both in the text and legend, contradicts the figure the expert**
22 **panel assumes that Figure 3a in the study is incorrectly labeled.**] In open-field testing, vehicle control
23 males displayed significantly more rearing and crossing behaviors following injection with *d*-
24 amphetamine, but an increase in rearing and crossing behavior following *d*-amphetamine injection did not
25 occur in males exposed perinatally to bisphenol A. The study authors concluded that perinatal exposure of
26 rats to bisphenol A resulted in altered behavior in rats.
27

28 **Strengths/Weakness:** This is another generally well performed study using protocols well established by
29 this group. The use of only a single exposure level of bisphenol A is a weakness, with the proviso that the
30 dose used is directly comparable to other studies. The authors' conclusion that bisphenol A caused
31 demasculinization of behavior is not supported by the lack of a male-female difference in behavior in the
32 control animals. The major problem noted above is the question of labeling of figures. This would seem
33 to be a clear-cut case of incorrect labeling; however, doubt about this label must be noted. Assuming that
34 the labeling is incorrect, the data, as stated in the text, reinforce the idea of demasculinizing effects of
35 perinatal exposure to bisphenol A occurring in both juvenile and adult animals.

36 **Utility (Adequacy) for CERHR Evaluation Process:** The weight given to this work must be diminished
37 by the figure-labeling issue. The paper is nevertheless consistent with many others and is suitable for
38 qualified evaluation.
39

40 **Porrini et al. (321)**, supported by MURST, the University of Firenze, and the University of Siena,
41 examined the effects of perinatal bisphenol A exposure on play behavior of female rats. [**No information**
42 **was provided about the type of feed or bedding and caging materials.**] Female Sprague Dawley rats
43 were co-housed with males for 36 hours and then fed the peanut oil vehicle ($n = 10$) or 0.040 mg/kg
44 bw/day bisphenol A [**purity not stated**] ($n = 12$) by micropipette during the gestation and lactation
45 period. Two days following delivery, litters were adjusted to 4 pups/sex and pups were fostered by a dam
46 from the same treatment group. Pups were weaned on day 21 [**assumed to be PND 21; day of birth not**
47 **defined**]. Offspring were housed in cages containing 3 pairs of male-female siblings, with no siblings of
48 the same sex in the same cage. Each group contained 18 female pups. Prior to examination of behavior in
49 rats from the same cages at 35, 45, and 55 days of age, animals were individually identified with dye.
50 Behavior was observed in a neutral arena in which the floor was covered with clean sawdust. Animals
51 were allowed to familiarize themselves to the new environment for 1 minute and then behavior was

3.0 Developmental Toxicity

1 videorecorded for 6 minutes. Video recordings were analyzed by an investigator blinded to treatment
2 conditions. Only behavior of female rats was considered. Data were analyzed by ANOVA for repeated
3 measures.

4
5 Factors were defined by study authors based on groups of behaviors. Significant effects were reported for
6 3 factors. Social and non-social exploration was increased [by ~34%] at 35 days of age and [by ~25%] at
7 45 days of age. Frequency of play behavior with males was decreased [by ~100%] at 45 days of age.
8 Grooming behavior was also decreased [by ~63%] at 45 days of age. The study authors concluded that
9 bisphenol A does not clearly induce masculinization of female behavior, but some aspects of female
10 behavior were defeminized.

11
12 **Strengths/Weakness:** This paper reports a well-performed study with a poorly researched endpoint
13 (juvenile play behavior) that has implications for reproductive behavior later in life. As with all
14 behavioral studies, the results had to be compiled in a blinded fashion in order to prevent subjective
15 evaluations. The authors have done a good job of making sure that the data are as objective as possible,
16 giving the reader confidence that the findings have real meaning. The use of only a single dose level of
17 bisphenol A is a weakness. The fostering of pups within treatment group prevents the evaluation of
18 intrauterine effects. The evaluation of play with factor analysis is questionable because only rough-and-
19 tumble play is sexually dimorphic. In addition, this behavior is organized by androgens, not estrogens,
20 decreasing the biologic plausibility of the conclusions.

21
22 **Utility (Adequacy) for CERHR Evaluation Process:** This work is suitable for the evaluation process.
23 The study supports concern that perinatal bisphenol A exposure leads to behavioral changes, in this case
24 defeminization of behavior, which appear later in life.

25
26 **Carr et al. (322)**, supported by the National Science Foundation, the Mississippi Agricultural and
27 Forestry Experiment Station, and the College of Veterinary Medicine at Mississippi State University,
28 examined the effects of bisphenol A exposure on performance of rats in the Morris water maze. In this
29 study, F344 rat dams and pups were fed Purina Test Diet 8117, a casein-based rodent chow. [No
30 information was provided about caging or bedding materials.] Treatment groups were assembled by
31 including pups from different litters. Ten pups/sex/group were gavage dosed from PND 1 (day of birth =
32 PND 0) through PND 14 with bisphenol A (>99% purity) at 0 (safflower oil vehicle), 0.1, and 0.25 mg/kg
33 bw/day. An additional group of rats was gavaged with 17 β -estradiol 72 μ g/kg bw/day during the same
34 time period. Straight channel swimming was tested on PND 33. Spatial learning and memory were tested
35 by Morris water maze for 4 days beginning on PND 34. In the test, acquisition of maze solution occurred
36 when the rat found a platform. A probe trial measuring the amount of time spent in an escape quadrant
37 from which the platform had been removed was conducted on PND 40. Data were analyzed by ANOVA
38 followed by means separation by least squared means or Greenhouse-Geisser adjusted F ratios.

39
40 There were no significant effects of bisphenol A treatment on straight channel swimming or time to
41 acquisition of maze solution in the Morris maze test. Time spent in the escape quadrant was significantly
42 lower in females of the high-dose group [by ~38%] than in controls. The study authors noted that
43 acquisition of maze performance was significantly better in control males than control females. However,
44 no sex-related difference was observed following treatment with the low bisphenol A dose. Increased
45 time to acquisition in males on the third day of testing, and no sex-related differences in performance were
46 reported for the 17 β -estradiol group. The study authors concluded “These data indicate that [17 β -
47 estradiol] and low dosages of [bisphenol A] can alter the normal sex-dependent pattern of acquisition,
48 while higher dosages of [bisphenol A] alter the retention of spatial information without significantly
49 affecting acquisition.”

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1 **Strengths/Weaknesses:** Strengths are the additional behavioral dimensions captured by this paper and
2 the use of a positive control. Though the doses were high, they were not damagingly so. The analyses
3 appeared appropriate. A weakness is the limited number of endpoints investigated.
4

5 **Utility (Adequacy) for CERHR Evaluation Process:** This study appears adequate and useful for the
6 evaluation process. This study shows mild effects of bisphenol A and of itself does not give rise for
7 concern about this agent in relation to the limited criteria examined.
8

9 **Della Seta et al. (323)**, supported by MURST and the University of Siena, examined the effects of
10 pubertal bisphenol A exposure on behavior of male rats. **[No information was provided about feed,**
11 **caging, or bedding.]** Seventy-eight Sprague Dawley males were obtained from 16 dams. On PND 23–30
12 (day of birth not defined), the rats were fed by micropipette peanut oil vehicle, 0.040 mg/kg bw/day
13 bisphenol A **[purity not reported]**, or 0.4 µg/kg ethinyl estradiol. **[The number of rats treated in each**
14 **group was not indicated.]** On PND 45, 12 animals/group were tested for social and non-social behavior
15 in response to a black PVC tube introduced into the cage. Behaviors were grouped according to factors of
16 play and social interaction, environmental exploration and social investigation, and elements directed to
17 the object. Twelve adults/group (> 90 days of age) were tested for sexual behavior with a sexually
18 receptive female. Males that were not used in behavioral testing were killed on PND 37 and 105 to
19 measure plasma 17β-estradiol and testosterone levels by RIA (n = 5–8 animals/group/evaluation period).
20 Data were assessed by ANOVA and Fisher least significant difference test.
21

22 Bisphenol A exposure did not affect factors associated with environmental exploration and social
23 investigation or with play and social interaction. Behaviors directed to the object (biting, sniffing,
24 climbing) occurred at a significantly lower frequency in the bisphenol A than control group. Compared to
25 the vehicle controls, the ethinyl estradiol group exhibited lower frequencies of behaviors associated with
26 environmental exploration or social investigation and with behaviors directed to the object. In sexual
27 behavior testing, 9–10 of 12 animals/group were sexually active and considered in data analysis.
28 Decreased intromission latency was the only parameter significantly affected in males from the bisphenol
29 A group. Significant effects in the ethinyl estradiol compared to the control group included increased
30 frequency of intromission, ratio of intromission/mount, and duration of refractory period. Rats exposed to
31 ethinyl estradiol also experienced significant decreases in mount latency, intromission latency, and genital
32 sniffing than control rats. On PND 37, the plasma testosterone level was significantly lower in the
33 bisphenol A and ethinyl estradiol group than in controls. The plasma testosterone level was significantly
34 lower in the bisphenol A than control group on PND 105. No effects were observed on plasma 17β-
35 estradiol levels. The study authors concluded that the few behavioral effects observed in the bisphenol A-
36 exposed rats occurred in the same direction as those observed in the ethinyl estradiol group but magnitude
37 of the effects was weaker in the bisphenol A than ethinyl estradiol group.
38

39 **Strengths/Weaknesses:** To be added
40

41 **Utility (Adequacy) for CERHR Evaluation Process:** To be added
42

43 3.2.4 Rat—parenteral exposure postnatally 44

45 3.2.4.1 Reproductive endpoints

46 **Fisher et al. (324)**, supported by the European Centre for Ecotoxicology of Chemicals and Zeneca,
47 examined the effect of neonatal bisphenol A exposure on excurrent ducts of the rat testis. On PND 2–12
48 (PND 1 = day of birth), Wistar rat pups were sc injected with the corn oil vehicle or 37 mg/kg bw/day
49 bisphenol A **[purity not given]**. The dose was based on the solubility limit in oil. **[The number of rats**
50 **treated was not indicated, but based on the number of rats examined in each time period (~4–7 in**
51 **treated group and 5–20 in control group), it appears that there were ~25/group in the bisphenol A**

3.0 Developmental Toxicity

1 **group and ~48 in the vehicle control group. No information was provided about feed, caging, or**
2 **bedding materials.]** Seven other compounds were also examined but will not be discussed, with the
3 exception of a brief explanation of results obtained with 0.0037–0.37 mg/kg bw/day diethylstilbestrol.
4 Rats were killed at 10, 18, 25, 35, and 75 days of age. Testes and epididymides were removed and fixed
5 in Bouin solution. Immunohistochemistry techniques were used to examine water channel aquaporin-1
6 levels. Morphology of rete testis and efferent duct were examined. Data were analyzed by ANOVA.
7

8 In the bisphenol A group, the only effect on testis weight was a significant decrease [**~40%**] at 35 days of
9 age. Epithelial cell height in the efferent ducts was significantly reduced [**by ~15%**] at 18 and 25 days of
10 age, but not at later time periods. There was no effect on expression of water channel aquaporin-1 protein
11 or morphology of the rete testis. Treatment with most diethylstilbestrol doses resulted in reduced
12 testicular weights at all ages, decreased expression of water channel aquaporin-1 protein, and decreased
13 epithelial cell height in efferent ducts at 25 days of age and younger, and fluid retention and enlargement
14 of rete testis, which was most severe at PND 18 and 25. The study authors concluded that the magnitude
15 and duration of adverse effects induced by estrogenic compounds were broadly similar to the estrogenic
16 potencies of the compounds.
17

18 **Strengths/Weaknesses:** This is a carefully performed study, although the inclusion of many
19 methodologic details (*vide supra*) would have improved it. Strengths include the use of a wide range of
20 estrogenic compounds to alter testicular development. A limitation for the present purpose is that only a
21 single dose level of bisphenol A was examined. A weakness is that tissues other than the testis were not
22 examined.
23

24 **Utility (Adequacy) for CERHR Evaluation Process:** This is a clear, well performed study suitable for
25 this evaluation. The effects observed on testicular development in relation to bisphenol A were minor and
26 transient; for this reason, bisphenol A exposure in this dose range does not raise concerns in relation to
27 this endpoint.
28

29 **Nagao et al. (325)**, supported by the Japanese Ministry of Health and Welfare, examined the effects of
30 neonatal bisphenol A exposure on reproductive function of male and female Sprague Dawley given CE-2
31 feed (Clea Japan). [**No information was provided about caging or bedding materials.]** From PND 1 to
32 5 (birth by 16:00 considered PND 0), 28–31 pups/sex/group were sc injected with corn oil vehicle, 300
33 mg/kg bw/day bisphenol A [**purity not reported**], or 2 mg/kg bw/day estradiol benzoate. Doses were
34 based on results of preliminary studies that demonstrated no effect on growth or viability at bisphenol A
35 doses up to 300 mg/kg bw/day administered by sc injection in the neonatal period. Pups were examined
36 for viability from PND 6 to 21. On PND 21, 5 pups/sex/group were randomly selected and killed. Pups
37 were transcidentally perfused, and reproductive organs were collected for histopathological evaluation. At
38 12 weeks of age, 22–25 rats/sex were mated with untreated rats. Females were killed on GD 13 for an
39 evaluation of implant number and viability of embryos. After fertility evaluation, sexual behavior with a
40 sexually receptive female was assessed in 10 males/group. Following evaluation of sexual behavior, 15
41 male rats/group were killed for measurement of reproductive organ and brain weight. Histopathology of
42 reproductive organs and SDN-POA volume were measured in 5 males/group. Copulation and fertility
43 indices were analyzed by chi-squared and Fisher exact 1-tailed test. Data for other endpoints were
44 analyzed by Student *t*-test.
45

46 In rats treated with bisphenol A, there were no clinical signs of toxicity or effects on pup viability or body
47 weight gain during or following the lactation period [**data for pup viability not shown by study**
48 **authors**]. There were no effects on age of vaginal opening or preputial separation. Copulation and
49 fertility indices and numbers of live embryos/litter were not affected in male or female rats treated with
50 bisphenol A. Bisphenol A treatment did not affect sexual behaviors of males, as determined by number of
51 mounts, intromissions, and ejaculations. No histopathological alterations were observed in the ovaries of

3.0 Developmental Toxicity

1 treated females at 21 days of age or in the epididymis, prostate, or seminal vesicles of treated male rats at
2 21 days or 14 weeks of age. **[The prostatic lobe not specified; based on the figure provided, the lobe
3 seems to have been ventral prostate. The Expert Panel notes that the number of apically located
4 nuclei may be elevated by 14 weeks of age over what would normally be expected; however, this
5 observation cannot be determined definitively based on a single high power field and in the absence
6 of a matched control.]** No effect of treatment was observed on the SDN-POA of males. In contrast to the
7 bisphenol A groups, rats treated with estradiol benzoate experienced decreased body weight gain,
8 compromised male sexual behavior, infertility, lesions in reproductive organs, and reduced volume of the
9 SDN-POA. The study authors concluded that neonatal exposure to a relatively high dose of bisphenol A
10 had no effect on morphological development or function of the reproductive system.

11
12 **Strengths/Weaknesses:** Strengths include a well performed and documented study that compared effects
13 of bisphenol A and estradiol benzoate. Additional strengths include documentation of both behavioral
14 (mating behavior) and biological (genital tract development) endpoints in both male and female rats.
15 Weaknesses include the use of only a single high dose level of bisphenol A and the choice of PND 1–5
16 for exposure, which might have excluded the most sensitive time periods.

17
18 **Utility (Adequacy) for CERHR Evaluation Process:** This paper is suitable for evaluation.

19
20 **Stoker et al. (326)**, support not indicated, examined the effects of prepubertal bisphenol A exposure on
21 prolactin secretion and prostate size in rats. **[No information was provided about feed, bedding, or
22 caging materials.]** On PND 22–32 (day of birth = PND 0), 15–17 male Wistar rats/group were sc
23 injected with bisphenol A **[purity not reported]** at 0 (sesame oil vehicle) or 50 mg/kg bw **[assumed to
24 be 50 mg/kg bw/day]**. Another group of rats was administered 17 β -estradiol through a sc Silastic tube
25 implant **[dose administered not clear]**. On PND 29, 6 animals/dose were killed and blood was collected
26 for measurement of serum prolactin concentration. The remaining rats (n = 9–11/group) were killed at
27 120 days of age. Prolactin levels were measured in serum and anterior pituitary by RIA. Inflammation
28 was visually examined in the ventral and lateral prostate. Left lateral and ventral prostates were weighed
29 and lateral prostate was analyzed for myeloperoxidase (an indicator of neutrophil numbers) and DNA.
30 The right lateral prostate was subjected to histological examination. Statistical analyses included
31 ANOVA, Dunnet *t*-test for multiple comparison, and Fisher exact probability test.

32
33 On PND 29, serum prolactin levels were significantly increased by ~210% in rats of the bisphenol A
34 group compared to the control group. On PND 120, there was no effect on prolactin levels in serum or
35 pituitary in the bisphenol A group. Ventral prostate weight was unaffected but lateral prostate weight was
36 increased **[by ~25%]** in the bisphenol A group. Exposure to bisphenol A had no effect on body or testis
37 weight. **[Data were not shown by study authors.]** The myeloperoxidase assay was reported to show a
38 “trend” for lateral prostate inflammation in the bisphenol A group. **[Trend was not defined; there was
39 no statistical difference between the bisphenol A group and the control in the myeloperoxidase
40 assay.]** No histological evidence of inflammation was observed in prostates from the control group. In the
41 bisphenol A group, histopathological analyses revealed that 44.4% of prostates contained increased a
42 focal luminal polymorphonuclear cellular infiltrate that was milder in severity compared to prostates from
43 the 17 β -estradiol group. The study authors noted the discrepancy between the results obtained by
44 myeloperoxidase assay and histological observation in the bisphenol A group and stated that the
45 discrepancy may have been due to evaluation of the whole tissue by myeloperoxidase assay versus only
46 one section of the tissue by histological evaluation. Bisphenol A had no effect on prostate DNA content.
47 In addition to prostate inflammation, effects observed in the 17 β -estradiol group were increased serum
48 prolactin levels on PND 29 and elevated myeloperoxidase and DNA content in lateral prostate on PND
49 120. Based on these findings, the study authors concluded that chemically induced, transient increases in
50 prolactin secretion in the prepubertal period can lead to increased incidence of lateral prostate
51 inflammation in 120-day-old rats.

3.0 Developmental Toxicity

1
2 **Strengths/Weaknesses:** Strengths include appropriately performed experiments and dependable data.
3 Comparison with other agents is also a strength. A weakness is that links between the prostatic changes
4 and prolactin levels were not definitive; although, there is certainly good evidence to suggest a link. The
5 single dose level of bisphenol A used in this study to some extent limits its applicability.
6

7 **Utility (Adequacy) for CERHR Evaluation Process:** This well performed study is suitable for inclusion
8 in this review.
9

10 **Atanassova et al. (327)**, supported by the European Center for the Ecotoxicology of Chemicals and
11 AstraZeneca, examined the effects of neonatal bisphenol A exposure on the reproductive system of male
12 rats. Wistar rats were fed rat and mouse breeding diet No. 3, which contains 15.5% soy meal flour. **[No**
13 **information was provided about caging and bedding materials.]** Litters of 8–12 male rats were
14 assembled by cross-fostering pups on PND 1 (day of birth). On PND 2–12, rats were sc injected with corn
15 oil vehicle or bisphenol A **[purity not given]** 0.5 mg/day. **[Assuming a 5–25 g body weight during this**
16 **interval, this dose would be ~100 mg/kg bw/day at the beginning of the interval and ~20 mg/kg**
17 **bw/day at the end of the interval.]** Other groups of rats were sc injected with 0.01–10 µg
18 diethylstilbestrol every other day between PND 2 and 12 or 2 mg 4-tert-octylphenol/day during PND 2–
19 12. Rats were killed on PND 18, 25, and 90–100. At PND 18 and 25, testes were weighed and fixed in
20 Bouin solution. Testicular cell numbers and seminiferous tubule lumen formation were determined by
21 standard point counting of cell nuclei. Apoptosis was assessed by DNA fragmentation detected by in situ
22 DNA 3'-end labeling. Spermatocyte nuclear volume as a fraction of Sertoli cell nuclear volume was
23 calculated as “an index of spermatogenic efficiency.” Plasma FSH and inhibin B were measured by RIA
24 and ELISA methods, respectively. Fertility was assessed at 80–90 days of age; rats were mated for 7 days
25 and number of pups was counted at birth. The number of rats/group examined was 7–14 at 18 days of age,
26 4–12 at 25 days of age, and 6 in fertility testing. Data were analyzed by ANOVA.
27

28 Significant effects observed on PND 18 were advanced testicular lumen formation and increases in testis
29 weight, Sertoli cell volume/testis, and spermatocyte nuclear volume/unit Sertoli cell. A decrease in germ
30 cell apoptosis was also described on PND 18 but was not statistically significant. Plasma FSH levels were
31 significantly increased on PND 18, but there was no effect on plasma inhibin B concentration. The only
32 significant effect observed on PND 25 was increased plasma FSH levels. Testis weight was increased in
33 adulthood, but there were no effects on fertility or litter size. Effects observed with octylphenol were
34 similar to those observed with bisphenol A. In contrast, exposure to one or more doses of
35 diethylstilbestrol resulted in increased apoptosis, decreased plasma inhibin levels, decreased Sertoli cell
36 nuclear volume, and changes in spermatocyte/Sertoli cell ratios. The study authors concluded that the
37 effect of bisphenol A on spermatogenic processes is benign.
38

39 **Strengths/Weaknesses:** Strengths include comparisons of bisphenol A with other estrogenic compounds
40 on the endpoints tested. The experiments and data recording were apparently good as was the
41 interpretation of the data. This group has extensive expertise in testis biology and male fertility in general,
42 therefore they are likely to identify even subtle problems. A significant weakness is that only 1 dose level
43 of bisphenol A was used and this dose level was variable on a weight basis, although always very high
44 because it was a set mass of drug per day applied to a growing animal and was not adjusted to body
45 weight.
46

47 **Utility (Adequacy) for CERHR Evaluation Process:** This work is suitable for the evaluation process.
48

49 **Williams et al. (328)**, supported by the European Centre for Ecotoxicology, examined the effect of
50 neonatal bisphenol A exposure on seminal vesicle structure and expression of sex steroid receptors in rats.
51 On PND 2 (day of birth = PND 1), litters consisting of 8–14 male Wistar rat pups were derived through

3.0 Developmental Toxicity

1 cross-fostering. Rats were sc injected with corn oil vehicle or 0.5 mg/day bisphenol A on PND 2–12.
2 **[Assuming a 5–25 g body weight during this interval, the dose would be ~100 mg/kg/day at the**
3 **beginning of the interval and ~20 mg/kg bw/day at the end of the interval.]** The dose was based on
4 the highest amount that could remain in solution. A positive control group was injected with
5 diethylstilbestrol at 0.1, 1, or 10 µg/day on PND 2, 4, 6, 8, 10, and 12. Ethinyl estradiol was administered
6 at 10 µg/day, according to the protocol for diethylstilbestrol. Control animals for each compound were
7 dosed with vehicle on the appropriate days, and because no differences were noted for controls, data were
8 pooled. The effects of 4-tert-octylphenol, genistein, Antarelix, flutamide, and tamoxifen were also
9 examined but will not be discussed. **[No information was provided about feed, caging or bedding**
10 **materials, or purity of compounds.]** Animals were killed on PND 18, and seminal vesicles from 11–15
11 animals/group were collected and stored in Bouin solution. Seminal vesicles were examined for gross
12 abnormalities in stroma and epithelium. Immunolocalization studies were conducted to assess ERβ, ERα,
13 androgen receptor, and progesterone receptor proteins in the seminal vesicle. Studies were replicated 3–5
14 times using samples from at least 6 animals/group. Results were subjectively scored.

15
16 The gross structure of the seminal vesicles from bisphenol A-treated rats appeared normal, and there were
17 no changes in ERβ, ERα, androgen receptor, or progesterone receptor proteins in the seminal vesicle. In
18 contrast, diethylstilbestrol induced changes in seminal vesicle morphology, increased ERα and
19 progesterone receptor, and decreased androgen receptor. Effects of ethinyl estradiol were similar to those
20 observed with diethylstilbestrol. The study authors concluded that the lack of bisphenol A effects
21 suggested that only high doses of potent estrogens induce gross abnormalities in the male reproductive
22 system; and that only agents that suppress androgen receptor while increasing ERα and progesterone
23 receptor are likely to cause gross developmental abnormalities in the male reproductive system.

24
25 **Strengths/Weaknesses:** Strengths include expertise of the group coupled to well-performed experiments,
26 data recording, and interpretation. However, a significant weakness is that only one dose level of
27 bisphenol A was used and this dose level was variable, although always very high because it was a set
28 mass per day given to a growing animal. Bisphenol A was not a primary target in this study but was one
29 of a series of estrogenic compounds, allowing comparison with other similar compounds. The target of
30 this investigation was the seminal vesicle which is not a major disease site; however, the focus of the
31 work is on appropriate expression of sex steroid receptors and therefore this organ can be considered to be
32 a good reporter for the male genital tract.

33
34 **Utility (Adequacy) for CERHR Evaluation:** This work is suitable for the evaluative process. Data
35 presented do not give rise for concern in relation to these endpoints.

36
37 **Rivas et al. (329)**, supported by the European Union and the Spanish Ministry of Education, examined
38 the effects of bisphenol A exposure on reproductive tract development of male rats. The main focus of the
39 study was determining the effects of decreased androgen production in combination with a low dose of
40 diethylstilbestrol. Effects of flutamide were also examined but will not be discussed. Wistar rats were fed
41 a soy-free diet (rat and mouse soya-free breeding diet, SDS, Dundee, Scotland). **[No information was**
42 **provided about caging and bedding materials.]** Litters of 8–12 male pups were assembled by cross-
43 fostering on PND 1 (day of birth). Male rats were sc injected with the corn oil vehicle or 0.1 mg bisphenol
44 A on PND 2, 4, 6, 8, 10, and 12 with and without coadministration of 10 mg/kg GnRH antagonist (a
45 suppressor of androgen production). **[Assuming a 5–25 g body weight during this interval, the**
46 **bisphenol A dose would be ~20 mg/kg bw/day at the beginning of the interval and ~4 mg/kg bw/day**
47 **at the end of the interval.]** Additional rats were sc injected with diethylstilbestrol at doses of 0.1 or 10
48 µg on PND 2, 4, 6, 8, 10, and 12 with and without administration of GnRH antagonist. Rats were killed
49 on PND 15. The testis was fixed in Bouin solution and testicular structures were measured. Plasma
50 testosterone levels were measured using an ELISA technique. From 3 to 10 animals/group were examined
51 for each endpoint. Data were analyzed by ANOVA.

3.0 Developmental Toxicity

1
2 Treatment with bisphenol A alone did not affect plasma testosterone levels but treatment with GnRH
3 antagonist alone and in combination with bisphenol A significantly lowered plasma testosterone levels.
4 Treatment of rats with bisphenol A alone or in combination with GnRH antagonist had no significant
5 effect on rete testis luminal area, efferent duct luminal area, efferent duct epithelial cell height, or vas
6 deferens epithelial cell height. Exposure to the high diethylstilbestrol dose increased rete area, and both
7 doses of diethylstilbestrol decreased plasma testosterone levels, increased efferent duct luminal area, and
8 decreased epithelial cell height in efferent duct and vas deferens. The study authors concluded that the
9 estrogenicity of bisphenol A when injected at a moderately high dose was insufficient for disrupting the
10 estrogen-androgen balance in rats.

11
12 **Strengths/Weaknesses:** This study was carefully performed and well documented. A minor weakness for
13 this evaluation is that bisphenol A was not the primary target of the work. The dose used was high and
14 only a single dose level was examined. That being said, the data generated can be considered reliable. The
15 testes and the Wolffian duct derivative structures are reasonable targets for estrogenic chemicals and are,
16 therefore, logical choices to examine.

17
18 **Utility (Adequacy) for CERHR Evaluation Process:** The data presented are suitable for the evaluation
19 process.

20
21 **Sharpe et al. (330)**, supported in part by the European Union and the Spanish Ministry of Education,
22 examined the effects of neonatal exposure of rats to bisphenol A on Leydig cell development and
23 function. Wistar rat dams were fed a standard soy-containing feed (rat and mouse breeding diet, SDS,
24 Dundee, UK). [No information was provided on feed given to male offspring following weaning or
25 bedding and caging materials.] Litters of 9–12 male pups were created by cross fostering pups on PND
26 1 (day of birth). Male pups were sc injected with the corn oil vehicle or 0.5 mg/day bisphenol A [purity
27 not reported] on PND 2–12. [Assuming 5–25 g body weight during this interval, the dose would be
28 ~100 mg/kg bw/day at the beginning of the interval and ~20 mg/kg bw/day at the end of the
29 interval.] Other groups of rats received diethylstilbestrol at 0.1–10 µg/day on PND 2, 4, 6, 8, 10, and 12.
30 Additional rats were treated with GnRH antagonist Antarelix or 4-tert-octylphenol, but those results will
31 not be discussed. Rats were killed on PND 18, 25, 35, or 90. Testes were weighed and fixed in Bouin
32 solution. Sections of testes were immunostained with the Leydig cell marker 3β-hydroxysteroid
33 dehydrogenase to evaluate Leydig cell development in 5–7 animals/group. Plasma testosterone levels
34 were measured by ELISA. Group sizes for evaluation of testes weight and plasma testosterone were 2–23,
35 with most groups containing at least 8 animals. Data were analyzed by ANOVA.

36
37 The only significant effect on plasma testosterone level following exposure to bisphenol A was an
38 increase on PND 18. In rats of the bisphenol A group examined at each time period, there were no
39 significant effects on testis weight, percent Leydig cell nuclear volume/testis, Leydig cell nuclear
40 volume/testis, or total Leydig cell volume (nuclear + cytoplasmic volume/testis). Significant results in rats
41 exposed to diethylstilbestrol included decreased Leydig nuclear cell volume at the mid and/or high dose
42 on or before PND 35 and reduced plasma testosterone level and testis weight at all doses and most time
43 points of evaluation. The study authors concluded that there were no consistent changes in Leydig cell
44 development following exposure to bisphenol A.

45
46 **Strengths/Weaknesses:** This paper reports a well performed and documented study. Limitations include
47 use of a single high but variable dose of bisphenol A. A strength is that bisphenol A was one of a number
48 of compounds examined enabling internal comparison with other similar molecules. Confidence in the
49 results is high.

50
51 **Utility (Adequacy) for CERHR Evaluation Process:** This study is suitable for the evaluation process.

3.0 Developmental Toxicity

1
2 **Khurana et al. (331)**, supported by NIH, March of Dimes, and Pardee Foundation, examined the effects
3 of neonatal bisphenol A exposure on prolactin levels in rats. [**The type of chow used and composition of**
4 **bedding and caging materials were not reported.**] On PND 1–5 (day of birth = PND 0), 8–10 Fischer
5 344 rat pups/sex/group were sc injected with the tocopherol-stripped corn oil vehicle, bisphenol A at 0.1
6 or 0.5 mg/day, diethylstilbestrol at 5 µg/day, or octylphenol at 0.1 or 0.5 mg/day. [**Assuming a pup body**
7 **weight of 5 g, bisphenol A intakes were estimated at 20 and 100 mg/kg bw/day.**] On PND 15, 20, and
8 25, blood was collected for measurement of serum prolactin level by RIA. A final sample for prolactin
9 analysis was obtained when animals were killed on PND 30. Medial basal hypothalamus, anterior
10 pituitary, uterus, and prostate were collected for measurement of *ERα* and *ERβ* mRNA expression by RT-
11 PCR in animals of the low-dose group. Statistical analyses included ANOVA followed by Student-
12 Newman-Keuls test.

13
14 In male and female rats, hyperprolactemia was observed on PND 25 and 30. [**On PND 30, prolactin**
15 **levels in the low- and high-dose groups compared to the control group were ~ 150 and 95% higher**
16 **in females and 120 and 80% higher in males**]. In females exposed to the low dose, *ERα* mRNA in the
17 medial basal hypothalamus was higher [**by 25%**] than control levels. In anterior pituitary of low-dose
18 males, *ERα* mRNA was higher [**by ~80%**] and *ERβ* mRNA was higher by 35–40% compared to control
19 levels. There were no effects on *ERβ* mRNA in female tissues. Most effects observed with octylphenol
20 exposure were similar to those observed with bisphenol A exposure. Diethylstilbestrol induced transient
21 increases in prolactin levels, decreased expression of *ERα* in medial basal hypothalamus of males,
22 upregulated *ERα* and *ERβ* expression in the pituitary of males, decreased expression of *ERα* in the uterus,
23 and upregulated *ERβ* expression in prostate. The study authors concluded that exposure of neonatal rats to
24 bisphenol A resulted in delayed and sustained hyperprolactemia and changes in *ER* mRNA expression.

25
26 **Strengths/Weaknesses:** This paper describes well performed and documented work. One of the strengths
27 of the paper is the use of two moderate dose levels of bisphenol A to assess effects. Another strength is
28 that both male and female animals were assessed. The fact that the lower of the two doses often produced
29 more marked effects than the higher dose is important. The observation of changes in receptor status in
30 the hypothalamus is consistent with studies linking brain structure and organismal behavior in relation to
31 this compound.

32
33 **Utility (Adequacy) for CERHR Evaluation Process:** This paper is suitable for the evaluation process.
34 The data raise concerns that bisphenol A exposure can have effects on the endpoints examined.

35
36 **Fukumori et al. (332)**, support not indicated, examined the effect of postnatal bisphenol A exposure on
37 ultrastructure of the prostate in rats. [**The study was published in Japanese; a translation was**
38 **provided by the American Plastics Council.**] On day 1–21 following birth, F344 rats were sc injected
39 with bisphenol A 5 days/week at doses of 0 (DMSO vehicle), 0.0008, 0.004, 0.020, and 0.500 mg/kg
40 bw/day. A positive control group received 100 µg/kg bw 17β-estradiol by sc injection during the same
41 time period. Rats were killed at 22 days of age. Ventral prostates were fixed in glutaraldehyde, sectioned,
42 and examined by electron microscopy. [**The number of rats treated and examined/group and the**
43 **number of litters represented were not reported. No information was provided on purity of**
44 **bisphenol A, type of feed, or composition of bedding and caging. The translated version of the**
45 **report did not include figures from the original report.**] In ventral prostates obtained from rats
46 exposed to 17β-estradiol, there was an increase in secretory granules accompanied by reductions in
47 microvilli on the surface of the glandular epithelium. Proliferation of fibroblasts was observed in the
48 fibromuscular layer of the stroma in rats from the 17β-estradiol group. In the 0.020 and 0.500 mg/kg
49 bw/day bisphenol A groups, a slight increase in secretory granules and slight decrease in microvilli was
50 observed in glandular epithelium. Effects in stroma were described as unremarkable for the bisphenol A

3.0 Developmental Toxicity

1 groups. The study authors concluded that bisphenol A may have ultrastructural effects on the ventral
2 prostates of suckling rats.

3
4 **Strengths/Weaknesses:** This is a translation of an apparently carefully performed study to assess the
5 effects of low doses of perinatal bisphenol A on prostatic structure. A major weakness is that the original
6 figures were not provided, so one is left to extract data from the text without pictures for comparison. The
7 young age at which the animals were sacrificed is also a concern because prostatic development is not
8 complete at 22 days of age making comparisons with the bulk of established data problematic. It may be
9 that some of the effects observed simply reflect mild retardation of development in the treated animals,
10 which would be corrected with the passage of time.

11
12 **Utility (Adequacy) for CERHR Evaluation Process:** This paper is suitable for inclusion with the
13 proviso that this is a translation provided by a group with a financial interest in this evaluation process.
14 Also, the primary data, in the form of figures, were not available for inspection.

15
16 **Kato et al. (333)**, supported by the Japanese Ministry of Education, Culture, Sports, Science, and
17 Technology and the Ministry of Health, Labor, and Welfare, examined the effects of neonatal bisphenol A
18 exposure on the reproductive organs of rats. Sprague Dawley rats were fed CRF-1 diet. **[No information**
19 **was provided on caging or bedding materials.]** Female offspring from 8 dams were grouped to achieve
20 equal distribution of body weight. At least 8 female offspring/group were sc injected with 0 (ethanol/corn
21 oil vehicle), 0.25, 1, or 4 mg/day bisphenol A **[purity not reported]** from PND 0 to 9 (day of delivery =
22 PND 0). **[Based on body weights reported on PND 0 and 9, CERHR calculated mean bisphenol A**
23 **intakes of ~26, 105, and 427 mg/kg bw/day.]** A positive control group was given 10 µg/day 17β-
24 estradiol **[~3 mg/kg bw/day]** during the same time period. Rats were weighed during and following the
25 lactation period and examined for day of vaginal opening. External reproductive organs were examined
26 on PND 60, and estrous cycles were assessed from PND 61 to 94. One group of rats was ovariectomized
27 on PND 80; ovaries were weighed, and fixed in 10% neutral buffered formalin for evaluation of corpora
28 lutea and polyovular follicles. Another group of bisphenol A-exposed and the vehicle-treated control
29 females were given 1 µg/kg 17β-estradiol from PND 94 to 96 and killed the day following final injection;
30 uterus and vagina were weighed, and fixed in 10% formalin. For all endpoints, 5–8 rats/group were
31 examined. Statistical analyses included Student *t*-test and Fisher exact probability test.

32
33 Treatment-related results are summarized in Table 79. Two rats of the high-dose group died. Body
34 weights of rats in the high-dose group were lower than controls on PND 9–30 but higher than controls on
35 PND 61–97. Effects observed at the mid and high dose included accelerated vaginal opening, increased
36 incidence of polycystic ovaries, decreased area of corpora lutea, and decreased uterine fluid weight. All
37 rats of the mid-dose group had partial clefts in the clitoris, and all rats of the high-dose group had deep
38 clefts in the clitoris. Additional effects observed in rats of the high-dose group included disrupted estrous
39 cycles (e.g., irregular cycles or persistent estrous) and decreased relative (to body weight) ovary and wet
40 or blotted uterus weights. Absolute weights of wet uterus and ovary were also reduced in the high-dose
41 group. No corpora lutea were observed in rats of the high-dose group. Qualitatively similar effects were
42 observed in the group treated with 17β-estradiol. The study authors concluded that exposure of rats to
43 bisphenol A during the neonatal period resulted in changes in female reproductive organs.

3.0 Developmental Toxicity

1 **Table 79. Effects in Female Rats Exposed to Bisphenol A During the Neonatal Period**

Endpoint	Dose, mg/kg bw/day [CERHR estimate]						
	26	105	427	BMD ₁₀	BMDL ₁₀	BMD _{1SD}	BMDL _{1SD}
Body weight gain							
PND 9	↔	↔	↓16%	286	200	233	156
PND 97	↔	↔	↑10%	432	261	430	253
Day of vaginal opening	↔	↓2.9 days	↓4.1 days	345	267	159	116
No. with normal estrous cycles ^a	↔ (8/8)	↔ (2/8)	↓ (0/6)	81	28		
No. with cleft clitoris ^b	↔ (0/8)	↑ (0/8)	↑ (6/6)	299	failed		
Relative organ weight							
Ovary	↔	↔	↓59%	85	59	140	93
Uterus, wet	↔	↔	↓60%	66	55	128	96
Uterus, blotted	↔	↔	↓21%	273	128	318	168
Uterine fluid weight	↔	↓42%	↓97%	42	34	139	104
No. with polycystic ovaries ^b	No data	↑ (4/8)	↑ (5/5)	81	24		
No. with corpora lutea ^a	No data	↔ (8/8)	↓ (0/5)	238	90		
No. of corpora lutea	No data	↔	↓ (none)	65	38	137	83
Corpora lutea area	No data	↓ 30%	↓ (none)	42	37	84	66

↑,↓ Statistically significant increase compared to controls; ↔ no statistically significant effect.

^aControl rate 8/8.

^bControl rate 0/8.

From Kato et al. (333).

2
3 **Strengths/Weaknesses:** The strengths are the carefully performed and documented experiments. The
4 major limitation is that the doses of bisphenol A used were massive. The changes in the female
5 reproductive organs seen are well documented, but given the extremely high dose of agent used, broadly
6 unsurprising.

7
8 **Utility (Adequacy) for CERHR Evaluation Process:** The results of this study reflect a careful
9 documentation of the experiments performed, so the study is relevant and useful for the evaluation
10 process.

11
12 **Toyama and Yuasa (334)**, supported in part by the Japanese Ministry of Environment and Ministry of
13 Education, Science, Sports and Culture, examined the effects of neonatal bisphenol A [**purity not**
14 **reported**] exposure on spermatogenesis during puberty and adulthood in rats and mice. [**No information**
15 **was provided about chow or bedding and caging materials. The mouse data are reported in Section**
16 **3.2.8.**] Wistar rats were sc injected with bisphenol A in a DMSO and olive oil vehicle on PND 1, 3, 5, 7,
17 9, and 11 (PND 0 = day of birth). Bisphenol A doses were 0.001, 0.010, 0.100, and 0.600 mg/kg bw in
18 rats. Additional animals were treated with 17β-estradiol and estradiol benzoate. Animals were killed
19 weekly at 2–10 weeks of age, and other pups were killed at 24 and 31 days of age. There were 5
20 animals/dose/time point in bisphenol A groups and apparently 5 vehicle control rats/time period. Testes
21 were examined by light and electron microscopy. Males from each experimental group (a total of 11 rats)
22 were mated with 2 females [**number tested in each dose group not reported**]. A total of 11 rat dams
23 were allowed to complete pregnancy. [**It does not appear that statistical analyses were conducted.**]

24
25 All rats treated given 0.600 mg/kg bw bisphenol A died before 20 days of age and were excluded from
26 analysis. In mature spermatids of 8-week-old rats in the vehicle control group, the incidences of deformed
27 acrosome, deformed nucleus, and abnormal ectoplasmic specialization were below 0.3%. In 8-week-old
28 rats treated with ≥0.010 mg/kg bw bisphenol A, the incidence of deformed acrosome was >50–60%, the
29 incidence of deformed nucleus was >40%, and the incidence of abnormal ectoplasmic specialization was
30 >60–70%. [**Data were not shown for individual dose levels.**] Similar effects were observed in the

3.0 Developmental Toxicity

1 groups treated with 17 β -estradiol and estradiol benzoate. No effects were reported at other ages. **[Data**
2 **were not shown by study authors.]** The blood-testis barrier remained intact based on histologic
3 observations. All tested males from the bisphenol A group were fertile, and sex ratio, litter sizes, and pup
4 weights were reported to be normal. **[No results were shown for individual dose levels. Fertility data**
5 **presented in Table 4 and 5 of the study, were not clearly identified by dose level.]** The study authors
6 concluded that bisphenol A acts as an estrogen and induces transient changes in the male reproductive
7 system of rodents that resolve in adulthood.

8
9 **Strengths/Weaknesses:** This study appears to have been well performed and documented. The strengths
10 include the use of multiple doses of bisphenol A and the use of both rats and mice, allowing interspecies
11 comparisons. Weaknesses include selective data presentation and failure to examine sperm morphology in
12 the fertile 15 week old animals to determine whether the changes in sperm maturation seen at earlier time
13 points had resolved or whether the animals were fertile in the face of such abnormalities.

14 **Utility (adequacy) for CERHR Evaluation Process:** This study is suitable for evaluation and shows
15 that high perinatal doses of bisphenol A result in toxicity notable in rats but not in mice given the same
16 dose of agent.

17
18 **Kato et al. (335)**, supported by the Japanese Ministry of Education, Culture, Sports, Science and
19 Technology and Ministry of Health, Labor and Welfare, examined the effects of neonatal exposure to
20 bisphenol A on reproductive function of male rats. Sprague Dawley rats were fed CRF-1 diet, which was
21 described as having relatively low estrogenic activity compared to other Japanese rodent feeds. **[No**
22 **information was provided on caging or bedding materials.]** Male rats used in this study were born to
23 12 dams, assigned to 8 foster dams in groups of 7 based upon body weights, and distributed to dose
24 groups. From PND 0 to 9 (PND 0 = day of birth), 24 male pups/group were sc injected with bisphenol A
25 at 0 (ethanol/corn oil vehicle), 0.000024, 0.000120, 0.000600, 0.003, or 1 mg/day bisphenol A. Study
26 authors calculated average exposures of 0.002, 0.011, 0.056, 0.277, or 97 mg/kg bw/day. An additional
27 group was treated with 10 μ g/day 17 β -estradiol (0.9 mg/kg bw/day) during the same time period. Eight
28 rats/group were killed and necropsied at PND 10, 35, and 150. At the PND 10 necropsy, serum
29 testosterone levels were measured by RIA, the testis was weighed and examined histologically, and
30 expression changes in genes for hormone receptors and steroidogenic enzymes were determined by RT-
31 PCR. The same endpoints were examined at the PND 35 necropsy in addition to measuring seminal
32 vesicle, ventral prostate, and epididymis weights. The remaining rats were assessed for day of preputial
33 separation. From PND 105 to 130, they were mated for 1 day a maximum of 4 times with an untreated
34 female in proestrus. Females were killed on GD 13 (day of sperm = GD 0) and examined for corpora
35 lutea, embryonic mortality, and implantation sites. Male rats were killed on PND 150. In addition to
36 endpoints examined at earlier time periods, sperm endpoints and histopathology of ventral prostate were
37 assessed. Statistical analyses included Bartlett method for homogeneity of variance followed by Dunnett
38 method for homogeneous variances or Dunnett-type method with rank order for heterogeneous variances.
39 Reproductive data were analyzed by Fisher exact probability test. Data obtained from the 17 β -estradiol
40 group were analyzed by Student *t*-test.

41
42 There were no deaths or decreases in body weight in animals of the bisphenol A group. There were no
43 effects on age of preputial separation, copulation rate, or fertility. In dams impregnated by bisphenol A-
44 treated males, there were no effects on numbers of implantation sites, implantation losses, or live fetuses.
45 Bisphenol A treatment had no adverse effects on sperm count, motility, or morphology. There were no
46 effects on serum testosterone levels, histopathology of testis or prostate, or weights of testis, epididymis,
47 seminal vesicle, ventral prostate, or penis. No significant changes were observed in mRNA for estrogen,
48 androgen, or progesterone receptor or steroidogenic enzymes. In contrast to the bisphenol A groups, rats
49 treated with 17 β -estradiol experienced decreases in reproductive organ weights, altered gene expression,
50 delayed and incomplete preputial separation, decreased copulatory rate, and decreased sperm numbers.

3.0 Developmental Toxicity

1 The study authors concluded that neonatal bisphenol A exposure caused no adverse effects on
2 reproductive function or gene expression of steroidogenic enzymes in the rat testis.

3
4 **Strengths/Weaknesses:** This paper has a number of major strengths, notably the wide range of levels of
5 bisphenol A administered allowing a good picture of the effects of these doses on immediate postnatal
6 male animals. The endpoints measured are all very relevant to the overall topic. As always, one could
7 identify as a weakness a number of endpoints that were not determined, however on balance this is a
8 useful document.

9
10 **Utility (Adequacy) for CERHR Evaluation Process:** This paper is suitable for evaluation and provides
11 a well documented data set that suggests no cause for concern about early postnatal exposure to bisphenol
12 A in these concentrations in relation to the criteria examined.

13
14 **Ho et al. (336)**, supported by NIH and Department of Defense, examined the effect of developmental
15 exposure to bisphenol A on susceptibility of Sprague Dawley rats to prostate cancer. The dams and
16 offspring used in this study were fed a soybean-free phytoestrogen-reduced diet (Zeigler Reduced Rodent
17 Diet 2, Ziegler Brothers, Inc), housed in polysulfone cages [**with unspecified bedding**], and provided
18 drinking water in glass bottles. On PND 1, 3, and 5 (day of birth = PND 0), 20–30 male pups/group were
19 sc injected with tocopherol-stripped corn oil vehicle, bisphenol A at 0.1 µg/pup (0.010 mg/kg bw), or
20 estradiol benzoate at 0.001 µg/pup (0.1 µg/kg bw) or 25 µg/pup (2500 µg/kg bw). Male rats from each
21 litter were randomly assigned to treatment groups, but the total number of litters from which the pups
22 were selected was not reported. Pups were weaned on PND 21. At PND 90, half the rats from each
23 treatment group were implanted with Silastic capsules containing 17β-estradiol and testosterone and the
24 other half were implanted with empty capsules; the capsules were left in place for 16 weeks. The
25 treatment was designed to result in a serum 17β-estradiol level of ~75 ng/L and testosterone level of ~3
26 µg/L, levels reported to induce prostatic intraepithelial neoplasia in 33% of Sprague Dawley rats. Rats
27 were killed at 28 weeks of age. Prostates were removed, and histopathological evaluations were
28 conducted on each lobe. Immunohistochemistry techniques were used to measure proliferation. Apoptosis
29 was measured using the terminal deoxynucleotidyl transferase-mediated dUTP nick-end labeling
30 (TUNEL) technique. PCR techniques were used to study methylation pattern and expression changes in
31 prostate cell signaling proteins on PND 10, 90, and 200. Statistical analyses included chi-squared test,
32 ANOVA, Fisher exact test, and Bonferroni test.

33
34 The study authors stated that similar responses were observed in each of the 3 prostate lobes; and thus
35 results were presented only for dorsal prostate. In bisphenol A-exposed compared to vehicle controls rats
36 that did not receive 17β-estradiol/testosterone exposure in adulthood, there were no effects on dorsal
37 prostate weight, histopathology alterations, proliferation index, or apoptotic index. In bisphenol A-treated
38 compared to vehicle control rats that received 17β-estradiol/testosterone exposure in adulthood, there was
39 increased incidence and severity of prostatic intraepithelial neoplasia (100 vs. 40% incidence), a precursor
40 lesion to prostate cancer. In the bisphenol A/17β-estradiol/testosterone group, proliferation and apoptosis
41 indices were increased in regions where prostatic intraepithelial neoplasia was observed. Changes
42 observed in rats exposed to the high estradiol benzoate dose in the neonatal period but not 17β-
43 estradiol/testosterone during adulthood included increased incidence and severity of prostatic
44 intraepithelial neoplasia and elevated apoptosis and proliferation indices. The same effects, in addition to
45 decreased prostate weight, were observed in rats receiving neonatal exposure to the high estradiol
46 benzoate dose and adult exposure to 17β-estradiol/testosterone.

47
48 In the investigation of a molecular basis for increased susceptibility to prostate cancer, exposure to
49 estrogenic compounds altered methylation pattern in several cell signaling genes. Phosphodiesterase type
50 4 variant, an enzyme involved in cyclic AMP breakdown, was selected for further study. Neonatal

3.0 Developmental Toxicity

1 bisphenol A exposure resulted in hypomethylation of the phosphodiesterase type 4 variant gene and
2 increased expression of that gene at 90 and 200 days of age, with or without 17 β -estradiol/testosterone
3 exposure in adulthood. Similar responses in phosphodiesterase type 4 variant gene methylation and
4 expression were observed with exposure to the low and high 17 β estradiol benzoate doses. The study
5 authors concluded that developmental exposures of rats to bisphenol A increased susceptibility to
6 precancerous prostate lesions resulting from prostate epigenome alteration.

7
8 **Strengths/Weaknesses:** This is a carefully performed study by a group with significant expertise in this
9 area of work. The paper has many strengths, from the use of multiple, biologically relevant doses of
10 bisphenol A to the search to identify molecular mechanisms underlying the observations made. This is
11 one of a very limited number of studies that have been carried out to look at the longer term consequences
12 of neonatal bisphenol A exposure. It could be suggested that carrying the study further in terms of animal
13 age might have produced more dramatic phenotypes, and failure to do this could be considered a
14 weakness of the work.

15
16 **Utility (Adequacy) for CERHR Evaluation Process:** This paper makes important contributions and is
17 suitable for the evaluation process.

18 19 3.2.4.2 Neurobehavioral endpoints

20 **Ishido et al. (337)**, supported by the National Institute for Environmental Studies and the Ministry of
21 Economy, Trade, and Industry, examined the effects of postnatal intracisternal bisphenol A exposure on
22 behavior of rats. Dams in this study were fed Standard laboratory chow (MF Diet; Oriental Yeast Corp.).
23 **[No information was provided about caging or bedding materials.]** At 5 days of age, 5–7 male Wistar
24 rat pups/group were injected intracisternally with a bisphenol A dose of 0 (ethanol/olive oil vehicle),
25 0.00002, 0.0002, 0.002, or 0.020 mg. Pups were weaned at 3 weeks of age. Spontaneous motor activity
26 was measured over a 12–24-hour period at 4–5 weeks of age. Rats were killed at 4 and 8 weeks of age,
27 and brains were removed. RNA was isolated from midbrain and striatum for DNA microarray analysis.
28 Expression of the gene for dopamine transporter in midbrain was studied by RT-PCR. Tyrosine
29 hydroxylase expression in brain was measured at 8 weeks of age using an immunostaining method.
30 Statistical analyses included ANOVA and Student *t*-test.

31
32 In 4–5-week-old rats from the 0.020 mg bisphenol A group, motor activity was significantly increased
33 and was 1.6 times higher than in control rats during the nocturnal period. In a dose response experiment,
34 it was noted that hyperactivity was significantly increased at doses ≥ 0.0002 mg. Microarray analysis
35 revealed that bisphenol A **[at an unspecified dose]** downregulated expression of dopamine D4 receptor
36 gene 2-fold at 4 weeks of age and dopamine transporter gene 2.8-fold at 8 weeks of age. Numerous other
37 gene expression changes were observed but not discussed in detail by study authors. Analysis by RT-PCR
38 confirmed that expression of the dopamine transporter gene was downregulated 3-fold in the midbrain of
39 8-week-old rats treated with bisphenol A in the neonatal period. In rats from the 0.020 mg bisphenol A
40 group, tyrosine hydroxylase immunoreactivity was reduced in the substantia nigra at 8 weeks of age. The
41 study authors interpreted the decrease in tyrosine hydroxylase immunoreactivity as degeneration of
42 dopaminergic neurons. They concluded that bisphenol A affected the central dopaminergic system,
43 resulting in hyperactivity that most likely occurred as a result of decreased tyrosine hydroxylase activity
44 in midbrain.

45
46 **Strengths/Weaknesses:** Strengths of this paper include the use of a range of concentrations of bisphenol
47 A. The correlation of changes in behavior patterns induced by bisphenol A with expression of specific
48 dopamine receptor sets is also a strength. A significant weakness is the inability to correlate the doses of
49 bisphenol A provided by this dosing mechanism with those seen by more common sc or oral routes, as
50 well as uncertainty about the disposition of the bisphenol A that is injected into the cerebrospinal fluid.
51 The behavioral data provided are stronger and more convincing than the somewhat cursory molecular

3.0 Developmental Toxicity

1 study, where a single, probably relevant, receptor was chosen from a microarray study and subjected to
2 minimal publishable follow-up.

3
4 **Utility (Adequacy) for CERHR Evaluation Process:** This paper is suitable for the evaluation process.

5
6 **Patisaul et al. (338)**, supported by the American Chemistry Council, evaluated the effect of neonatal
7 bisphenol A on the anteroventral periventricular nucleus of the Sprague Dawley rat. Pregnant rats (n = 5)
8 were fed a phytoestrogen-free diet (Purina 5K96) during the last week of gestation. **[No information was
9 provided about caging or bedding.]** Dams were permitted to litter. Pups were cross-fostered among all
10 dams so that 4 dams reared 6 females and 6 males and 1 dam reared 5 males. Pups (n = 5–8/group) were
11 randomly assigned to receive sc injections of 17β-estradiol 50 μg/pup, genistein 250 μg/pup, bisphenol A
12 250 μg/pup, or sesame oil vehicle every 12 hours for 48 hours. The authors estimated that the twice daily
13 dosing with 250 μg/pup was approximately equivalent to 100 mg/kg bw/day. Injections began the
14 morning of PND 1 (delivery = PND 0). On PND 19, the pups were transcardially perfused with ice-cold
15 saline followed by paraformaldehyde. Brains were post-fixed in 20% sucrose in paraformaldehyde,
16 sectioned coronally, and processed for immunohistochemistry for ERα and tyrosine hydroxylase.
17 Sections were counterstained with Nissl stain. Cells of the anteroventral periventricular nucleus positive
18 for ERα, tyrosine hydroxylase, or both were counted. Statistical analysis used 2-way ANOVA with sex
19 and treatment as factors, followed by 1-way ANOVA and post hoc Fisher least significant difference test.

20
21 There was a significant, sex-related effect on tyrosine hydroxylase-positive cells in the anteroventral
22 periventricular nucleus with the number in males about 29% that of females **[estimated from a graph]**.
23 Treatment effects are summarized in Table 80. The authors concluded that neonatal treatment with
24 bisphenol A interfered with the normal testosterone-associated masculinization of the anteroventral
25 periventricular nucleus. Because 17β-estradiol is aromatized to testosterone in the brain, the authors
26 interpreted this effect of bisphenol A as anti-estrogenic. Cells staining for both ERα and tyrosine
27 hydroxylase are not present in rodents after puberty, and the authors stated that these cells may play a role
28 in the organization of the LH-surge. They postulated that the decrease in these cells with neonatal
29 exposure to bisphenol A may result in cycle disruption in adulthood.

30
31 **Table 80. Effects of Neonatal Treatments on the Rat Anteroventral Periventricular Nucleus.**

Endpoint	Females		Males	
	17β-Estradiol	Bisphenol A	17β-Estradiol	Bisphenol A
Number of cells positive for				
tyrosine hydroxylase	↓50%	↔	↔	↑1.9-fold
ERα	↔	↔	↔	↔
both	↓38%	↓41%	↔	↔
Percent of cells positive for				
ERα + tyrosine hydroxylase	↔	↓41%	↔	↓63%

↑,↓,↔ Statistically significantly increased, decreased, unchanged compared to within-sex sesame oil control.
Estimated from figures in Patisaul et al. (338).

32
33 **Strengths/Weaknesses:** Strengths of this study are the use of 17β-estradiol as a positive control and the
34 measurement of ERα receptors. Weaknesses are the relatively high dose level of bisphenol A and the use
35 of the injection route of exposure of newborn pups.

36
37 **Utility (Adequacy) for CERHR Evaluation Process:** This study is moderately useful for the evaluation
38 process.

3.0 Developmental Toxicity

1 **Shikimi et al. (339)**, supported by the Japan Society for the Promotion of Science for Young Scientists,
2 examined the effects of bisphenol A exposure on Purkinje cell development in rats. **[No information was**
3 **provided about feed or composition of caging and bedding materials.]** At 6–9 days of age, 4 male or
4 female Fisher rats/group received bisphenol at 0 (sesame oil vehicle), 0.050, or 0.500 mg/day by injection
5 into the cerebrospinal fluid near the region of the cerebellum. During the same time period, additional
6 groups of 4 rats received 0.500 mg/day tamoxifen, 0.500 mg/day bisphenol A + 0.500 mg/day tamoxifen,
7 or 5 µg/day estradiol benzoate through the same exposure route. **[Both male and female rats were**
8 **treated, but it was not indicated if there were equal numbers in each group; both sexes were**
9 **apparently evaluated together.]** At 10 days of age, pups were killed and vermal cerebella were removed
10 and sectioned. Purkinje cells were examined morphologically following identification by calbindin-D_{28K}
11 immunostaining. Data were analyzed by ANOVA, followed by Duncan multiple range test. Treatment
12 with the high dose of bisphenol A increased Purkinje fiber length. There was no effect on cross-sectional
13 soma area or Purkinje cell number as a result of bisphenol A treatment. Cotreatment with tamoxifen
14 inhibited the increase in dendritic length that was observed following treatment with bisphenol A alone.
15 Estradiol benzoate also induced an increase in dendritic length of Purkinje fibers that was blocked by
16 tamoxifen. Treatment with tamoxifen alone also reduced dendritic fiber length. The effects of octylphenol
17 were also examined and an increase in dendrite length was observed. The study authors concluded that
18 bisphenol A induced Purkinje dendritic growth, possibly through the ER.

19
20 **Strengths/Weaknesses:** The use of 17β-estradiol as a positive control is a strength of this study.
21 Weaknesses are the injection into cerebrospinal fluid and the expression of dose as mg/day, both of which
22 prevent comparison with other studies.

23
24 **Utility (Adequacy) for CERHR Evaluation Process:** This paper has little utility in the evaluation
25 process.

26
27 **Zsarnovszky et al. (340)**, supported by NIH, NIEHS, and the American Heart Association, evaluated the
28 effect of intracerebellar injection of bisphenol A on the development of activated extracellular signal-
29 regulated kinase (ERK)-positive cells in cerebellar sections in Sprague Dawley rats. Neonatal rats on
30 PND 4–19 underwent a single direct injection under anesthesia of bisphenol A or 17β-estradiol under
31 stereotactic guidance into cerebellar folia 6 and 7. **[For bisphenol A, only PND 10 results were given.**
32 **The number of animals at each age was not specified, but a figure legend indicated at least 6/dose**
33 **group. The purity of the chemicals was not specified. The day of birth was not defined.]**
34 Concentrations of the chemicals were 10⁻¹² to 10⁻⁶ M **[bisphenol A concentrations of 0.23 ng/L to 0.23**
35 **mg/L]**. Uninjected, mock-injected, and vehicle-injected controls were used. Brains were removed and
36 fixed 6 minutes after the onset of the injection. Sections were processed for immunohistochemistry using
37 an antibody that recognized activated ERK. Quantitative analysis was performed on images of folium 9.
38 Statistical analysis was performed using ANOVA with post hoc Tukey-Kramer multiple comparison test.
39 Response to different chemicals and different concentrations on PND 10 were compared using 2-factor
40 ANOVA with post hoc Bonferroni test. Adult rats were also treated but were not included in the
41 quantitative analysis.

42
43 The qualitative appearance of the immunostained sections was similar after bisphenol A and 17β-
44 estradiol. In the 10⁻¹² to 10⁻⁹ M dose range, the quantitative responses to the 2 chemicals were similar.
45 Activated ERK-positive cells increased with a median effect concentration of 7.46 pM for 17β-estradiol
46 and 3.25 pM **[0.74 ng/L]** for bisphenol A. Both chemicals were described as having an inhibitory effect at
47 higher doses. **[The data graph shows drop-offs to control densities at 10⁻⁹ and 10⁻¹⁰ M, with a second**
48 **increase in density at 10⁻⁷ and 10⁻⁵ M.]** Coadministration of 10⁻¹⁰ M 17β-estradiol with bisphenol A
49 10⁻¹²–10⁻¹⁰ M **[0.23–23 ng/L]** resulted in a concentration-dependent decrease in activated ERK-positive
50 cells compared to the administration of 17β-estradiol alone. The authors concluded that 17β-estradiol

3.0 Developmental Toxicity

1 regulates ERK signaling in the developing cerebellum and that bisphenol A can mimic and also inhibit
2 this estrogenic effect, with potentially adverse affects on brain development and function.

3
4 **Strengths/Weaknesses:** Although the bisphenol A dose was not completely clear, it was probably low.
5 The use of 17 β -estradiol as a positive control is a strength.

6
7 **Utility (Adequacy) for CERHR Evaluation Process:** This study is useful in the evaluation and shows
8 that ERK signaling in the developing cerebellum is disrupted by bisphenol A.

9 10 3.2.5 Mouse—oral exposure only during pregnancy

11 3.2.5.1 Studies without neurobehavioral endpoints

12 **Morrissey et al. (273)**, supported by NTP/NCTR, examined the effects of prenatal bisphenol A exposure
13 in rats and mice in studies conducted according to GLP. The studies are also available as NTP
14 publications for rats (274) and mice (275). The study was conducted in two sets of rats and mice and data
15 were pooled for each species. [The data for rats were discussed in Section 3.2.1.] Pregnant CD-1 mice
16 were randomly assigned to groups of ≥ 10 animals in each set of the study, for a total of ≥ 20 animals/dose.
17 On GD 6–15 (GD 0 = sperm or plug), mice were gavaged with bisphenol A at 0, 500, 750, 1000, or 1250
18 mg/kg bw/day. Doses were based on results of preliminary studies and were expected to result in 10%
19 maternal mortality at the high dose and no toxicity at the low dose. The purity of bisphenol A was >95%,
20 and 2,4'-bisphenol A was reported as an impurity. Concentrations of dosing solutions were verified.
21 Pregnant animals were weighed during the study. Mice were killed on GD 17. Liver and uteri were
22 weighed, and corpora lutea and implantation sites were examined. Fetuses were sexed, weighed, and
23 examined for viability and external, visceral, and skeletal malformations. Data were analyzed by Bartlett
24 test for homogeneity of variance, ANOVA and/or William multiple comparison, Dunnett, and/or Fisher
25 exact probability tests.

26
27
28 Clinical signs reported in mice treated with bisphenol A included arched back, lethargy, piloerection,
29 rough coat, vaginal bleeding, vocalization, alopecia, weight loss, and wheezing. One or 2 of 29–34 dams
30 died in each of the 3 lowest dose groups and 6 of 33 dams died in the 1250 mg/kg bw/day group.
31 Statistically significant effects are summarized in Table 81. Absolute liver weight was increased in the
32 500, 750, and 1000 mg/kg bw/day dose groups, and relative liver weights were increased in all bisphenol
33 A dose groups. Decreased gravid uterine weight and dam body weight gain during the gestation and
34 treatment periods attained statistical significance at the 1250 mg/kg bw/day dose. The number of litters
35 available for evaluation in the control and each dose group was 26, 23, 21, 23, and 21. Increased
36 resorptions/litter and decreased fetal body weights/litter attained statistical significance in the high-dose
37 group. There was no effect on the number of live fetuses/litter at birth or on fetal malformations/litter.
38 The study authors concluded that bisphenol A is not teratogenic in mice at doses that result in maternal
39 toxicity.

40
41 **Table 81. Maternal and Developmental Toxicity in Mice Gavaged with Bisphenol A**

Endpoint	Dose in mg/kg bw/day				BMD ₁₀	BMDL ₁₀	BMD _{1SD}	BMDL _{1SD}
	500	750	1000	1250				
Dam weight in treatment period	↔	↔	↔	↓43%	881	661	1159	1039
Gravid uterine weight	↔	↔	↔	↓32%	983	690	1243	1123
Relative dam liver weight	↑9%	↑13%	↑17%	↑26%	618	411	755	541
Resorptions/litter	↔	↔	↔	↑2.8-fold	817	377	1245	1162
Fetal body weight/litter	↔	↔	↔	↓15%	1079	785	1249	1024

↑,↓ Statistically significant increase, decrease; ↔ no statistically significant change. From Morrissey et al. (273).

3.0 Developmental Toxicity

Strengths/Weaknesses: The oral route of exposure is a strength. The use of very high doses is a weakness.

Utility (Adequacy) for CERHR Evaluation Process: This paper is of moderate utility in the evaluation.

vom Saal et al. (341), supported by NIH, examined the effects of bisphenol A exposure on male reproductive organs and sperm production in mice. The CF-1 mice used in this study were purchased in 1979 and maintained as an outbred stock in a closed colony. Dams were fed Purina breeder chow (#5008) during pregnancy and lactation, and male offspring were fed Purina #5001 standard lab chow after weaning. Housing consisted of polypropylene cages with corn cob bedding. Bisphenol A [**purity not reported**] in tocopherol-stripped corn oil vehicle was fed to 7 mice/group by electronic micropipette at 0.002 or 0.020 mg/kg bw/day on GD 11–17 (day of vaginal plug = GD 0). One group of 6 mice was given the vehicle control, and a group of 5 mice was not handled. Based on results of in vitro assays conducted by the study authors, the 0.02 mg/kg bw/day bisphenol A dose was predicted to be bioactive in mice. Additional mice were treated with the same doses of octylphenol. Females delivered pups naturally on GD 19, and pups were weaned on PND 23 (day of birth not defined). Male siblings were housed 3/cage until 5 months of age. Randomly selected males were housed individually at 5 months of age and killed 1 month later. Body, testes, epididymides, preputial glands, and seminal vesicles were weighed in 11 control mice and 7 treated mice/group. Data from the two control groups did not differ significantly and were combined for analyses of organ and body weight. Data for prostate weight were reported by Nagel et al. (205). Daily sperm production was determined in 8 control males/group and 5 treated males/group. [**It was not stated how data from the 2 control groups were handled for sperm analyses.**] Sperm data were analyzed by ANOVA. Organ weight data were analyzed by ANCOVA, Pearson's correlation analysis, ANOVA, and least significant means test.

Statistically significant findings are summarized in Table 82. Exposure to bisphenol A resulted in dose-related reductions in daily sperm production efficiency (i.e., per g testis) that attained statistical significance at the highest dose level. Some significant but non-dose related effects were observed for body and organ weights. Epididymal weights were reduced at both doses. At the low dose, body and seminal vesicle weights were reduced and preputial weight was increased. In mice treated with octylphenol, daily sperm production was reduced at the low dose but there was no effect on reproductive organ weights. The study authors concluded that exposure of the fetus to low doses of endocrine-disrupting chemicals can affect the size and function of reproductive organs.

Table 82. Sperm Production and Male Reproductive Organ Weights in Mice Exposed to Bisphenol A During Gestation

Endpoint	Dose in mg/kg bw/day ^b					
	0.002	0.020	BMD ₁₀	BMDL ₁₀	BMD _{1SD}	BMDL _{1SD}
Sperm production efficiency	↔	↓19%	0.011	0.007	0.010	0.007
Body weight	↓9%	↔				
Preputial weight	↑36%	↔				
Seminal vesicle weight	↓12%	↔				
Epididymal weight	↓12%	↓8%				

^aEstimated by CERHR from a graph.

^bBenchmark doses were not estimated for values obtained from graphs and non-dose-related effects; errors were assumed to be SEM, as reported earlier in the paper. From vom Saal et al. (341).

[**The NTP Statistics Subpanel (295) noted that vom Saal et al. (341) did not apparently require overall differences by ANOVA to be significant before applying the least significant difference test,**

1 **which is prone to false positive findings without the overall protection of ANOVA. The NTP**
2 **Subpanel was not able to confirm any of the significant findings reported for bisphenol A. The NTP**
3 **Subpanel noted that in theory, their reanalysis of organ weights was not necessarily in conflict with**
4 **the findings of the study authors because of the use of different statistical methods (Dunnett test**
5 **versus Fisher least significant difference test).]**
6

7 **Strengths/Weaknesses:** Strengths are the use of oral delivery and low dose levels. Weaknesses are the
8 lack of clarity concerning the strain of mouse, failure to weight-adjust the maternal dose daily, the lack of
9 consideration of litter of origin in randomly selected males, the lack of information on testis weight
10 (which is needed for consideration of daily sperm production), and the questions about the statistical
11 analysis. In spite of these statistical questions, the 36% increase in preputial weight at 0.002 mg/kg
12 bw/day seems robust.
13

14 **Utility (Adequacy) for CERHR Evaluation Process:** The Expert Panel was divided on the utility of this
15 study.
16

17 **Nagel et al. (205)**, supported by NIH and the University of Missouri-Columbia, examined the effect of
18 prenatal bisphenol A exposure on mouse prostate weight. The mice used in this study were the same ones
19 used in the study by vom Saal et al. (341), and experimental details are provided in the summary of that
20 study. CF-1 mice were fed Purina Laboratory Chow 5001 and housed in polypropylene cages with corn
21 cob bedding. The mice (7/group) were dosed with bisphenol A [**purity not reported**] at 0.002 and 0.020
22 mg/kg bw/day on GD 11–17. The study authors stated that doses were within ranges of human exposure.
23 A control group of 6 mice was given the tocopherol-stripped corn oil vehicle during the same time period.
24 Vehicle and dosing solutions were fed to the mice using a micropipette. A second control group of 5 dams
25 was unhandled. Because there were no significant differences between the 2 control groups, data from the
26 2 groups were pooled. Females were allowed to litter. Pups were weaned at 23 days of age and housed
27 3/cage. One male/litter was selected and housed individually for 1 month. Body weights of males were
28 measured throughout the study. Selected males were killed at 6 months of age for measurement of
29 prostate weight. Data for prostate weight were analyzed by ANCOVA using body weight as the covariate.
30 If it was determined that body weight did not account for differences in prostate weight, data were
31 reanalyzed by ANOVA without adjustment for body weight. Body weights were lower in males from the
32 0.002 mg/kg bw/day group than in controls. Statistical analyses revealed that prostate weight was not
33 related to body weight. Compared to control values, prostate weights were 30% higher in the 0.002 mg/kg
34 bw/day group and 35% higher in the 0.020 mg/kg bw/day group. The study authors concluded that
35 bisphenol A alters the reproductive system of mice at doses near reported ranges of human exposure.
36

37 **[The NTP Statistics Subpanel (295) concluded that Nagel et al. (205) used appropriate statistical**
38 **methods, and the Subpanel reached essentially the same conclusions as the study authors regarding**
39 **elevated prostate weight.]**
40

41 **Strengths/Weaknesses:** Strengths are the use of the same methods as vom Saal et al. (341) and the use of
42 dose levels in the range of human exposure. The independent confirmation of the data analysis by the
43 NTP Statistics Subpanel is another strength. The lack of clarity on the mouse strain that was used is a
44 weakness. The Purina 5001 chow has high and variable levels of soy phytoestrogens, and the corn cob
45 bedding may be problematic due to antiestrogenic constituents. The method of selection of males is not
46 clear, and it appears that litter of origin was not considered. This study did not use a positive control,
47 although there are earlier reports from this laboratory using diethylstilbestrol.
48

49 **Utility (Adequacy) for CERHR Evaluation Process:** This paper is useful in the evaluation.
50

3.0 Developmental Toxicity

1 **Cagen et al. (342)**, support not indicated [**authors noted to work in industry**], examined the effects of
2 prenatal bisphenol A exposure on the developing reproductive system of male mice. The study attempted
3 to duplicate the findings by vom Saal et al. (341) and Nagel et al. (205) by repeating their procedures.
4 Exceptions were (1) use of larger group sizes to increase statistical power; (2) use of 4 dose levels instead
5 of 2; (3) use of 2 methods to determine sperm counts; (4) killing of male offspring at 90 instead of 180
6 days; (5) conducting the study according to GLP (6) obtaining mice from a commercial source instead of
7 an in-house bred colony; and (7) housing males individually after weaning. In the study by Cagen et al.,
8 CF-1 mice gaining more than 4.5 g weight from GD 0 to 10 (day of vaginal plug not stated) were
9 randomly assigned to groups of 28 animals and administered bisphenol A (>99% pure) 0.0002, 0.002,
10 0.020, or 0.2 mg/kg bw/day on GD 11–17. Two negative control groups with 28 dams each were given
11 the tocopherol-stripped corn oil vehicle. Because results from the two vehicle control groups were
12 statistically equivalent, data from the two groups were pooled. A positive control group of 28 mice was
13 given 0.2 µg/kg bw/day diethylstilbestrol. Dosing solutions were dripped into the animals' mouths using a
14 micropipette. Concentrations of dosing solutions were verified prior to dosing. Animals were fed certified
15 rodent chow #5002. Water was provided in glass bottles with Teflon seals. Cages were made of
16 polypropylene with steel lids. Corn cob bedding was used. Music was played at low volume to provide
17 background noise. Dams were monitored for clinical signs, food intake, body weight gain, and fertility
18 endpoints. Pups were counted and sexed at birth (PND 0) and monitored for survival and weight gain
19 until weaning on PND 22. Litters were culled to 8 pups on PND 4, leaving as many males as possible. At
20 weaning, no more than 4 males/litter (65–95 males/group) were randomly selected to continue in the
21 study and housed individually. The males were monitored for body weight gain and feed intake until they
22 were killed on PND 90. Brain, liver, kidneys, and reproductive organs were weighed. Daily sperm
23 production and epididymal sperm counts were determined and a histopathological examination of testes
24 was conducted. The litter was considered the experimental unit in statistical analyses. Data were analyzed
25 by Levene test, ANOVA, Dunnett test, rank transformation, Wilcoxon rank sum test with Bonferroni
26 correction, Fisher exact probability test, and binomial distribution test.

27
28 There were no clinical signs or significant differences in body weight gain or feed intake in dams. The
29 numbers of dams that died of unknown causes during the study were: 2 receiving vehicle controls; 1
30 dosed with diethylstilbestrol; 3 dosed with 0.0002 mg/kg bw/day bisphenol A; and 1 each in the 0.002
31 and 0.020 mg/kg bw/day bisphenol A groups. The number of total pups/litter was significantly lower than
32 controls in the 0.2 mg/kg bw/day bisphenol group (mean ± SD = 9.60 ± 3.85 compared to 12.37 ± 3.02 in
33 the control group). In communications with the animal vendor, it was determined that litter size in the
34 control group exceeded typical litter sizes (9–10 pups), and the study authors therefore concluded that the
35 effect was not treatment related. Bisphenol A had no significant effects on gestation index or duration,
36 percentage of male pups at birth, or pup survival and body weight during the lactation period. The same
37 endpoints were unaffected in the diethylstilbestrol group.

38
39 Terminal body weights were increased [**by 7%**] in the 0.020 mg/kg bw/day group and [**by 5%**] in the 2
40 mg/kg bw/day group. Bisphenol A did not affect absolute or relative (to body or brain) weights of
41 reproductive organs including prostate, preputial gland, seminal vesicle, or epididymis. Non-dose-related
42 effects were observed for brain and kidney weights, and the study authors concluded that the effects were
43 not treatment-related. There were no significant effects on cauda epididymal sperm concentration, daily
44 sperm production, or efficiency of sperm production. Testicular histopathology was not affected by
45 bisphenol A treatment. [**Data were not shown by authors.**] Reproductive development of male offspring
46 was also unaffected by diethylstilbestrol. The study authors noted that the diethylstilbestrol dose was
47 considered the “maximum effect” oral dose by vom Saal but was lower than doses affecting male
48 offspring in other studies. The study authors also noted that the effects of bisphenol A on prostate weight
49 and sperm production reported by vom Saal et al. (341) and Nagel et al. (205) were not repeated in this
50 study. They concluded that bisphenol A should not be considered a selective reproductive or
51 developmental toxicant.

3.0 Developmental Toxicity

1
2 [The NTP Statistics Subpanel (295) concluded that the statistical methods used by Cagen et al. (342)
3 were appropriate. Although the Subpanel agreed with the study author conclusions, they noted that
4 (1) a significant ANOVA is not a requirement for Dunnett test and (2) a Bonferroni correction of
5 the Wilcoxon-rank sum test was not needed because the study authors already required significance
6 by ANOVA, which was sufficient.]
7

8 **Strengths/Weaknesses:** The attempt to replicate the studies of vom Saal et al. (341) and Nagel et al.
9 (205), the use of litter analysis, the large sample sizes, and the agreement of the NTP Subpanel with the
10 author conclusions are strengths. Weaknesses include some of the differences between this study and the
11 studies being replicated, specifically the possible strain differences (since the mice came from a different
12 source), termination at 90 instead of 180 days, and the use of solo housing rather than small group
13 housing. The lack of response of the positive control is a weakness, but it is unclear why this dose of
14 diethylstilbestrol was expected to be positive.
15

16 **Utility (Adequacy) for CERHR Evaluation Process:** This paper is useful for the evaluation process.
17

18 **Ashby et al. (343)**, support not indicated, examined the effects of prenatal bisphenol A exposure on the
19 mouse reproductive system. The study attempted to duplicate the findings reported by vom Saal et al.
20 (341) and Nagel et al. (205). Both generations of CF-1 mice were fed RM1 diet containing 6.5% soy
21 during periods when they were not pregnant or lactating, and dams were fed RM3 diet containing 18.5%
22 soy during pregnancy and lactation. On postconception days 11–17, 8 dams/group were dosed with
23 bisphenol A (99% pure) at 0, 0.002, or 0.020 mg/kg bw/day. The negative control group was administered
24 the tocopherol stripped corn oil vehicle. A positive control group of 7 dams received diethylstilbestrol at
25 0.2 µg/kg bw/day. A naïve group of 7 dams was not weighed or dosed. The dosing solution was slowly
26 expelled from a pipette placed in the animals' mouths. Day of vaginal plug detection was designated
27 postconception day 1, which was stated to be consistent with GD 0 as the day of mating. Females that had
28 no vaginal plugs but gained >3.5 g were considered to be 10 days pregnant. Females with vaginal plugs
29 and those that gained >3.5 g were distributed evenly among treatment and control groups. Females that
30 gained >1 but <3.5 g were considered to be pregnant, but because the day of pregnancy could not be
31 determined, they were assigned to the naïve control group. Dams were allowed to litter. All female
32 offspring were weighed and monitored for vaginal opening. Females were killed at ~44 weeks of age, and
33 liver, kidney, and reproductive organs were weighed. Male pups were housed as littermates until PND
34 112 (day of birth designated as PND 1). To determine the effects of housing, ~3 males from 4–7
35 litters/group (11–21 males/group) were randomly selected and housed separately from PND 112 until
36 study termination, which occurred ~71 days later. The remaining male pups from 4–5 litters/group from
37 each litter (11–17/group) were housed together. Singly housed males were weighed and killed on PND
38 183–185, and group-housed males were weighed and killed on PND 186–187. Equal numbers of males
39 from each group were killed each day. Liver, kidney, and reproductive organs were weighed, and
40 testicular sperm count and efficiency were determined. Technicians were blinded to experimental
41 conditions. Measures taken to reduce stress to animals included administering test agents by drip feeding,
42 minimal handling of pups, and minimal environmental noise. Selection of 3 males from each litter
43 increased statistical power compared to previous studies (205, 344). Statistical analyses were conducted
44 using the individual offspring and the litter as the statistical unit. Data were evaluated by ANOVA and
45 Dunnett test. Results from vehicle-treated and naïve controls were pooled when there was no evidence of
46 a vehicle effect. Data from individually housed and group housed-males were pooled when they did not
47 differ significantly.
48

49 There were no significant differences in litter sizes or percentage of males/litter. In female offspring from
50 the bisphenol A groups, there were no significant effects on body weight or organ weights, including
51 cervix, uterus, vagina, and ovary. Age and weight at vaginal opening were also unaffected in groups

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1 exposed to bisphenol A. Vaginal opening was delayed in the diethylstilbestrol-treated group. The authors
 2 noted that because delayed vaginal opening is not an estrogenic effect and because vaginal opening was
 3 also delayed in the naïve control group, the effect was most likely not biologically significant.

4
 5 Significant effects observed in male offspring are summarized in Table 83. Significant effects included
 6 increased terminal body weights in the low-dose group, increased testis weight in both dose groups, and
 7 increased epididymis weight in the high-dose group. Because testis and epididymis weights relative to
 8 body weights were nearly identical to controls [**data not shown by study authors**], the authors
 9 considered the finding equivocal. Although prostate weights were slightly higher in the bisphenol A
 10 groups, there were no statistically significant effects on prostate weight when adjusted for body weight
 11 and litter effects. Daily sperm production was increased in both dose groups, but the study authors
 12 considered the finding equivocal due to low biological significance. The study authors noted that the
 13 study failed to confirm the increase in prostate weight and decrease in sperm production reported in the
 14 studies by vom Saal et al. (344) and Nagel et al. (205), but results were consistent with those reported by
 15 Cagen et al. (342). Possible reasons for variability between studies were stated as differences in
 16 background sound level, diet, and animal body weights. The study authors also mentioned the possibility
 17 of genetic drift occurring in mice bred in-house in the vom Saal laboratory.

18
 19 **Table 83. Treatment-related Findings in Male Mice Exposed to Bisphenol A during Prenatal**
 20 **Development**

Endpoint	Naïve control	Bisphenol A in mg/kg bw/day		Diethylstilbestrol 0.2 µg/kg bw/day
		0.002	0.020	
Terminal body weight	↔	↑(litter) 12%	↔	↔
Right or left testis weight	↔	↑(litter) 8–9%	↑(litter) 10–12%	↔
Left epididymis weight	↔	↔	↑(litter) 9%	↔
Right epididymis weight	↔	↔	↑(litter) 9%	↔
covaried with body weight				
Sperm/testis/day	↔	↑(individual) 11%	↑(individual) 17%	↔

↑, ↓ Statistically significant increase, decrease compared to vehicle controls or pooled controls in cases where the vehicle-treated and naïve controls did not differ; ↔ no statistically significant effect.
 From Ashby et al. (343).

21
 22 **[The NTP Statistics Subpanel (295) essentially reproduced the findings reported by Ashby et al.**
 23 **(343).]**

24
 25 **Strengths/Weaknesses:** Strengths are the rather close replication of the designs of the studies by vom
 26 Saal et al. (341) and Nagel et al. (205) with diet as the only major difference, the use of both solo and
 27 group housed mice, the use of diethylstilbestrol as a positive control, and the support of the conclusions
 28 by the NTP Statistics Subpanel. The lack of effect of the positive control is a weakness, but it is unclear
 29 why this dose of diethylstilbestrol was expected to give a positive response.

30
 31 **Utility (Adequacy) for CERHR Evaluation Process:** This paper is very useful in the evaluation.

32
 33 **Howdeshell et al. (345)**, support not indicated, examined the effect of prenatal bisphenol A exposure on
 34 age of puberty in female mice. [**No information was provided about chow or composition of bedding**
 35 **and cage materials.**] CF-1 mice (n = 21/group) were fed oil vehicle [**type of oil not specified**] or
 36 bisphenol A [**purity not reported**] at 0.0024 mg/kg bw/day on GD 11–17 [**day of vaginal plug not**
 37 **defined**]. On GD 19, pups were obtained by cesarean section. Intrauterine position of pups (i.e., located
 38 next to male or female pups) was noted at that time. Pups were raised by untreated mothers and weaned
 39 on PND 22. Body weights were measured, and pups were monitored for vaginal opening and time to
 40 estrus. Results were analyzed according to all pups from each dose group or in relation to intrauterine

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1 position. The study authors stated that fetuses positioned between 2 male mice were exposed to the lowest
2 levels of 17 β -estradiol, while exposures to 17 β -estradiol are highest in fetuses positioned next to female
3 fetuses. Data were analyzed on a litter basis to control for maternal effects. Age of vaginal opening was
4 covaried with weight at weaning. Numbers of female offspring evaluated were 75–111/group for body
5 weight and 51–58/group for vaginal opening. The study authors attempted to evaluate females from each
6 intrauterine position in each litter. **[No additional information was provided for statistical analysis in
7 this brief communication.]**
8

9 Body weight at weaning was significantly increased in females in the bisphenol A group. When analyzed
10 according to intrauterine position, body weights were 22% higher than controls in females who were not
11 positioned next to a male fetus and 9% higher in females who had been positioned next to 1 male in utero.
12 There were no significant effects on age of vaginal opening. **[It was not clear if the data presented were
13 covaried with body weight.]** Bisphenol A treatment significantly reduced the period between vaginal
14 opening and first estrus by ~2.5 days. When evaluated according to intrauterine position, a significant
15 decrease in time to first estrus was observed in females who were not positioned next to a male pup
16 (accelerated by ~5 days) and in females positioned next to 1 male [**~2 days**]. No statistically significant
17 findings were observed in females who had been positioned next to 2 males in utero. The study authors
18 concluded that prenatal exposure to bisphenol A at environmentally relevant levels altered postnatal
19 growth and reproductive function in female mice but that natural variations in individual endogenous
20 17 β -estradiol levels influenced the response to bisphenol A.
21

22 The results of this study were also discussed in a publication by Howdeshell and vom Saal (346), which
23 indicated that the work was supported by NIH and reported additional findings. There was a bisphenol A-
24 associated reduction in pup survival between birth and weaning. Complete litter death occurred in 6 of 21
25 litters in the bisphenol A group compared to 1 of 21 litters in the control group. Significantly increased
26 body weight of male pups at weaning was also reported for the bisphenol A group. Body weights were
27 highest in males who were positioned next to 2 female pups in utero and were 10% higher than body
28 weights of control males positioned next to 2 female fetuses in utero. No increase in body weight
29 occurred in males that were positioned between two male fetuses in utero.
30

31 **[The NTP Statistics Subpanel (295) requested the Howdeshell et al. (345) dataset for reanalysis, but
32 it was not provided by study authors.]**
33

34 **Strengths/Weaknesses:** Strengths are the oral route of exposure, the use of an environmentally-relevant
35 dose level of bisphenol A, and the assessment of puberty onset using vaginal smears. The omission of a
36 description of husbandry conditions and the lack of a positive control are weaknesses. Time from vaginal
37 opening to first estrus is not a standard endpoint for assessing puberty in mice. Although the authors
38 identified a litter-based analysis, in Study figure 1, the n values exceed the number of dams, suggesting
39 that some of the data were analyzed on a per pup basis.
40

41 **Utility (adequacy) for CERHR evaluation process:** This paper is useful for the evaluation process, but
42 the utility of the study is reduced by the analytic question and the lack of clear significance of time from
43 vaginal opening to first estrus.
44

45 **Gupta (347)**, supported by NIH, examined the effects of bisphenol A exposure on the reproductive
46 system of male mice. CD-1 mice were received on GD 12 (GD 0 = day of breeding). The mice were fed
47 Purina Chow-5 L9 at the Charles Rivers Laboratory and Purina Chow 5012 at the study author's
48 laboratory. **[No information was provided on bedding or caging materials.]** On GD 16–18, 15
49 mice/group were fed the corn oil/12% ethanol vehicle or 0.050 mg/kg bw/day bisphenol A **[purity not
50 reported]**. Additional groups of mice were administered diethylstilbestrol at 0.1 and 200 μ g/kg bw/day
51 and Aroclor at 0.050 mg/kg bw/day during the same time period. The bisphenol A dose level was based

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1 on a level reportedly considered safe by the FDA. Following delivery, litters were culled to 8 pups, with
2 at least 3 males. Body weight and anogenital distance were examined in 3 pups/litter (45 pups) on PND 3,
3 2 pups/litter (30 pups) on PND 21, and 1 offspring/litter (15 offspring) on PND 60. **[Although Table 1 of**
4 **the study lists the n value as 15–45/group, a statement in the methods section indicated that an**
5 **equal number of pups (n=1–3) were pooled from each litter.]** Prostate and epididymis were weighed in
6 15 offspring/group on PND 3, 21, and 60. Whole-tissue mounts of prostate were examined for growth in
7 15-day-old offspring (n = 4/group). Androgen binding was measured in prostates isolated at 3, 21, and 60
8 days of age, with 2–6 prostates pooled, depending upon age; an n of 5 was reported in Figure 2 of the
9 study. Data were analyzed by ANOVA.

10
11 Body weights of male offspring were not affected by bisphenol A treatment. In male pups of the
12 bisphenol A group compared to the control group, anogenital distance adjusted for body weight was
13 significantly increased **[by 22%]** on PND 3, **[by 25%]** on PND 21, and **[by 33%]** on PND 60. Prostate
14 weights in males of the bisphenol A group were significantly increased **[by 56%]** on day 3, **[by 39%]** on
15 day 21, and **[by 101%]** on day 60. Relative (to body weight) epididymis weight in the bisphenol A group
16 was significantly reduced **[by 35%]** on PND 60. Prostate growth was reported to be qualitatively
17 increased by bisphenol A exposure. Androgen receptor binding was increased on PND 21 and 60 **[by**
18 **~344% on PND 21 and 358% on PND 60, estimated from a graph]**. Similar effects were reported
19 following treatment with the low dose of diethylstilbestrol and Aroclor. In contrast, the high dose of
20 diethylstilbestrol reduced body weights, anogenital distance, prostate weight, and androgen receptor
21 binding.

22
23 The report also included an in vitro study to examine the effects of bisphenol A on prostate growth. The
24 urogenital sinus was dissected from GD 17 fetuses and cultured for 7 days in media containing 0, 5, or 50
25 ng/L bisphenol A with and without the addition of testosterone. The urogenital sinus was also incubated
26 in 0.1 or 0.5 ng/L diethylstilbestrol and 5 or 30 ng/L Aroclor. Prostates obtained from cultures were then
27 fixed in Bouin solution and examined histologically. A similar protocol was used to examine androgen
28 binding in cultured prostates, except that only the high doses of each compound were examined, and cells
29 were cultured for 6 days. Bisphenol A at 50 ng/L increased prostate size **[by 140%]** in the absence of
30 testosterone and **[by 150%]** in the presence of testosterone. Androgen binding in prostate was increased
31 **[by 200%]** following treatment with bisphenol A. Similar effects were reported with diethylstilbestrol
32 and the high Aroclor dose. The study authors concluded that the effects of in vivo studies were
33 reproduced in in vitro studies, which suggests a direct effect on reproductive organs of fetal mice.

34
35 In a subsequent commentary, Elswick et al. (348) noted several concerns and requested clarification of
36 the data analysis performed by Gupta. It was noted that statistical analyses were insufficiently described
37 to determine if analyses in addition to ANOVA were conducted. It was not indicated if post hoc tests
38 were used or if corrections were made for multiple comparisons. Table 1 of the study was noted to
39 contain a footnote indicating $P < 0.05$ (larger) or $P < 0.05$ (smaller). It was stated that determining a mean
40 and conducting a one-tailed post hoc test based upon whether the mean is larger or smaller is a source of
41 potential bias in the statistical analyses. Analyses conducted by Elswick et al. indicated that the
42 assumption of homogeneity of variance, a requirement for ANOVA, was not met for some data such as
43 anogenital distance on PND 3 (Table 1 of the study) and prostate size (Table 3 of the study). Therefore,
44 questions were raised about whether homogeneity testing was done or if data were transformed to account
45 for lack of homogenous variances prior to ANOVA. Failure to consider the litter as the experimental unit
46 was noted in cases where the sample size was listed as 30 and 45, while only 15 dams/group were treated.
47 It was noted that if anogenital distance was measured in the same animal at different time points, a
48 repeated-measures ANOVA would have been the appropriate statistical test. It was stated that correction
49 of anogenital distance by the cube root of body weight instead of body weight would have been preferred
50 to avoid overcorrection; ANCOVA with body weight as a covariate would have been a better method for
51 correcting anogenital distance, and the best method would have been a nested ANCOVA (dam within

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1 treatment). Questions were raised about whether sampling 1 pup/litter on PND 60 provided a reliable
2 estimate, especially for highly variable endpoints such as anogenital distance, which can be affected by
3 sex of the adjacent fetuses in the uterus. Organ weights were also stated to be variable, and it was
4 questioned whether sampling 1 offspring/litter on PND 60 resulted in a reliable estimate.

5
6 Gupta (349) responded to the questions raised by Elswick et al. Regarding the question of post hoc tests
7 for data analyzed by ANOVA, Gupta stated that comparisons using the least significant difference test
8 support the effect reported in the original paper. Gupta stated that the use of 1-tailed tests was never
9 mentioned and that the criticism was unfounded. The numbers of offspring examined at each age was
10 reiterated [**with no mention of considering the litter the statistical unit**]. It was stated that individual
11 animals were not identified because it would have required using a toe clip or tattoo, which is stressful to
12 the animals. Therefore, it was not known if the same animals were examined for anogenital distance at the
13 different time points and use of the repeated-measures ANOVA would not have been appropriate.
14 Regarding use of 1 animal/litter, it was stated that it is the standard procedure accepted by NIEHS to
15 control for litter effects. Correction of anogenital distance by body weight was stated to be appropriate
16 because of a significant correlation between body weight and anogenital distance ($r = 0.47$, $P < 0.001$).
17 Adjustment for litter effects was stated to occur because litter was nested within treatment in the
18 ANOVA. Gupta noted a typographical error in Table 3 of the original paper. Standard deviations for the
19 50 ng/L bisphenol A and Aroclor groups were mistakenly indicated to be 10-fold higher than the actual
20 values (i.e., the actual values were 0.024 for bisphenol A and 0.032 for Aroclor). The errors made it
21 appear that there were differences in variances between groups, when actually there were not. Gupta
22 stood by his original conclusion that low levels of bisphenol A alter the development of the male
23 reproductive tract. Lastly, Gupta noted a disparity between conclusions made in industry-funded studies
24 and studies conducted at independent academic laboratories.

25
26 **Strengths/Weaknesses:** Strengths are the oral route of administration, the use of a reasonable dose level
27 of bisphenol A, the use of diethylstilbestrol as a positive control, the prostate measurements at 3 postnatal
28 time points, and the use of an in vitro study to support the in vivo results. The question about the
29 statistical analyses is a weakness, but the response by the authors to the Elswick et al. letter was
30 satisfactory. The apparent lack of attention to possible litter effects and the unexpected direction of the
31 effect of bisphenol A on anogenital distance are additional weaknesses.

32
33 **Utility (Adequacy) for CERHR Evaluation Process:** This study is very useful in the evaluation.

34
35 **Iida et al. (350)**, supported by the Japan Society for Promotion of Science, examined the effect of
36 prenatal bisphenol A exposure on spermatogenesis in adult mice. [**No information was provided about**
37 **composition of feed, caging, or bedding.**] On GD 10–17 [**day of vaginal plug not defined**], ≥ 3 ddY
38 mice/group were orally administered bisphenol A [**purity not reported**] at 0 (corn oil vehicle), 1, 10, or
39 100 mg/kg bw/day. [**The specific method of oral dosing was not stated.**] At 60 days of age, 4–5 male
40 mice/dose group (obtained from 3 litters/dose group) were weighed and killed. Testes were removed and
41 fixed in paraformaldehyde for histopathological evaluation by light microscopy. At 120 days of age,
42 testicular histopathology was examined by light and electron microscopy in 3 mice/group from the control
43 and 10 mg/kg bw/day groups. Data were analyzed by ANOVA.

44
45 No effects on body weight were observed in 60-day-old mice. Significant and dose-related increases in
46 the incidence of abnormal seminiferous tubules were observed in mice exposed to bisphenol A. The
47 incidence of abnormal seminiferous tubules in the control and each respective treatment group was 3.7,
48 15.2, 17.7, and 31.5%. [**Benchmark dose analysis using a probit model and n = 3 litters gave a**
49 **BMD₁₀ = 44 and a BMDL₁₀ = 17 mg/kg bw/day.**] Examples of seminiferous tubule lesions included
50 luminal space loss in tubules, reduced numbers of maturing elongate spermatids, decreased tubular
51 diameter, aberrant distribution of spermatogenic cells in epithelium, and accumulation of material within

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1 tubules. In the 120-day-old mice exposed to 10 mg/kg bw/day, the same types of lesions were observed at
2 a higher incidence than controls (28.3 compared to 5.14%). Electron microscopic examinations of 2
3 abnormal seminiferous tubules from exposed 120-day-old mice revealed the presence of round but not
4 elongated spermatids, leading study authors to suggest disrupted spermatogenesis. Disorganized
5 arrangement of Sertoli cells was also observed in the 120-day-old mice of the 10 mg/kg bw/day group.
6 The study authors noted that degeneration of Sertoli cells may be the cause of aberrant distribution of
7 spermatogenic cells.
8

9 **Strengths/Weaknesses:** The oral route of delivery, the detailed study of the seminiferous tubules, and the
10 identification of a dose-related effect are strengths of this study. The lack of information on details of
11 husbandry, the small sample size, and the lack of adjustment for litter effects are weaknesses.
12

13 **Utility (Adequacy) for CERHR Evaluation Process:** This paper by itself is not useful based on the
14 small sample size and apparently inappropriate analysis. The paper might be used to support other studies
15 showing a similar effect with comparable dosing regimens.
16

17 **Timms et al. (351)**, supported by NIEHS and US EPA, examined the effects of bisphenol A exposure on
18 development of the prostate in mice. CD-1 mice were fed soy-based Purina 5008 chow, provided drinking
19 water in glass bottles, and housed in polypropylene cages. **[The type of bedding material was not**
20 **indicated.]** On GD 14–18 (day of mating = GD 0), pregnant mice were fed by micropipette with 0.010
21 mg/kg bw/day bisphenol A (n = 6), the tocopherol-stripped corn oil vehicle (n = 5), 0.1 µg/kg bw/day
22 ethinyl estradiol (n = 5), or 0.1 µg/kg bw/day diethylstilbestrol (n = 5), the positive control. The dose of
23 bisphenol A was based on previous findings that suggested bisphenol A was 100-fold less potent than
24 diethylstilbestrol in permanently increasing prostate size in mice. On GD 19, fetuses were removed by
25 cesarean section, and during the removal process, intrauterine position of male fetus relative to sex of
26 adjacent fetuses was recorded. To reduce effects associated with sex hormone exposure from the adjacent
27 fetus, 1 male/litter that developed between a male and female fetus was examined. Prostate morphology
28 was determined by a 3D computer reconstruction technique. Immunohistochemistry techniques were used
29 to measure levels of proliferating cell nuclear antigen and mouse keratin 5. Statistical analyses included
30 ANOVA, followed by Fisher least-squares mean test when statistical significance was obtained. In a
31 separate study, prostate morphology was examined in 4 pregnant mice/group that were dosed with vehicle
32 or 200 µg/kg bw/day diethylstilbestrol according to the procedures described above.
33

34 Bisphenol A increased numbers of ducts, volume, and proliferation in one or more prostate regions, as
35 outlined in Table 84. The pattern of proliferating cell nuclear antigen staining was similar to that observed
36 with mouse keratin 5, a basal epithelial cell maker. The study authors also reported a 56% increase in the
37 volume of the coagulating glands. **[Data were not shown by study authors.]** An abnormal narrowing
38 was observed in the portion of the urethra near the neck of the bladder. **[The volume of the cranial**
39 **urethra was reduced by 35% compared to controls. Malformation of prostatic sulci was reported,**
40 **but no information was provided on incidence or severity.]** Similar effects on the prostate were
41 reported in mice exposed to ethinyl estradiol and the low dose of diethylstilbestrol. Narrowing of the
42 cranial urethra was observed in mice exposed to ethinyl estradiol. In contrast, exposure to the high
43 diethylstilbestrol dose resulted in inhibited morphogenesis of the prostate. The study authors concluded
44 that the differentiating urogenital system of male mice is very sensitive to a low dose of bisphenol A.
45

1 **Table 84. Effects on Prostate Development in Mice Following Prenatal Exposure to 0.010 mg/kg**
 2 **bw/day Bisphenol A**

Endpoint ^a	Prostate region		
	Dorsolateral	Ventral	Dorsolateral and ventral
No. of prostate ducts	↑41%	↔	↑40%
Prostate duct volume	↑99%	↑78%	↑91%
Proliferating cell nuclear antigen staining	↑44%	↔	No data

↑,↓ Statistically significant increase, decrease; ↔ no statistically significant effect.

^aPercent changes calculated by CERHR differed slightly from values presented by authors; it was not clear which part of the prostate the authors' values represented.

From Timms et al. (351).

3
 4 **Strengths/Weaknesses:** Strengths are the oral route of administration, the reasonable dose level of
 5 bisphenol A, the use of diethylstilbestrol and 17β-estradiol as positive controls, and the sophisticated
 6 measures applied to the prostate.

7
 8 **Utility (Adequacy) for CERHR Evaluation Process:** This paper is very useful for the evaluation.

9
 10 **Palanza et al. (352)**, supported by NIEHS, NIH, MURST, the University of Parma, and the National
 11 Council for Research, examined the effects of bisphenol A treatment on maternal behavior following
 12 exposure of mice during prenatal development and/or adulthood. The CD-1 mice used in this study were
 13 maintained as an outbred colony. Mice were housed in polypropylene cages with corn cob bedding.
 14 During pregnancy and lactation, mice were fed Purina 5008 (soy-based) chow. After weaning, mice were
 15 fed Purina 5001 (soy-based) chow. Water was provided in glass bottles. On GD 14–18 (GD 0 = day of
 16 vaginal plug), 14 mice were fed the tocopherol-stripped corn oil vehicle and 9 mice were fed 0.010 mg/kg
 17 bw/day bisphenol A [**purity not reported**] using an electronic micropipette. Dams were housed 3/cage
 18 after mating and individually housed on GD 17. Body weights of dams were measured during gestation.
 19 The day of birth was considered PND 1, and offspring were weaned on PND 20. At 2–2.5 months of age,
 20 F₁ female offspring from vehicle- and bisphenol A-treated dams were mated and exposed to vehicle or
 21 0.010 mg/kg bw/day bisphenol A on GD 14–18. There were 4 groups of F₁ females that were exposed
 22 during gestation-adulthood to vehicle-vehicle (n = 20), vehicle-bisphenol A (n = 15), bisphenol A-vehicle
 23 (n=15), and bisphenol A–bisphenol A (n=15). Maternal behavior was observed in F₁ dams every 4
 24 minutes during a 120-minute period on PND 2–15. On PND 1, F₂ pups were weighed, sexed, and
 25 counted. Litters were then culled to 10 pups, with equal numbers of male and female pups when possible.
 26 Pups were weighed during the lactation period and cliff-drop aversion and righting reflex were evaluated
 27 in all pups of a subset of 8 litters/group on PND 3, 5, 7, and 9. For statistical analyses, all pup data were
 28 adjusted for litter. Data were analyzed by ANOVA, Holms *t*-test, and/or Fisher protected least-squared
 29 difference test.

30
 31 Bisphenol A treatment did not affect gestational body weight gain in F₀ or F₁ dams. Statistically
 32 significant effects for F₁ maternal behavior collapsed across 14 observation days are presented in Table
 33 85. Exposure to bisphenol A either in gestation or in adulthood resulted in decreases in the percentage of
 34 time the dams spent nursing and in the nest and increases in the percentage of time the dams spent nest
 35 building, resting alone, grooming, and out of the nest. Increase activity was also observed in the group
 36 exposed to bisphenol A in adulthood. The only significant effect observed in mice exposed to bisphenol A
 37 during gestation and adulthood was increased time resting. When data were presented for individual
 38 evaluation days, time resting was significantly increased on PNDs 9, 10, 11, 12, and 14 in the group
 39 exposed to bisphenol A during gestation. Time spent resting was significantly increased on PND 9 and 14
 40 in the group exposed to bisphenol A during gestation and adulthood. No other significant effects were
 41 observed on specific evaluation days. There were no significant differences in the number of live F₂
 42 pups/litter, sex ratio, or body weight at birth or in weight gain during the lactation period. [**Data were not**

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1 **shown]**. No significant effects were observed for cliff aversion or righting reflexes. The study authors
 2 concluded that reduced levels of nursing behavior were observed in mice exposed to bisphenol A only as
 3 fetuses or only as adults. **[Because this study involves effects of adult exposure on maternal**
 4 **behaviors, it is also discussed in Section 4.2]**

6 **Table 85. Maternal Behavior Effects in Mice Exposed to Bisphenol A During Gestation and/or**
 7 **Adulthood**

Percent time ^a	Bisphenol A exposure during gestation/adulthood		
	Bisphenol A/vehicle	Vehicle/bisphenol A	Bisphenol A/bisphenol A
Nursing	↓18%	↓13%	↔
Nest building	↑67%	↑150%	↔
Resting alone	↑71%	↑14%	↑43%
Grooming	↑27%	↑18%	↔
Active	↔	↑17%	↔
In nest	↓12%	↓8%	↔
Out of nest	↑9%	↑5%	↔

^aValues were estimated from a graph by CERHR.

↑,↓ Statistically significant increase/decrease compared to vehicle-vehicle group, ↔ no statistically significant effect.

From Palanza et al. (352).

8
 9 **Strengths/Weaknesses:** Strengths are the oral route of administration, the low dose level of bisphenol A,
 10 and the exploration of effects on complex maternal behaviors. It is unusual that pre- and postnatal
 11 exposure had effects but not the combination of pre- and postnatal exposure, and failure to explain this
 12 finding is a weakness. Only 1 of 6 maternal behaviors was affected in the mice exposed during both time
 13 periods. The use of a diet high in soy isoflavones is an additional weakness.

15 **Utility (Adequacy) for CERHR Evaluation Process:** This paper is very useful in the evaluation.

16
 17 **Nishizawa et al. (353)**, supported by the Japanese Ministry of Education, Culture, Sports, Science, and
 18 Technology, examined the effects of prenatal bisphenol A exposure on expression of retinoic acid
 19 receptor α and retinoid X receptor α in mouse embryos. ICR mice were fed standard feed (CM, Oriental
 20 Yeast, Tokyo). **[No information was provided about caging and bedding materials.]** Mice were orally
 21 dosed with bisphenol A at 0 (olive oil vehicle) or 0.002 mg/kg bw/day on 6.5–11.5, 6.5–13.5, 6.5–15.5,
 22 and 6.5–17.5 days post coitum. Day of vaginal plug was considered 0.5 days post coitum. **[No**
 23 **information was provided about the specific method of oral dosing.]** Twelve dams/group were killed
 24 at 12.5, 14.5, 16.5, and 18.5 days post coitum, 24 hours after receiving the last dose. Expression of
 25 mRNA for retinoic acid receptor α and retinoid X receptor α was measured by RT-PCR in fetal cerebrum,
 26 cerebellum, and gonads. Data were analyzed by ANOVA. Changes in gene expression are summarized in
 27 Table 86. Numerous changes in mRNA expression were observed following in utero exposure to
 28 bisphenol A, and they varied according to sex, tissue, and dosing period. The study authors concluded
 29 that these findings suggest a novel mechanism of bisphenol A toxicity mediation by disruption of the
 30 expression of retinoic acid receptor α and retinoid X receptor α .

1 **Table 86. Expression of Retinoic Receptor α and Retinoid X Receptor α in Mouse Embryos**
 2 **Exposed to 0.002 mg/kg bw/day Bisphenol A In Utero**

Days post coitum	Gene and Tissue					
	Retinoic acid receptor α			Retinoid X receptor α		
	Cerebrum	Cerebellum	Testis/Ovary	Cerebrum	Cerebellum	Testis/Ovary
<i>Males</i>						
12.5	↔	↑	↔	↓	↓	↔
14.5	↓	↔	↓	↔	↔	↓
16.5	↔	↔	↔	↔	↔	↔
18.5	↔	↔	↓	↔	↔	↔
<i>Females</i>						
12.5	↔	↑	↔	↔	↓	↔
14.5	↓	↔	↔	↔	↓	↓
16.5	↔	↔	↔	↔	↔	↔
18.5	↔	↔	↔	↔	↓	↔

↑,↓ Statistically significant increase, decrease compared to controls; ↔ no statistically significant effect compared to controls.

From Nishizawa et al. (353).

3

4 **Strengths/Weaknesses:** Strengths are the oral route of delivery, the use of a low dose level of bisphenol
 5 A, and the exposure at different time periods. The results did not permit the easy identification of a
 6 critical period.

7

8 **Utility (Adequacy) for CERHR Evaluation Process:** This paper is slightly useful in the evaluation.

9

10 **Nishizawa et al. (354)**, supported by the Japanese Ministry of Education, Culture, Sports, Science, and
 11 Technology and by the Japan Society for the Promotion of Science, examined the effects of bisphenol A
 12 exposure on expression of mRNA for arylhydrocarbon and retinoid receptors in mouse embryos. ICR
 13 mice were fed standard diet (CM; Oriental Yeast, Tokyo). **[No information was provided about caging**
 14 **or bedding materials.]** Pregnant mice were orally dosed with bisphenol A at 0 (olive oil vehicle),
 15 0.00002, 0.002, 0.20, or 20 mg/kg bw/day from 6.5 to 13.5 days post coitum or 6.5 to 17.5 days post
 16 coitum. Day of vaginal plug detection was considered 0.5 days post coitum. **[No information was**
 17 **provided about the specific method of oral dosing.]** Twelve pregnant mice/group were killed on 14 and
 18 18.5 days post coitum, 24 hours after the last bisphenol A dose was administered. RT-PCR analyses were
 19 conducted to determine expression of mRNA for retinoic acid, retinoid X, and arylhydrocarbon receptors
 20 in fetal cerebrum, cerebellum, ovary, and testis. Data were analyzed by ANOVA. Changes in gene
 21 expression are summarized in Table 87. Numerous changes in mRNA expression were observed
 22 following bisphenol A exposure and they varied according to dose, sex, tissue, and exposure period. The
 23 study authors concluded the this study demonstrates a novel mechanism by which bisphenol can induce
 24 endocrine disruption through upregulation of arylhydrocarbon receptor (a key factor in the metabolism of
 25 some xenobiotics compounds) and retinoid receptors (key factors in nuclear receptor signal transduction).

26

3.0 Developmental Toxicity

1 **Table 87. Changes in mRNA Gene Expression in Mice Following In Utero Exposure to Bisphenol A**

Gene	Tissue	Days post coitum	Dose in mg/kg bw/day			
			0.00002	0.002	0.20	20
<i>Males</i>						
Arylhydrocarbon receptor	Cerebrum	14.5	↑	↔	↑	↑
		18.5	↑	↔	↑	↑
	Cerebellum	14.5	↑	↔	↑	↑
		18.5	↑	↔	↑	↑
	Testis	14.5	↑	↔	↔	↑
		18.5	↑	↑	↑	↔
Retinoic acid receptor α	Cerebrum	14.5	↔	↔	↑	↑
		18.5	↔	↔	↑	↑
	Cerebellum	14.5	↑	↔	↔	↔
		18.5	↑	↔	↑	↑
	Testis	14.5	↔	↔	↔	↔
		18.5	↔	↔	↔	↑
Retinoid X receptor α	Cerebrum	14.5	↑	↔	↑	↑
		18.5	↔	↔	↑	↑
	Cerebellum	14.5	↑	↔	↑	↑
		18.5	↔	↔	↔	↔
	Testis	14.5	↔	↔	↔	↑
		18.5	↔	↔	↑	↔
<i>Females</i>						
Arylhydrocarbon receptor	Cerebrum	14.5	↑	↔	↔	↑
		18.5	↑	↔	↑	↑
	Cerebellum	14.5	↑	↔	↑	↑
		18.5	↑	↔	↑	↑
	Ovary	14.5	↑	↔	↔	↑
		18.5	↑	↑	↑	↔
Retinoic acid receptor α	Cerebrum	14.5	↔	↔	↑	↑
		18.5	↔	↔	↑	↑
	Cerebellum	14.5	↑	↔	↔	↑
		18.5	↑	↔	↑	↑
	Ovary	14.5	↑	↔	↑	↑
		18.5	↔	↔	↔	↑
Retinoid X receptor α	Cerebrum	14.5	↑	↔	↑	↑
		18.5	↔	↔	↑	↑
	Cerebellum	14.5	↑	↔	↔	↑
		18.5	↔	↔	↔	↔
	Ovary	14.5	↔	↔	↔	↑
		18.5	↔	↔	↑	↔

↑Statistically significant increase compared to controls; ↔ No statistically significant effect compared to controls.

Nishizawa et al. (354).

2
3 **Strengths/Weaknesses:** The wide dose range from 0.00002 to 20 mg/kg bw/day and the oral route are
4 strengths, although the lack of specification of the method of oral dosing is a weakness.

5
6 **Utility (Adequacy) for CERHR Evaluation Process:** This paper is moderately useful and suggests a
7 new mechanism of action of bisphenol A, upregulation of arylhydrocarbon receptor and retinoic

3.0 Developmental Toxicity

1 receptors. The very low dose had an effect on mRNA expression related to retinoic acid, the next dose
2 had little effect, then effects were again seen at higher doses.

3
4 **Nishizawa et al. (355)**, supported by the Japan Society for the Promotion of Science, examined the
5 effects of bisphenol A exposure on expression of aryl hydrocarbon receptors, related factors, and
6 metabolizing enzymes in mouse embryos. ICR mice were fed standard diet (CM, Oriental Yeast, Tokyo).
7 **[No information was provided about caging and bedding materials.]** Mice were orally dosed with
8 bisphenol A at 0 (olive oil vehicle), 0.00002, 0.002, 0.2, or 20 mg/kg bw/day from 6.5–13.5 days post
9 coitum and 6.5 to 17.5 days post coitum. Day of vaginal plug was considered 0.5 days post partum. **[No**
10 **information was provided about the method of oral dosing.]** Another group of mice was dosed with 5
11 µg/kg bw/day 17β-estradiol during the same time periods. Twelve mice/group were killed at 14.5 and
12 18.5 days post coitum, 24 hours after receiving the final dose. Embryos were dissected to obtain
13 cerebrum, cerebellum, ovary, testis, and liver. RT-PCR analysis was used to measure mRNA levels of
14 genes listed in Table 88. Western immunoblotting was used to measure protein levels of CYP1A1 and
15 glutathione-S-transferase in liver. Data were analyzed by ANOVA.

16
17 Changes in gene expression are summarized in Table 88. Numerous changes in mRNA expression were
18 observed following bisphenol A exposure, and they varied according to dose, sex, tissue, and exposure
19 period. In at least one sex and time period, exposure to 17β-estradiol increased expression of mRNA
20 arylhydrocarbon receptor in all tissues, arylhydrocarbon receptor repressor in testes and ovaries,
21 arylhydrocarbon receptor nuclear translocator in brain or testes, *CYP1A1* in brain, and glutathione S-
22 transferase in brain. Changes in protein levels of CYP1A1 and glutathione S-transferase in liver were also
23 examined in embryos at 18.5 days post coitum and levels of both proteins were increased with exposure
24 to bisphenol A at doses ≥0.2 mg/kg bw/day and with exposure to 17β-estradiol. The study authors
25 proposed a novel mechanism of toxicity involving up-regulation of mRNA for arylhydrocarbon receptor
26 and other factors by bisphenol A.

27
28 **Table 88. Changes in mRNA Expression in Mice Following In Utero Exposure to Bisphenol A**

Gene	Tissue	Days post coitum	Dose in mg/kg bw/day			
			0.00002	0.002	0.20	20
<i>Males</i>						
Arylhydrocarbon receptor	Cerebrum	14.5	↑	↔	↑	↑
		18.5	↑	↔	↑	↑
	Cerebellum	14.5	↑	↔	↑	↑
		18.5	↑	↔	↑	↑
	Testis	14.5	↑	↔	↔	↑
		18.5	↑	↑	↑	↔
Arylhydrocarbon receptor repressor	Cerebrum	14.5	↑	↔	↔	↔
		18.5	↑	↔	↑	↑
	Cerebellum	14.5	↑	↔	↑	↑
		18.5	↑	↔	↑	↑
	Testis	14.5	↔	↑	↑	↔
		18.5	↑	↔	↔	↑
Arylhydrocarbon receptor nuclear translocator	Cerebrum	14.5	↔	↔	↑	↑
		18.5	↑	↔	↔	↔
	Cerebellum	14.5	↑	↔	↔	↑
		18.5	↑	↔	↔	↑
	Testis	14.5	↔	↔	↔	↔
		18.5	↑	↑	↔	↔
<i>CYP1A1</i>	Cerebrum	14.5	↔	↔	↔	↔
		18.5	↔	↔	↑	↑

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Gene	Tissue	Days post coitum	Dose in mg/kg bw/day				
			0.00002	0.002	0.20	20	
Glutathione S-transferase	Cerebellum	14.5	↔	↔	↔	↑	
		18.5	↔	↔	↑	↑	
	Testis	14.5	↔	↔	↔	↔	
		18.5	↔	↔	↔	↔	
	Cerebrum	14.5	↔	↔	↔	↔	
		18.5	↔	↔	↔	↑	
	Cerebellum	14.5	↔	↔	↔	↔	
		18.5	↔	↔	↑	↑	
	Testis	14.5	↔	↔	↔	↔	
		18.5	↔	↔	↑	↑	
	<i>Females</i>						
	Arylhydrocarbon receptor	Cerebrum	14.5	↔	↔	↔	↔
18.5			↑	↔	↑	↑	
Cerebellum		14.5	↑	↔	↑	↑	
		18.5	↑	↔	↑	↑	
Arylhydrocarbon receptor repressor	Ovary	14.5	↑	↔	↔	↑	
		18.5	↑	↑	↑	↔	
	Cerebrum	14.5	↑	↔	↔	↔	
		18.5	↑	↔	↑	↑	
Arylhydrocarbon receptor nuclear translocator	Cerebellum	14.5	↑	↔	↑	↑	
		18.5	↑	↔	↑	↑	
	Ovary	14.5	↔	↑	↑	↔	
		18.5	↑	↔	↔	↑	
CYP1A1	Cerebrum	14.5	↔	↔	↔	↔	
		18.5	↔	↔	↑	↑	
	Cerebellum	14.5	↔	↔	↔	↑	
		18.5	↔	↔	↑	↑	
	Ovary	14.5	↔	↔	↔	↔	
		18.5	↔	↔	↔	↔	
Glutathione S-transferase	Cerebrum	14.5	↔	↔	↔	↔	
		18.5	↔	↔	↔	↑	
	Cerebellum	14.5	↔	↔	↔	↔	
		18.5	↔	↔	↑	↑	
	Ovary	14.5	↔	↔	↔	↔	
		18.5	↔	↔	↑	↑	

↑Statistically significant increase compared to controls; ↔ No statistically significant effect compared to controls.

From: Nishizawa et al. (355).

1
2
3
4

Strengths/Weaknesses: This paper had the same strengths and weaknesses as the previous study from this laboratory (354).

3.0 Developmental Toxicity

1 **Utility (Adequacy) for CERHR Evaluation Process:** This paper is useful and replicates many of the
2 findings of the previous study. Again, the 0.002 mg/kg bw dose gave the weakest receptor mRNA
3 response.
4

5 **Yoshino et al. (356)**, supported by the Japanese Ministry of Education, Science, Sports, and Culture and
6 the Japan Private School Promotion Foundation, examined the effect of prenatal bisphenol A exposure on
7 immune response in mice. **[No information was provided about feed or caging and bedding**
8 **materials.]** DBA/1 J mice were fed bisphenol A **[purity not indicated]** at doses of 0 (ethanol/corn oil
9 vehicle), 0.003, 0.030, 0.300, or 3 mg/kg bw/day for 18 days **[stated to be 17 days in the Methods**
10 **section but 18 days in other parts of the report]**, beginning on the day of a 24-hour mating period (day
11 0). Twelve mice/group were treated and 7–9/group became pregnant. **[The specific method of oral**
12 **dosing was not described.]** At 8 weeks of age (day 77) 5 mice/group/sex were randomly selected and
13 immunized by ip injection with hen egg lysozyme. Blood was collected and spleens were removed 3
14 weeks following immunization (day 98). Serum levels of hen egg lysozyme-specific immunoglobulin G
15 (IgG), IgG1, and IgG2A were measured by ELISA. Spleen cell suspensions were prepared, and
16 proliferation was assessed by incorporation of ³H-thymidine following a 72-hour incubation with hen egg
17 lysozyme. Spleen cell suspensions were also prepared for measurement of interferon- γ and interleukin-4
18 secretion by ELISA. An additional 6 mice/group/sex were killed at 8 weeks of age (day 77). Spleens were
19 removed and expression of CD3⁺CD8⁺ and CD3⁺CD4⁺ molecules on splenic lymphocytes was examined
20 using monoclonal antibodies and flow cytometry. Thymus and spleen were fixed in 4% formaldehyde and
21 examined histologically. Data were analyzed by Mann-Whitney *U* test.
22

23 Bisphenol A treatment had no significant effect on pregnancy rate, sex ratio, or body weight of offspring.
24 Statistically significant immune responses for male mice are summarized in Table 89. **[Results in female**
25 **mice were said to be similar to those observed in male mice but the data were not show by study**
26 **authors.]** At bisphenol A doses ≥ 0.03 mg/kg bw/day, production of anti-hen egg lysozyme IgG2a
27 following immunization was increased. Effects observed at ≥ 0.3 mg/kg bw/day included increases in
28 production of anti-hen egg lysozyme IgG and secretion of interferon- γ and interleukin-4. Additional
29 findings at the high dose (3 mg/kg bw/day) were increases in spleen cell proliferation and production of
30 anti-hen egg lysozyme IgG1 following immunization. Augmentation of interferon- γ and interleukin-4
31 secretion following incubation of spleen cells with hen egg lysozyme was examined in the high-dose
32 group only and found to be increased. **[Increases in CD3⁺CD8⁺ and CD3⁺CD4⁺ expression on**
33 **lymphocytes were reported in males and females exposed to bisphenol A, but the doses at which the**
34 **effects occurred were not specified.]** No histopathological alterations were reported for the spleen or
35 thymus. The study authors explained that effects on IgG2a and interferon- γ were indicators of T helper 1
36 immune responses and effects on IgG1 and interleukin-4 were indicators of T helper 2 responses. They
37 concluded that the findings suggest that prenatal exposure to bisphenol A may up-regulate immune
38 responses in mice.
39

1 **Table 89. Immune Responses in Mice Following Prenatal Exposure to Bisphenol A Hen Egg**
 2 **Lysozyme**

Endpoint	Dose in mg/kg bw/day ^a			
	0.003	0.030	0.300	3
Quantity of anti-hen egg lysozyme IgG following immunization	↔	↔	↑	↑72%
Proliferative response to spleen cells following exposure to hen egg lysozyme	↔	↔	↔	↑65%
Production of anti-hen egg lysozyme IgG2a following immunization	↔	↑	↑	↑134%
Production of anti-hen egg lysozyme IgG1 following immunization	↔	↔	↔	↑51%
Secretion of interferon- γ	↔	↔	↑	↑200%
Secretion of interleukin-4	↔	↔	↑	↑62%
Augmentation of interferon- γ secretion following incubation with hen egg lysozyme	Not examined	Not examined	Not examined	↑
Augmentation of interleukin-4 secretion following incubation with hen egg lysozyme	Not examined	Not examined	Not examined	↑

↑ Statistically significant increase compared to control values; ↔ no significant difference compared to controls.

^aPercent changes compared to the control were only included when presented by study authors because CERHR attempts to estimate the values from graphs did not match the study author estimates. From Yoshino et al. (356).

3
 4 **Strengths/Weaknesses:** The oral route of administration and the wide range of doses are strengths.

5
 6 **Utility (Adequacy) of CERHR Evaluation:** This study is moderately useful and showed up-regulated
 7 immune response.

8
 9 *3.2.5.2 Studies with neurobehavioral endpoints*
 10 **Narita et al. (357)**, supported by the Japanese Ministry of Health, Labor, and Welfare, and Ministry of
 11 Education, Culture, Sports, Science, and Technology, conducted a series of studies to examine the effects
 12 of bisphenol A on the dopaminergic system of mice exposed during development. Only brief details were
 13 provided about the studies. In each study, ddy mice received feed containing bisphenol A from mating to
 14 weaning of their offspring. **[No information was provided on purity of bisphenol A, type of feed,**
 15 **caging and bedding materials, the number of dams treated, or the ages or sexes of offspring that**
 16 **were tested.]** Statistical analyses included ANOVA with Bonferroni/Dunnett test. In a place
 17 conditioning-study, testing was conducted in 6–14 mice/group born to dams exposed to bisphenol A at 0,
 18 0.03, 0.3, 3, 500, or 2000 mg/kg food. **[Assuming a female mouse eats ~0.2 kg feed/kg bw/day (81),**
 19 **bisphenol A intake would have been 0.006, 0.06, 0.6, 100, or 400 mg/kg bw/day.]** During the
 20 preconditioning period, mice were placed in one section of a cage following injection with saline **[specific**
 21 **route not reported]** and in another section of the cage following sc injection with 1 mg/kg bw morphine.
 22 On the day of testing, the amount of time spent in each section of the cage was recorded. Mice from the
 23 lowest dose group (0.03 mg/kg food) and 2 highest dose groups (500 and 2000 mg/kg) food spent more
 24 time in the section of the cage associated with morphine injection. **[Compared to controls, the time**
 25 **spent in the morphine-associated section of the cage was~ 9.5-, 7-, and 9-fold longer in each of the**
 26 **respective dose groups.]** Total locomotor activity was measured for 3 hours in 5–15 mice/group born to
 27 dams exposed to 0, 0.03, 3, or 2000 mg/kg food. Following sc injection with 10 mg/kg bw morphine,

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1 activity was increased in mice from the low- (0.03 mg/kg food) and high- (2000 mg/kg food) dose groups
2 compared to the control group [**increased by ~9-fold in the low dose group and 12-fold in the high-**
3 **dose group**]. Binding of ³⁵S-guanosine-5' [γ-thio]-triphosphate in the limbic system was measured in 3
4 samples/group obtained from offspring of dams exposed to 0.03, 3, or 2000 mg/kg food. Dopamine-
5 induced binding of ³⁵S-guanosine-5' [γ-thio]-triphosphate in the limbic system was increased at each dose
6 level compared to controls [**by ~32, 18, and 56%**]. Based on their findings, the study authors concluded
7 that prenatal and neonatal exposures to low bisphenol A doses can potentiate central dopamine receptor-
8 dependent neurotransmission in the mouse.

9
10 **Strengths/Weaknesses:** This paper is so poorly written that it is extremely difficult to understand many
11 sentences (let alone paragraphs) and to determine precisely what was done, why, and what happened. The
12 main weakness of the paper is therefore its inability to pass its message to the reader. Given this
13 limitation, it is difficult to determine whether the paper has any strengths, and if so what they might be.

14
15 **Utility (Adequacy) for CERHR Evaluation:** This paper is inadequate for the evaluation process because
16 of the lack of methodologic details and the poor communication of the study results.

17
18 **Kawai et al. (358)**, supported by Core Research for Evolutional Science and Technology, Japan Science
19 and Technology, examined the effects of prenatal bisphenol A exposure on aggressive behavior in male
20 mice. [**No information was provided about feed or bedding and caging materials.**] Pregnant CD-1
21 mice were randomly assigned to groups of 7 and orally dosed by micropipette with 0.002 or 0.020 mg/kg
22 bw/day bisphenol A [**purity not reported**] on GD 11–17. A control group of 9 mice received the corn oil
23 vehicle by micropipette during the same time period. Doses were said to be within the range of human
24 exposures. Pups were weaned on PND 21 (day of birth = PND 0), and randomly selected males from the
25 same litter were housed in groups of 4 or 5. Aggression testing was conducted at 8, 12, and 16 weeks of
26 age. For the testing, 15 control male mice from the 9 litters were randomly selected to be opponents and
27 housed 5/cage. Opponents were used only once/day for testing. During testing of mice from the control
28 and treated groups, the subject was housed alone for 5 minutes prior to placing the opponent mouse into
29 the cage. Behavior with the opponent mouse was observed for 7 minutes. The numbers of mice evaluated
30 were 26–32/group at 8 weeks of age, 18–24/group at 12 weeks of age, and 10–16/group at 16 weeks of
31 age. Randomly selected mice were killed at 9, 13, and 17 weeks of age, one week following behavior
32 testing, for measurement of testis weight and serum testosterone level. [**The results section states that**
33 **testis weights and serum testosterone levels were obtained at 8, 12, and 16 weeks of age.**] Eight
34 mice/group were killed after the first 2 test periods and 10–16 mice/group were killed after the last test
35 period. Mice that were not killed were tested at the next evaluation period, so that mice killed after 16
36 weeks of age were tested a total of 3 times. Statistical analyses included ANOVA and Spearman rank
37 correlation test.

38
39 Aggression scores, as determined by contact time, were significantly increased compared to the control
40 group at 8 weeks of age in both the low- (124% increase) and high- (146% increase) dose bisphenol A
41 groups. No treatment-related effects on aggression score were observed at 12 and 16 weeks of age. In the
42 low-dose group, relative (to body weight) testis weight was 10% lower than controls at 8 weeks of age
43 and 18% lower than controls at 12 weeks of age. Relative testis weight was 11% lower than control
44 values in the high-dose group at 12 weeks of age. No significant effects were observed for serum
45 testosterone levels. There were no correlations between serum testosterone levels and contact time in
46 aggression testing. The study authors concluded that prenatal bisphenol A exposure of mice resulted in
47 behavioral changes and decreased relative testis weight that was more pronounced at the lower dose.

48
49 **Strengths/Weaknesses:** Strengths are the use of 2 low dose levels and the oral route of administration.
50 The lack of husbandry information and the apparent lack of consideration of possible litter effects are
51 weaknesses.

1
2 **Utility (adequacy) for CERHR Evaluation Process:** This study is moderately useful and showed
3 increased aggression at both low doses at 8 weeks but not at 12 or 16 weeks. Testis weight was increased
4 at low dose levels.

5
6 **Laviola et al. (359)**, supported by Italian Ministry of Health, Ministry of Universities and Research, and
7 the University of Parma, examined the effect of prenatal bisphenol A exposure on *d*-amphetamine-
8 reinforcing effects in mice. **[No information was provided about feed, housing, or bedding**
9 **composition.]** CD-1 mice were trained to drink the “tocopherole”-purified corn oil vehicle through a
10 syringe. The mice were randomly assigned to groups, and 10–12/group were exposed to bisphenol A
11 **[purity not reported]** at 0 (vehicle) or 0.010 mg/kg bw by feeding from a syringe on GD 11–18 **[day of**
12 **vaginal plug not defined]**. Another group of mice was exposed to methoxychlor; those findings will not
13 be discussed. Litters were culled to 10 pups (5 ± 1 of each sex) within 12 hours of parturition. Offspring
14 were weaned and group housed with littermates of the same sex on PND 25. At 60 days of age, 3
15 offspring/sex/litter (1 sex/litter at each *d*-amphetamine dose) were subjected to conditioned place-
16 preference testing. For the test, animals were acclimated to the apparatus on the first day of testing. On
17 alternate days over a 4-day period, animals were ip injected with 0, 1, or 2 mg/kg bw *d*-amphetamine and
18 confined to one compartment of the apparatus for 20 minutes. On the other days of the 4-day period,
19 animals were injected with saline and confined in another section of the apparatus for 20 minutes. On the
20 fifth day of testing, animals were not treated and were given free access to the entire apparatus for 10
21 minutes. The amount of time spent in the compartment associated with *d*-amphetamine treatment was
22 measured. Data were analyzed by a split-plot ANOVA, in which the litter was considered the block
23 variable, and Tukey HSD test.

24
25 No differences were reported for birth weight and sex ratio at birth. **[Data were not shown by authors.]**
26 There were no significant effects of bisphenol A treatment on locomotive activity. Conditioned place-
27 preference occurred in control females following injection with either *d*-amphetamine dose, but was not
28 observed in females treated with bisphenol A. In males, both the vehicle control and the bisphenol A
29 group displayed a preference for the *d*-amphetamine-associated compartment following treatment with
30 the high *d*-amphetamine dose. Therefore, there was no change in preference following bisphenol A
31 treatment of males. The study authors concluded that prenatal bisphenol A exposure affected organization
32 of the brain dopaminergic system in female mice leading to long-term alterations in neurobehavioral
33 function.

34
35 **Strengths/Weaknesses:** The use of only 1 dose level and the small sample size are weaknesses.

36
37 **Utility (Adequacy) for CERHR Evaluation Process:** This study is slightly useful in the evaluation.

38 39 *3.2.6 Mouse—parenteral exposure only during pregnancy*

40 **Markey et al. (360)**, supported by NIH, examined the effect of prenatal bisphenol A exposure on
41 mammary gland development in mice. CD-1 mice were fed RMH 3000 rodent diet, which showed
42 negligible activity in estrogenicity testing. Caging and bedding were also reported to test negative in
43 estrogenicity assays. Dams (6–10/group) received the DMSO vehicle or bisphenol A **[purity not**
44 **reported]** at 0.000025 or 0.000250 mg/kg bw/day through a sc pump from GD 9–20 (GD 1 = day of
45 vaginal plug). **[The original publication stated that bisphenol A doses were 25 and 250 µg/kg**
46 **bw/day, but units were corrected to ng/kg bw/day in an addendum released for the study].** Doses
47 were said to decrease as dams gained weight during pregnancy. Dams were allowed to litter and offspring
48 were weaned at 19 days of age. At 10 days, 1 month, and 6 months of age, 6–10 female offspring/group
49 were killed during each time period. **[Number of litters represented was not stated but there may**
50 **have been 1 offspring/litter based on the numbers examined.]** Vaginal smears were assessed in mice
51 following puberty, and post-pubertal mice were killed during proestrus. Prior to being killed, females

3.0 Developmental Toxicity

were injected with bromodeoxyuridine, and incorporation of bromodeoxyuridine in mammary glands was determined by an immunohistochemistry method. Histological and morphometric analyses of mammary glands were also conducted. Data were analyzed by ANOVA, least significant difference test, and *t*-test.

At 1 month of age, the rate of ductal migration into the stroma was increased in the low-dose group and decreased in the high-dose group; values in the 2 treatment groups were significantly different from one another but neither dose group was significantly different from the control group. Statistically significant findings compared to the control group are summarized in Table 90. Bisphenol A treatment increased percentages of ducts and buds at 6 months of age. Bromodeoxyuridine incorporation was decreased in epithelial cells at both doses at 10 days of age, decreased in stromal cells at the high dose at 1 month of age, and increased in stromal cells at both dose levels at 6 months of age. At 1 month of age, the ratio of bromodeoxyuridine-positive epithelial to stromal cells was 4:1 in the control group, 2:1 in the 0.000025 mg/kg bw/day group, and 6:1 in the 0.000250 mg/kg/bw/day group. The percentage of alveoli containing secretory products was increased at the low dose at 6 months of age. The study authors concluded gestational exposure to low doses of bisphenol A alters timing of DNA synthesis in mammary epithelium and stroma, resulting in a histoarchitecture that is not typical for a virgin mouse.

Table 90. Effects of Prenatal Bisphenol A Exposure on Mammary Gland Development in Mice

Endpoint	Bisphenol A dose in mg/kg bw/day	
	0.000025	0.000250
10 Days of age		
Epithelial cells incorporating bromodeoxyuridine, %	↓52%	↓36%
1 month of age		
Stromal cells incorporating bromodeoxyuridine, %	↔	↓32%
6 months of age		
Duct area	↑29%	↑25%
Terminal duct area	↑237%	↑219%
Terminal end bud area	↑192%	“↑”139%
Alveolar bud area	↑288%	↑361%
Stromal cells incorporating bromodeoxyuridine, %	↑56% ^a	↑95%
Percent alveoli-containing secretory products	↑60%	↔

↑,↓ Statistically significant increase, decrease compared to controls; ↔ no statistically significant effect compared to controls. “↑” Increase identified in the text, although not statistically significant according to the data figure (3B in the study).

^aValue given in the text. The data figure (4B in the study) suggests a 2-fold increase.

From Markey et al. (360).

Strengths/Weaknesses: The administration of very low doses by subcutaneous pump and the examination of the mammary gland, a system not often studied, are strengths of this study.

Utility (adequacy) for CERHR Evaluation Process: This paper is useful and shows tissue effect at extremely low dose levels.

Markey et al. (361), supported by NIH and the Massachusetts Department of Public Health, examined the effects of prenatal bisphenol A exposure on development of the female reproductive system and mammary gland in mice. CD-1 mice were fed Purina Rodent Chow that tested as having negligible estrogenicity. Cages and bedding tested negative for estrogenicity in the E-SCREEN assay. Water was provided in glass bottles. Mice (n = 6–10/group) were administered bisphenol A at 0 (DMSO vehicle), 0.025, or 0.250 mg/kg bw/day by sc pump from GD 9 through the remainder of pregnancy (GD 1 = day of vaginal plug). Number of offspring, sex ratio, body weight, and age at vaginal opening were assessed.

3.0 Developmental Toxicity

Beginning at 3 months of age and continuing for 2 weeks, estrous cyclicity was assessed by visual examination of the external vagina and confirmation by vaginal smears. Female offspring (6–10/group) were killed at 1, 3, 4, 6, 9, and 12 months of age on the afternoon of proestrus. Reproductive organs were grossly assessed, and morphometric measurements were obtained for ovary and mammary gland.

[Although the methods section suggests that morphometric measurements were obtained at each time period of sacrifice, it does not appear that the measurements were taken at 1 and 12 months of age.] A histopathological evaluation of the ovary was conducted at 3 months of age. Reproductive organ weights were obtained at 1, 3, and 6 months of age. **[Though not stated, it is assumed that as in other studies reported from this laboratory, different litters were represented at each time period.]**

Statistical analyses included ANOVA, Kruskal-Wallis, and Mann-Whitney tests.

Statistically significant findings are reported in Table 91. Bisphenol A exposure had no significant effect on litter size or sex ratio. A significant interaction between age for body weight and treatment was reported from 2 to 12 months of age but the effect on body weight was not explained. No significant effects were observed for vaginal opening in treated mice. Significant increases were observed in percentages of 3-month-old mice with estrus/metestrus for ≥ 4 or 8 days. At 6 months of age, the incidence of fluid-filled ovarian bursae was increased in both treatment groups. Reproductive organ weights were not affected at 1 or 6 months of age, but at 3 months of age, absolute and relative (to body weight) weights of vagina were decreased in the high-dose group. The percentage of ovary tissue consisting of antral follicles was increased in the high-dose group at 3 months of age. No significant differences were observed for mammary structures at 4 months of age. At 6 months of age, the percentage of alveolar buds/lobulo-alveoli was increased in both dose groups compared to the control group. The percentage of alveolar buds/lobulo-alveoli was decreased in the low-dose group compared to control group at 9 months of age. The study authors concluded that exposure of mice to environmentally relevant doses of bisphenol A during the development of estrogen-sensitive tissues results in effects that are manifested in adulthood.

Table 91. Reproductive Effects Observed in Mice Exposed to Bisphenol A During Prenatal Development

Endpoint	Bisphenol A, mg/kg bw/day	
	0.025	0.250
3 months		
Estrus/metestrus for 4 or more days, % of animals	↑50%	↑50%
Estrus/metestrus for 8 or more days, % of animals	↑4.9-fold	↑4.3-fold
Relative vaginal weight	↔	↓26%
Antral ovarian follicles, %	↔	↑2.4-fold
6 months		
Fluid-filled ovarian bursae (control = 0)	↑ to 11.5 %	↑ to 16 %
Alveolar buds/lobulo-alveoli in mammary, %	↑3.9-fold	↑4.1-fold
9 months		
Alveolar buds/lobulo-alveoli in mammary ^a	↓38%	↔

↑,↓ Statistically significant increase, decrease compared to controls; ↔ no significant effects compared to controls.

^aValue from treatment group was estimated from a graph. From Markey et al. (361).

Strengths/Weaknesses: The administration of very low, environmentally relevant doses by subcutaneous pump and the multiple measures of female ovarian cycle and tissues are strengths of this study.

Utility (Adequacy) for CERHR Evaluation Process: This study is useful and showed significant effects, especially at the higher dose level.

3.0 Developmental Toxicity

1 **Vandenberg et al. (362)**, support not indicated, examined the effects of prenatal bisphenol A exposure on
2 mouse mammary gland development. CD-1 mice were fed Harlan Teklad 2008, which was reported to
3 contain 20 fmol/g estrogen equivalents. The type of caging and bedding used was not reported but they
4 were stated to test negative for estrogenicity in the E-SCREEN. Water was supplied in glass bottles. On
5 GD 8 (GD 1 = day of vaginal plug) mice were implanted [**assumed sc**] with osmotic pumps that delivered
6 the 50% DMSO vehicle or bisphenol A [**purity not reported**] at 0.000250 mg kg bw/day. The bisphenol
7 A dose was selected because it was thought to be environmentally relevant and shown to alter mammary
8 endpoints (360, 363). Pumps were left in place until dams were killed on GD 18. [**The number of dams**
9 **treated was not reported.**] Fetal mammary glands were mounted whole or sectioned to examine
10 mammary gland development in 36–40 offspring/group. Immunohistochemistry techniques were used to
11 measure expression of Ki67 and Bax in mammary structures from 4–8 offspring/group. Mammary
12 collagen localization was assessed using Masson Trichrome stain in 6–17 mice/group. Expression of
13 mRNA for ER α , ER β , adipocyte lipid binding protein, Col-1, and PPAR γ were measured by RT-PCR in
14 mammary glands from 4–6 offspring/group. Litter was considered in analyses by assigning 1
15 individual/litter to each group or endpoint. Statistical analyses included *t*-tests, ANOVA, Mann-Whitney
16 U non-parametric tests, and/or Chi-squared test.

17
18 Morphometric analysis revealed significantly higher ductal area and extension in the bisphenol A group
19 than in controls. In the control group, females positioned next to 2 females in utero had significantly
20 fewer branching points than females positioned next to 1 or 2 males; this difference was not observed in
21 the bisphenol A group. In fetuses that were not positioned next to a male, significantly more branching
22 points were observed in the bisphenol A than in the control group. Control females positioned next to 2
23 males had significantly larger epithelial duct area than control females not positioned next to a male; this
24 difference was not observed in the bisphenol A group. In bisphenol A-treated females positioned next to 1
25 male, ductal extension was significantly greater than in control females positioned next to 1 male.

26
27 In the bisphenol A group, epithelial cells were less rounded, more evenly spaced, and more numerous
28 than in controls. Bisphenol A did not significantly affect Ki67 (a proliferation marker) expression in
29 mammary epithelium. Lumen formation was observed in 6 of 16 control mice and 0 of 10 bisphenol A-
30 exposed mice. Significantly decreased numbers of Bax-positive (apoptotic) cells were observed in the
31 inner epithelial cord (not in contact with basement membrane) of bisphenol A-exposed than control mice.
32 Optical density of histological staining was significantly lower in the fat pad of the bisphenol A-exposed
33 than control group. Fat pads of the bisphenol A group compared to control group were found to be
34 significantly less cellular, contain more Bax-positive cells, and have more vacuoles at a distance <1 mm
35 from the epithelial compartment. Study authors interpreted the effect as increased epithelial penetration
36 and advanced maturation of fat pads. No significant differences were observed for PPAR γ or adipocyte
37 lipid binding protein mRNA expression. Density of collagen deposits was lower in the entire mammary
38 gland but higher in the periductal stroma (within 10 μ M of the epithelium) of the bisphenol A than the
39 control group. Bisphenol A exposure did affect collagen type I, ER α , or ER β mRNA expression. ER α
40 protein expression was also unaffected by bisphenol A exposure. Study authors concluded that advanced
41 maturation of fat pad and changes in extracellular matrix may be the cause of altered growth, cell size,
42 and lumen formation in mammary epithelium of mouse fetuses exposed to bisphenol A.

43
44 **Strengths/Weaknesses:** To be added.

45
46 **Utility (Adequacy) for CERHR Evaluation Process:** To be added.

47
48 **Honma et al. (364)**, supported by the Japanese Ministry of Education, Culture, Sports, Sciences, and
49 Technology, examined the effect of prenatal bisphenol A exposure on the reproductive system of female
50 mice. Mice were fed commercial diet (CE-2, CLEA, Tokyo, Japan). [**No information was provided**
51 **about bedding or caging materials.**] Ten ICR/Jcl mice/group were sc injected with bisphenol A [**purity**

3.0 Developmental Toxicity

not reported] in sesame oil at 0, 0.002, or 0.020 mg/kg bw/day on GD 11–17 (GD 0 = vaginal plug). Additional mice were injected with diethylstilbestrol at 0.02–2 µg/kg bw/day. Pups were sexed, counted, and weighed at birth. At 22 days of age, offspring were weaned and litter sizes were adjusted to 8 pups. Male and female offspring were weighed during the postnatal period. Anogenital distance was measured in males and females at 22 and 60 days of age. Females were monitored for vaginal opening. Vaginal smears were obtained for 30 days following vaginal opening. Female offspring were mated with untreated males from 90 to 120 days of age. F₂ pups were counted and sexed at birth. The litter was considered the experimental until in statistical analyses. Data were analyzed by ANOVA and Student or Welch *t*-test.

Statistically significant findings are summarized in Table 92. There were no effects on gestation duration, number of pups/litter, or sex ratio. Body weights were slightly lower in high-dose males at birth, both dose groups of females at weaning, and high-dose males and females at 60 days of age. Anogenital distance was increased in low-dose females at weaning and both dose groups of males at 60 days of age. Age of vaginal opening and 1st estrus was accelerated in the high-dose group, and body weight at vaginal opening was lower in both dose groups. Estrous cycle length was increased in both dose groups. Total days that cornified cells were present in vaginal smears was increased and total days that lymphocytes were detected was decreased in the low-dose group. In F₁ offspring there were no significant effects on mating, number of F₂ pups/litter, or sex ratio of F₂ pups. Results in mice dosed with diethylstilbestrol were similar to those observed in mice dosed with bisphenol A. The study authors concluded that prenatal exposure to low doses of bisphenol A results in early vaginal opening in mice but did not affect female reproductive function.

Table 92 . Effects in Mice Exposed to Bisphenol A During Prenatal Development

Endpoint	Dose (mg/kg bw/day)					
	0.002	0.020	BMD ₁₀	BMDL ₁₀	BMD _{1SD}	BMDL _{1SD}
Female body weight						
Weaning	↓10%	↓7%	0.065	0.017	0.088	0.021
PND 60	↔	↓4%	0.054	0.021	0.11	0.021
Male body weight						
Birth	↔	↓5%	0.054	0.020	0.031	0.015
PND 60	↔	↓6%	0.048	0.020	0.044	0.020
Anogenital distance						
Females at weaning	↑6%	↔				
Males on PND 60	↑6%	↑8%	0.035	0.020	0.035	0.020
Age at vaginal opening ^a	↔	↓1.3 days				
Body weight at vaginal opening ^a	↓10%	↓11%				
Age at 1 st estrus ^a	↔	↓1 day				
Estrous cycle length	↑1.3 day	↑1 day	0.021	0.007	0.12	0.021
Cornified cells in vaginal smear	↑3.1 days	↔	0.17	0.020	0.44	0.021
Lymphocytes in vaginal smear	↓2.2	↔	0.26	0.020	0.26	0.020

↑,↓ Statistically significant increase, decrease; ↔ no significant effect.

^aValue estimated from a graph by CERHR; data from graphs were not modeled.

From Honma et al. (364).

Strengths/Weaknesses: The use of low dose levels of bisphenol A is a strength.

Utility (Adequacy) for CERHR Evaluation Process: The study is useful and found changes in anogenital distance at the low dose level. A delay in puberty occurred at the higher dose level.

Iwasaki and Totsukawa (365), support not indicated, examined the effect of prenatal bisphenol A exposure on reproductive development of female mice. ICR mice were fed F1 diet (Funabashi, Chiba,

3.0 Developmental Toxicity

1 Japan) and housed in polycarbonate cages containing an unspecified chip bedding. On GD 7–18 (GD 0 =
2 day of copulatory plug), 6 dams/group received bisphenol A [**purity not reported**] at 0 (DMSO vehicle)
3 0.00025, 0.025, or 2.5 mg/kg bw/day by sc injection. A positive control group of mice received 100 µg/kg
4 bw/day 17β-estradiol [**route not specified**]. Dams were weighed during the study. Pups were counted and
5 sexed on PND 0, and pup viability was determined on PND 4. Pups were weaned on PND 21, and male
6 pups were killed and discarded. Female pups (24–41/group) were observed for vaginal opening. On PND
7 21, 1 pup/litter(4/group) from the low- and mid-dose group was injected with 3 µg/kg bw/day 17β-
8 estradiol for two days and then killed. Uterine weights were assessed and expression of the *ERα* gene in
9 uterus was determined using a colorimetric method. Statistical analyses included ANOVA, ANOVA on
10 ranks (Kruskall-Wallis test), and Dunnett test.

11
12 Weight gain was described as increased in all treated dams compared to control dams, but there was no
13 evidence of a dose-response relationship and statistical significance was not achieved. Pup birth weight
14 was significantly lower [**6%**] in the low-dose group compared to the control group. There were no
15 differences in litter size at birth. Pup viability on PND 4 was significantly reduced [**by 26%**] in the low-
16 dose group. Age of vaginal opening was significantly delayed by 3 days in the low-dose group, but
17 significantly accelerated by 2.2 days in the high-dose group. Following 17β-estradiol exposure, uterine
18 weight was significantly decreased [**by ~85%**] in the low-dose bisphenol A group and significantly
19 increased [**by ~29%**] in the mid-dose bisphenol A group. Although expression of *ERα* mRNA was
20 observed at 132% of control levels in the mid-dose bisphenol A group following exposure to 17β-
21 estradiol, the effect did not attain statistical significance. Expression of *ERα* gene was not detectable in
22 the low-dose bisphenol A group following 17β-estradiol exposure. No significant effects were reported in
23 mice treated with 17β-estradiol. The study authors concluded that “The levels tested in this study appear
24 to be dangerous.”

25
26 **Strengths/Weaknesses:** The use of 3 dose levels, including low doses, and the use of 17β-estradiol as a
27 positive control are strengths of this study.

28
29 **Utility (Adequacy) for CERHR Evaluation Process:** The study is moderately useful and shows the low
30 dose affecting puberty and uterine weight.

31
32 **Nikaido et al. (366)**, supported by the Japanese Ministry of Health, Labor, and Welfare examined the
33 effects of bisphenol A exposure on mammary glands and reproductive systems of mice. Outbred CD-1
34 (ICR) mice were fed NIH-07 (a low-phytoestrogen diet) and provided with water supplied in
35 polycarbonate bottles with rubber stoppers. The mice were housed in polyisopentene cages with white
36 pine chip bedding. Beginning on GD 15 (plug day not specified), mice were sc injected with 0 (DMSO
37 vehicle), 0.5, or 10 mg/kg bw/day bisphenol A (≥99% purity) or 0.5 or 10 µg/kg bw/day diethylstilbestrol
38 for 4 days. [**The control group contained 6 dams/group, but the number of dams in treated groups
39 was not clear.**] Additional groups of mice were treated with the same doses of genistein, resveratrol, or
40 zearalenone. Female pups were weaned at 21 days of age. Onset of vaginal opening was monitored.
41 Estrous cyclicity was monitored in 12 mice/group at 9–11 weeks of age. At 4, 8, 12, and 16 weeks of age,
42 6 randomly selected mice/group were weighed and killed. Ovaries, uterus, vagina, and mammary glands
43 were preserved in 10% formalin for histopathological evaluation. Differentiation of mammary structures
44 was evaluated in whole mounts. Statistical analyses included homogeneity of variance tests followed by
45 ANOVA or Kruskal-Wallis test. When *P* values were below 0.05, Fisher protected least significant
46 difference test was conducted.

47
48 Body weight gain of offspring was increased by bisphenol A treatment, and at 16 weeks of age, body
49 weight compared to controls was higher [**by ~50%**] in the low-dose group and [**by ~23%**] in the high-
50 dose group. Vaginal opening was accelerated by 1.2 days at the high-dose group. Estrous cycle length
51 was increased by 2.8 days in the low-dose group and 3 days in the high-dose group as a result of

3.0 Developmental Toxicity

1 increased time spent in diestrus. Corpora lutea were observed in all control mice at each age. No corpora
2 lutea were observed in 2 of 6 mice of the low-dose group and 3 of 6 mice of the high-dose group at 4
3 weeks of age, but all mice had corpora lutea at 4, 8, 12, and 16 weeks of age. With the exception of
4 vaginal cornification observed in mice lacking corpora lutea, no histopathological abnormalities were
5 observed in the uterus or vagina. Two of three mice with corpora lutea in the high-dose bisphenol group
6 had greater mammary alveolar differentiation compared to control mice at 4 weeks of age. No differences
7 in mammary differentiation were observed at later ages. The study authors concluded that both the high
8 and low dose of bisphenol A produced transient changes in the mammary gland and reproductive tracts of
9 mice. Transient effects on the reproductive tract and mammary gland were also observed with genistein
10 and diethylstilbestrol, while prolonged effects were induced by zearalenone.

11 **Strengths/Weaknesses:** The lack of clarity regarding sample size and the weak description of the
12 histopathology findings are weaknesses.

13 **Utility (Adequacy) for CERHR Evaluation Process:** This paper is moderately useful.

14
15 **Park et al. (367)**, support not indicated, treated ICR mice during pregnancy. Bisphenol A [**purity not**
16 **indicated**] in corn oil was given ip at dose levels of 0, 0.05, 0.5, or 5 mg/kg bw on the day of mating and
17 every 3 days for a total of 6 doses (n = 12/group). Dams were killed on GD 18 (plug = GD 0) for
18 determination of litter size, fetal weight, and sex ratio. The uterus and right ovary were removed from
19 each dam, fixed in Bouin fluid, and sections were stained with hematoxylin and eosin for light
20 microscopy. Results were analyzed with least significant difference test [**apparently on a per fetus**
21 **basis**]. Maternal weight was not altered by treatment. Fetal body weight was decreased in the high-dose
22 group by 14% for males and 12% for females. There was no effect on litter size or sex ratio. There was no
23 treatment effect on dam uterine or ovarian weight. Histopathology of the dam ovary was reportedly not
24 affected by treatment. Histopathology of the dam uterus showed thickening of the endometrium in the
25 0.05 and 0.5 mg/kg bw groups and uterine muscle damage in the 5 mg/kg bw group. [**The damage is not**
26 **otherwise described. The photomicrographs available in the report were not interpretable due to**
27 **poor reproduction quality.**] The authors concluded that bisphenol A at low doses does not produce
28 reproductive toxicity in mice. [**This paper was written in Korean with an English abstract and tables.**
29 **A translation was provided to CERHR by the American Plastics Council.**]

30
31 **Strengths/Weaknesses:** The use of 3 dose levels is a strength. The lack of information on husbandry
32 conditions, the ip dose route, and the poor presentation of histopathology results are weaknesses.

33
34 **Utility (Adequacy) for CERHR Evaluation Process:** This paper has marginal utility.

35
36 **Park et al. (368)**, support not indicated, treated ICR mice during pregnancy. Bisphenol A [**purity not**
37 **indicated**] in corn oil was given ip at dose levels of 0, 0.05, 0.5, or 5 mg/kg bw on the day of mating, and
38 every 3 days for a total of 6 doses (n = 3–6/group). Offspring were evaluated on PND 45 for body weight,
39 reproductive organ weight and histopathology, semen analysis, complete blood count, and serum
40 chemistry. [**There were 24 female and male offspring evaluated per dose group (not indicated**
41 **whether 12 of each sex). Litter of origin appears not to have been considered. No information was**
42 **provided on standardization of litters, diet, or cage/bedding materials.**] Statistical analysis was
43 performed using the least significant difference test. There was a statistically significant 6% decrease in
44 male body weight in the high-dose group; a comparable body weight decrement in female offspring was
45 not statistically significant. There were no statistically significant treatment effects on the weights of the
46 testis, epididymis, seminal vesicles, coagulating glands, uterus, or ovary. Sperm concentration, viability,
47 motility, and morphology were not affected by treatment. Blood endpoints were not affected by treatment
48 except for a statistically significant 6% increase in erythrocyte count in male offspring and a 2% decrease
49 in serum albumin in female offspring. An 11% increase in blood urea nitrogen in mid-dose female
50
51

3.0 Developmental Toxicity

1 offspring was not dose related. Histopathology of the testis and ovaries was described as unaffected by
2 treatment. Uterine intimal proliferation was described in the mid- and high-dose female offspring. **[The**
3 **histological methods were not described. The photomicrographs available in the report were not**
4 **interpretable due to poor reproduction quality.]** The authors concluded that bisphenol A at low doses
5 does not produce reproductive toxicity in mice. **[This paper was written in Korean with an English**
6 **abstract and tables. A translation was provided to CERHR by the American Plastics Council.]**
7

8 **Strengths/Weaknesses:** The inadequate description of methods, the ip dosing, and the poor presentation
9 of histology results are weaknesses of this study.

10 **Utility (Adequacy) for CERHR Evaluation Process:** This paper has marginal utility.

11 3.2.7 Mouse—oral exposure postnatally with or without prenatal exposure

12 **Nagao et al. (369)**, support not indicated, examined the effects of bisphenol A in mice following
13 exposure during different life stages. An initial study compared the sensitivity of male juvenile
14 C57BL/6N and ICR mice to 17 β -estradiol. Following sc dosing of 10 mice/strain/group with 10 μ g/kg
15 bw/day 17 β -estradiol on PND 27–48, there were no weight changes or histopathological alterations in
16 reproductive organs of ICR mice. In contrast, C57BL/6N mice exposed to 17 β -estradiol experienced
17 significant decreases in absolute and relative weights of testes, epididymides, and seminal vesicles. In
18 addition, epididymal sperm was reduced and there was increased severity of seminal vesicle and Leydig
19 cell atrophy. The study authors concluded that C57BL/6N mice are sensitive to estrogen and this strain of
20 mice was used in the remaining experiments.
21
22
23

24 Life stages examined in experiments with bisphenol A included prenatal development, adolescence, and
25 adulthood. The studies conducted during prenatal development and adolescence are described here, and
26 the study conducted during adulthood is described in Section 4.2. C57BL/6N mice were fed PLD
27 (phytoestrogen-low diet, Oriental Japan). They were housed in polycarbonate cages with wood bedding.
28 Daidzein and genistein levels were analyzed in the diet, tap water, and bedding and found to be below 0.5
29 mg/100 g. Bisphenol A (stated to be 99% pure in the study with adult mice) was administered to juvenile
30 or pregnant mice by gavage at doses of 0.002, 0.020, or 0.200 mg/kg bw/day. Control animals were
31 gavaged with 0.5% carboxymethyl cellulose **[assumed to be the vehicle]**. Juvenile males (30 /group
32 (obtained from 10 litters) were treated on PND 21–43 (day of birth not defined). At six weeks of age, 25
33 mice/group were necropsied. Ten pregnant C57BL/6N mice/group were treated on GD 11–17 (GD 0 =
34 day of vaginal plug). Fetuses were removed by cesarean section on GD 18 and that day was considered
35 PND 0. Litters were fostered to untreated dams. On PND 4, females were disposed and litters were culled
36 to 3 males. Males were weaned on PND 21 and housed individually in polycarbonate cages. At 12 weeks
37 of age, males were weighed and 25 males/group were killed and necropsied. During necropsy of males
38 that had been exposed during prenatal development or during adolescence, testes, epididymis, and
39 seminal vesicles with coagulating glands were weighed. In the study conducted in adult mice, it was noted
40 that ventral prostates were not weighed due to difficulties in obtaining only prostate and determining the
41 precise weight of the organ. Epididymal sperm counts were obtained. Histopathological examinations
42 were conducted for reproductive organs fixed in Bouin solution. For males exposed during gestation, the
43 litter was considered a single sample. Data were analyzed by Bartlett's test to determine homogeneity of
44 variance, followed by ANOVA when homogeneity of variance was obtained or Wallace-Wallace analysis
45 of ranks when variance was not homogenous. Dunnett test was used for multiple comparisons.
46

47 There were no significant effects on embryo mortality after birth, body weight gain, or terminal body
48 weight. **[Data were not shown.]** The only reproductive organ weight effect was a significant, but non-
49 dose related **[6%]** decrease in absolute seminal vesicle weight in the low-dose bisphenol A group. Organ
50 weights were not affected in males exposed during adolescence. Sperm density was unaffected by
51 bisphenol A exposure. No treatment-related lesions were observed in testes or other reproductive organs

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1 including ventral prostate. **[Data were not shown.]** The study authors concluded that low-dose bisphenol
2 A exposure of mice did not reduce sperm density or disrupt male reproductive system development.
3

4 **Strengths/Weaknesses:** Strengths are the use of 3 low dose levels, the oral route of administration, the
5 careful description of methods, the use of a low-phytoestrogen diet, and the confirmation that the strain of
6 mice used was estrogen sensitive.

7 **Utility (Adequacy) for CERHR Evaluation Process:** This study is very useful and found no effects on
8 sperm count. The only affected organ was the seminal vesicle, and the seminal vesicle weight reduction
9 was not dose-related.
10

11 **Kabuto et al. (107)**, supported by the Kagawa Prefectural College of Health Sciences, examined the role
12 of oxidative stress in bisphenol A-induced toxicity in mice. ICR mice were fed standard laboratory chow
13 containing 24% protein (MF Oriental Yeast Co., Tokyo, Japan). **[No information was provided about**
14 **bedding or caging materials.]** From 1 week prior to mating through gestation and lactation, 6
15 mice/group were given drinking water containing the 1% ethanol vehicle or bisphenol A **[purity not**
16 **reported]** at 5 or 10 µg/L. **[Based on the reported water intake of 5 mL/day and an assumed body**
17 **weight of 0.02 kg (81), it is estimated that bisphenol A intake in mice at the start of pregnancy was**
18 **0.0013 or 0.0025 mg/kg bw/day.]** Mice gave birth about 3 weeks following mating and pups were
19 housed with dams for 4 weeks. **[Based on an assumed body weight of 0.0085 kg and assumed water**
20 **intake rate of 0.003 L/day (81), it is estimated that intake of bisphenol A in weanling males was**
21 **0.0018 or 0.0035 mg/kg bw/day].** At 4 weeks of age, male pups were killed and brain, kidney, liver, and
22 testis were weighed in 8–13 mice/group. Tissues were homogenized to determine activities of superoxide
23 dismutase, catalase, and glutathione peroxidase and concentrations of glutathione and L-ascorbic acid in
24 6–8 mice/group. Tissue level of thiobarbituric acid-reactive substance, a biogenic macromolecular
25 peroxidation indicator, was measured in 6 mice/group. Data were analyzed by ANOVA followed by
26 Scheffe F test.
27

28 Significant findings are summarized in Table 93. Organ weight effects included decreased brain weight at
29 the low dose, decreased kidney weight at the high dose, and decreased testis weight at both doses.
30 **[Relative organ weights were not determined.]** In the high-dose group, thiobarbituric acid-reactive
31 substance levels were increased in brain, kidney, and testis. Changes in antioxidant enzyme levels
32 included decreased catalase activity in testis and increased glutathione oxidase activity in kidney. No
33 significant effects were observed for superoxide dismutase activity or glutathione or ascorbic acid levels
34 in any of the tissues examined. The study authors concluded that bisphenol A exposure during gestation
35 and lactation results in oxidative stress and peroxidation in offspring that ultimately lead to
36 underdevelopment of brain, kidney, and testis.
37

1 **Table 93. Effects in Male Mice Exposed to Bisphenol A During Gestation and Lactation**

Endpoint	Bisphenol A in drinking water, µg/mL	
	5	10
Organ weight		
Brain	↓6%	↔
Kidney	↔	↓18%
Testis	↓18%	↓14%
Thiobarbituric acid-reactive substance level		
Brain	↔	↑43%
Kidney	↔	↑66%
Testis	↔	↑69%
Catalase activity		
Liver	↑22%	↔
Testis	↔	↓18%
Kidney glutathione peroxidase activity	↔	↑32%

↑,↓ Statistically significant increase, decrease compared to controls; ↔ no statistically significant effects compared to controls.

Benchmark doses were not estimated because in most cases the number of animals/group was not specifically indicated and it was not known if litters were equally represented among doses.

From Kabuto et al. (107).

2
3 **Strengths/Weaknesses:** The delivery of bisphenol A in drinking water at low dose levels is a strength.
4 The testing of males only is a weakness.

5
6 **Utility (Adequacy) for CERHR Evaluation Process:** This study is useful. The changes, especially at
7 low dose levels, were small and were more consistent at higher dose.

8
9 **Takao et al. (370)**, support not indicated, examined the effects of bisphenol A exposure on expression of
10 *ERα* and *ERβ* in the testis of young mice. [No information was provided about feed, caging, or
11 bedding materials.] Three-week-old male C57BL/6 mice (n = 7/group) were administered bisphenol A
12 [purity not indicated] through drinking water at 0 (ethanol vehicle), 0.5, or 50 mg/L for 8 weeks.
13 [Assuming a weanling mouse drinks ~0.35 L/kg bw/day (81), bisphenol A intake would have been
14 ~0, 0.175, or 17.5 mg/kg bw/day.] The stability of bisphenol A was not determined, but water bottles
15 were changed 2 times a week to maintain a stable concentration of bisphenol A in drinking water. Mice
16 were killed at an unspecified period following exposure, and the testis and spleen were weighed. The
17 testis was examined for ERα- and ERβ-positive cells using an immunohistochemistry method and *ERα*
18 and *ERβ* mRNA using a semi-quantitative RT-PCR technique. Data were analyzed by ANOVA followed
19 by Fisher protected least significant difference test. Exposure to 50 mg/L bisphenol A resulted in a
20 decreased number of ERβ-positive cells and increased number of ERα-positive cells. Expression of *ERβ*
21 mRNA was decreased and expression of *ERα* mRNA was increased following exposure to 50 mg/L
22 bisphenol A. There were no differences in body weight or absolute or relative weights of testis or spleen
23 following bisphenol A treatment. The study authors concluded that differential modulation of ERα and
24 ERβ could be involved in effects observed following bisphenol A exposure.

25
26 **Strengths/Weaknesses:** The delivery of bisphenol A in drinking water and the measurement of ER in the
27 testis are strengths. The lack of information on age at sacrifice is a weakness.

28
29 **Utility (Adequacy) for CERHR Evaluation Process:** This study is marginally useful and appears to be
30 very preliminary.

31

3.0 Developmental Toxicity

1 **Matsumoto et al. (371)**, support not indicated, examined the effect of maternal bisphenol A exposure on
2 growth of offspring in mice. Mice were fed standard rodent chow (CE-2, Japan Clea). **[No information**
3 **was provided on caging and bedding materials.]** Mice of the ddY strain were exposed to bisphenol A
4 ($\geq 97\%$ purity) through feed at 0 or 1% from GD 14 through PND 7. The study authors stated that the
5 bisphenol A dose was equivalent to 1000 mg/kg bw/day. **[The number of dams treated was not**
6 **indicated. Day of vaginal plug and day of birth were not defined].** Mice delivered pups on PND 21.
7 During the postnatal period, body weight was monitored in 31 pups from the control group and 61–89
8 pups from the bisphenol A group. Serum prolactin levels were measured by RIA in 3 dams/group 4 days
9 following delivery. Pups were killed on PND 7, and stomach weight was measured. Data were analyzed
10 by Student *t*-test.

11
12 No differences were reported for live pups at birth. During the postnatal period, body weights of pups in
13 the bisphenol A group were significantly lower **[by ~40%]** than control group pups. No deaths were
14 reported for pups in the control group, but 30% of pups in the bisphenol A group died before PND 7. On
15 PND 1, milk could be seen in stomachs of pups from the control group, but not the bisphenol A group.
16 **[The number of pups evaluated for milk in stomach was not reported].** On PND 7, stomach weight
17 was significantly lower **[by 40%]** in pups from the bisphenol A than control group. Serum prolactin level
18 was significantly reduced **[by 46%]** in dams from the bisphenol A group. The authors concluded that
19 administration of a high bisphenol A dose to mice resulted in suppressed postnatal growth of offspring
20 which probably resulted from an insufficient supply of milk, which might have been due to decreased
21 prolactin secretion. **[Because of the implications of this study for lactation competence, this paper**
22 **will be discussed again in Section 4.2.]**

23
24 **Strengths/Weaknesses:** Weaknesses of the study are the difficulty in calculating bisphenol A intake, the
25 likely high exposure level, the lack of information on dam number and husbandry, and the high level of
26 pup body weight decrement and mortality.

27
28 **Utility (Adequacy) for CERHR Evaluation Process:** This study is not useful in the evaluation.

29
30 **Suzuki et al. (372)**, supported by the Japanese Ministry of Health, Labor, and Welfare and the Ministry
31 of Education, Culture, Sports, Science, and Technology conducted a study to determine the effect of
32 prenatal bisphenol A exposure on dopamine-receptor mediated actions in mice. Female ddY mice were
33 fed chow containing bisphenol A at 0.002, 0.5, or 2 mg/kg feed from mating through weaning of
34 offspring. **[No information was provided on the number of dams treated, purity of bisphenol A, or**
35 **the type of chow, bedding, or caging materials. Assuming a female mouse eats ~0.2 kg feed/kg**
36 **bw/day (81), bisphenol A intake would have been 0.0004, 0.1, or 0.4 mg/kg bw/day.]** Male offspring
37 were subjected to a series of tests **[age at testing not stated]**. In a conditioned place-preference test,
38 groups of 6–10 mice were injected with 0.5 mg/kg bw methamphetamine and placed in either the dark or
39 light area of the test apparatus for 3 days. On the other 3 days, males were injected with saline and placed
40 in the other compartment of the testing apparatus. On the 7th day, the divider in the apparatus was raised
41 and the time spent in each compartment was measured. Activity was measured in groups of 9–10 mice for
42 3 hours following injection with saline or 2 mg/kg bw methamphetamine. Dopamine-induced binding of
43 ³⁵S-guanosine-5' [γ -thio]-triphosphate in the limbic system was measured (n = 3 samples/group). Protein
44 levels of dopamine and vesicle monoamine transporters in brain were determined by Western blot (n = 6
45 samples), and mRNA levels of dopamine receptor in brain were determined by RT-PCR. Data were
46 analyzed by ANOVA with Bonferroni/Dunnett test.

47
48 In conditioned-preference testing, exposure to all 3 bisphenol A doses resulted in a significant and dose-
49 related increase in preference for compartments associated with methamphetamine exposure. **[Control**
50 **mice showed no compartment preferences while the times spent in the methamphetamine-**
51 **associated compartment were ~150, 200, and 275 seconds by animals in each respective dose group.]**

3.0 Developmental Toxicity

1 Preference for the methamphetamine compartment was eliminated by injecting the animals with
2 SCH23390, A dopamine D₁ receptor antagonist. In mice exposed to the high dose of bisphenol A, activity
3 was significantly increased [**by ~80% at peak**] compared to the control group following
4 methamphetamine challenge, and sensitization to methamphetamine-induced activity was also enhanced.
5 Dopamine-induced binding of ³⁵S-guanosine-5' [γ-thio]-triphosphate in the limbic system was potentiated
6 [**increased by ~15%; not clear if statistically significant**] and G-protein activation was increased [**by**
7 **~75%**] in mice exposed to the high bisphenol A dose. The effects on G-protein activation were eliminated
8 following injection with SCH23390 or sulpiride, a dopamine D₂ receptor antagonist. No changes were
9 observed for expression of dopamine and vesicle monoamine transporter proteins. Expression of
10 dopamine D₁ receptor mRNA was significantly up-regulated to 130% of control levels in the high-dose
11 bisphenol A group. [**For all endpoints except for conditioned preference, only the data from the**
12 **high-dose bisphenol A group was shown. It was not clear if that was the only dose tested for those**
13 **endpoints or if the high-dose data were shown because it was the only dose that resulted in a**
14 **statistically significant effect.**] The study authors concluded that “prenatal and neonatal exposure to
15 bisphenol A can potentiate...central dopamine D₁ receptor-dependent neurotransmission, resulting in
16 supersensitivity of methamphetamine-induced pharmacological actions related to psychological
17 dependence on psychostimulants.”
18

19 **Strengths/Weaknesses:** This report contains inadequate description of what was done in the study.
20

21 **Utility (Adequacy) for CERHR Evaluation Process:** This report is not useful in the evaluation process.
22

23 **Mizuo et al. (373)**, supported by the Japanese Ministry of Health, Labor, and Welfare and the Ministry of
24 Education, Culture, Sports, Science, and Technology, examined the effect of perinatal bisphenol A
25 exposure on morphine-induced rewarding effects and hyperlocomotion in mice. Testing was conducted in
26 offspring of ddY mice that received chow containing 0, 0.002, 0.5, or 2 mg bisphenol A/g feed [**0, 2, 500,**
27 **or 2000 ppm**] during gestation and the neonatal period of pup development. [**No information was**
28 **provided on the number of dams treated/group, purity of bisphenol A, or feed, caging, or bedding**
29 **materials.**] In place-conditioning testing, 6–10 offspring/group were placed in one compartment of a
30 testing apparatus following saline injection and in a second compartment of the apparatus following
31 morphine injection; on the second day, mice were given free access to both compartments and the time
32 spent in each compartment was measured. Locomotor activity was measured after injecting 9–10
33 mice/bisphenol A group with saline or 10 mg/kg bw morphine. Guanosine-5'-diphosphate binding and
34 expression of μ-opioid receptor mRNA were measured in 3 independent samples/group. Statistical
35 analyses included 2-way ANOVA with Bonferroni/Dunnett test. [**No information was given on the ages**
36 **that testing was conducted and the sex of mice tested.**]
37

38 In place-preference conditioning testing, a dose-dependent increase was observed for the time spent in the
39 compartment associated with morphine exposure and statistical significance was attained at the two
40 highest dose levels. [**The time spent in the morphine-associated compartment was ~15 seconds for**
41 **controls, 150 seconds for the mid-dose group, and 175 seconds for the high-dose group.**] Locomotion
42 in the high-dose bisphenol A group was significantly increased following morphine injection [**~130**
43 **compared to 10 activity counts in high-dose bisphenol A group compared to the control**]. Bisphenol
44 A treatment had no effect on guanosine-5'-diphosphate binding (i.e., μ-opioid receptor mediated G-
45 protein activation) or expression of μ-opioid receptor mRNA. The study authors concluded that chronic
46 exposure to bisphenol A induces morphine-induced rewarding effect and hyperlocomotion that does not
47 occur through activation of the μ-opioid receptor.
48

49 **Strengths/Weaknesses:** The wide dose range used is a strength, but this report does not include essential
50 information.
51

3.0 Developmental Toxicity

1 **Utility (Adequacy) for CERHR Evaluation Process:** This report is not useful in the evaluation process.

2
3 **Miyatake et al. (374)**, supported by the Japanese Ministry of Health, Labor, and Welfare and the
4 Ministry of Education, examined the effects of developmental bisphenol A exposure on morphine-
5 induced rewarding effects in male ddy mice. Maternal mice were orally exposed to olive oil vehicle,
6 bisphenol A at 0.003 or 200 mg/kg bw/day, or 17 β -estradiol at 3 μ g/kg bw/day by gavage. The
7 compounds were administered 3 times a day from the mating period through weaning of offspring. Seven
8 male offspring/group were examined in a place-conditioning test at 7 weeks of age. During the
9 preconditioning period, mice were placed in one compartment of a cage following injection with saline
10 and in another compartment of the cage following sc injection with morphine. During testing, the amount
11 of time spent in each compartment of the cage was measured. Statistical analyses included ANOVA
12 followed by Bonferroni/Dunnet test. Developmental exposures to either bisphenol A dose resulted in a
13 preference for the cage compartment associated with morphine exposure. Developmental exposure to
14 17 β -estradiol at 3 μ g/kg did not affect place preference. Based on the findings of this study and in vitro
15 studies described in Section 3.2.1.1, the study authors concluded that bisphenol A alters dopamine
16 responsiveness in mouse neurons and astrocytes, which could potentially contribute to development of
17 psychological dependence on drugs of abuse.

18
19 **Strengths/Weaknesses:** It is a weakness that only 2 doses were used, 1 very low and 1 high. Both had
20 similar effects.

21
22 **Utility (Adequacy) for CERHR Evaluation Process:** This report is moderately useful.

23
24 **Ryan and Vandenberg (375)**, supported by North Carolina State University and EPA, evaluated the
25 effects in mice of prenatal and postnatal exposure to bisphenol A on sexually dimorphic behaviors.
26 C57BL/6 mice were maintained in polycarbonate cages (checked frequently for condition) with chip
27 bedding and were given Purina 5001 chow. Females were mated and the day a vaginal plug was identified
28 was considered GD 1. Beginning on GD 3, dams were treated with bisphenol A 2 or 200 μ g/kg bw/day,
29 ethinyl estradiol 5 μ g/kg bw/day, or the tocopherol-stripped corn oil vehicle. The dose was placed in the
30 back of the throat with a gavage needle. Daily dosing was continued to PND 21, when pups were weaned.
31 One female per litter was randomly selected for behavioral testing and was ovariectomized. Pup
32 anogenital distance was measured at weaning. Non-ovariectomized mice were checked for vaginal
33 opening and vaginal smears taken daily thereafter. Puberty was defined as the first day on which cornified
34 cells were detected in 4–7 females/group. Fourteen mice/treatment group were tested in an elevated plus
35 maze and a light-dark preference chamber. Sixteen mice/treatment group were tested in a radial arm maze
36 and a modified Barnes maze. Testing occurred 2 weeks after ovariectomy. Statistical analysis used
37 ANOVA with post-hoc Student *t*-test. The radial arm and Barnes mazes were run for 5 consecutive days
38 and a repeated measures design was added to the ANOVA.

39
40 There was no effect of treatment on anogenital distance or anogenital distance divided by body weight.
41 Other results are summarized in Table 94. Puberty was advanced by exposure to ethinyl estradiol or the
42 high dose of bisphenol A. The results of the elevated plus and light-dark preference tests led the authors
43 to conclude that bisphenol A and ethinyl estradiol increased anxiety. The improved performance in the
44 radial arm and Barnes mazes led the authors to conclude that ethinyl estradiol masculinized spatial ability.
45 **[The results from the elevated plus maze also suggest masculinization of behavior, because males**
46 **show more “anxiety” in this paradigm.]** Bisphenol A 200 μ g/kg bw/day resulted in a decrease in errors
47 on earlier trials than the control in the radial arm maze, but this effect was not characterized by the
48 authors as providing strong evidence of an alteration in spatial memory.

1 **Table 94. Behavior of Female Mice after Gestational and Lactational Exposures**

Endpoint ^a	Bisphenol A, µg/kg bw/day		Ethinyl estradiol
	2	200	
Puberty onset	↔	↓4.5 days	↓6.25 days
Time in open arms of plus maze	↔	“↓” 41% (<i>P</i> = 0.06)	↓73%
Time in light part of light/dark preference box	↔	↓52%	↓69%
Errors in radial arm and Barnes mazes	↔	↔	↓

^aThe size of the difference from control was estimated from graphs.

↓ Statistically significant decrease from control value; ↔ no statistical difference from control value, “↓” Decrease identified by authors although statistical difference from control not shown.

From Ryan and Vandenberg (375)

2
3 **Strengths/Weaknesses:** Selection of established measurements of sexually dimorphic behaviors is a
4 strength; however, behavioral evaluations were conducted only on ovariectomized females (at 2 weeks
5 post-surgery). These data were then interpreted with respect to established dimorphic patterns as opposed
6 to concurrent assessments of performance in males or intact females.

7
8 **Utility (Adequacy) for CERHR Evaluation Process:** This study is useful. The interpretation of
9 masculinizing effects on behavior is somewhat limited by the absence of concurrent data from males.

10
11 **Tyl et al. (376)**, sponsored by the American Plastics Council, conducted a 2-generation GLP study of
12 bisphenol A in CD-1 mice. **[This study is discussed in detail in Section 4.2.3.2. Results relevant to**
13 **developmental toxicity will be briefly presented here.]** Mice were fed Purina Certified Ground Rodent
14 Diet No. 5002 containing 177–213 ppm genistein, 173–181 ppm daidzein, and 39–55 ppm glycitein. Mice
15 were housed in polypropylene cages with Sani-Chip® bedding. F₀ and F₁ mice (28 sex/group/generation)
16 were fed diets containing bisphenol A (99.70–99.76% purity) at 0.018, 0.18, 1.8, 30, 300, or 3500 ppm.
17 Target intakes were 0.003, 0.03, 0.3, 5, 50, or 600 mg/kg bw/day. The study authors estimated bisphenol
18 A intake in males at 0.0024–0.0038, 0.024–0.037, 0.24–0.37, 3.98–6.13, 39.1–60.8, or 529–782 mg/kg
19 bw/day. Bisphenol A intakes (in mg/kg bw/day) by females were estimated at 0.0030–0.0041, 0.030–
20 0.042, 0.32–0.43, 5.12–7.12, 54.2–67.8, 653–910 during the pre-mating period; 0.0027–0.0029, 0.027–
21 0.028, 0.28–0.29, 4.65–4.80, 47.0–48.6, 552–598 during the gestation period; and 0.0087–0.0063, 0.062–
22 0.091, 0.61–0.89, 10.4–15.1, 103.2–146.4, 1264–1667 during the lactation period. In each generation,
23 there were 2 vehicle controls groups with 28 mice/sex/group. A positive control group was given feed
24 containing 17β-estradiol at 0.5 ppm (target intake of 0.08 mg/kg bw/day). Homogeneity, stability, and
25 concentration of bisphenol A in feed were verified. Exposure of F₀ mice began at ~6 weeks of age.
26 Exposure of F₁ animals began at weaning, although it was noted that pups began eating the dosed feed in
27 the late lactation period. F₀ and F₁ mice were fed the bisphenol A-containing diets for a minimum of 8
28 weeks prior to mating and during a 2-week mating period. Exposures of females continued through the
29 gestation and lactation period.

30
31 Live F₁ and F₂ pups and litters at birth, sex ratio, and survival during the lactation period were not
32 affected and there were no clinical or gross signs of toxicity in F₁ or F₂ offspring. A non-dose-related
33 decrease in PND 21 survival index and lactational index (pups surviving on PND 21/PND 4) was
34 described in F₂ pups of the 300 ppm group. **[The biological significance of the effect was not discussed**
35 **by the study authors, but because the effect was not dose-related it is unlikely to be of biological**
36 **significance.]** In F₁ pups from the 3500 ppm group, body weights were reduced during PND 7, 14, and 21
37 in F₁ females and both sexes combined and on PND 7 and 21 in F₁ males. An increase in male pup body
38 weight observed on PND 7 in the 1.8 ppm group was not considered to be treatment related by the study
39 authors because no dose-response relationship was observed. There was no effect on anogenital distance
40 in F₁ or F₂ males or females on PND 0. Anogenital distance was also unaffected in F₂ males and F₁ and F₂
41 females on PND 21. Anogenital distance adjusted for body weight was reduced in F₁ males from the 300

3.0 Developmental Toxicity

1 and 3500 ppm groups on PND 21. Based on the lack of effect on anogenital distance at birth and
2 inconsistencies between generations, the study authors did not consider the decreases in anogenital
3 distance in F₁ males to be treatment-related. An increase in anogenital distance in F₂ females from the
4 0.018 ppm group on PND 0 was not considered to be treatment related by the study authors. Preputial
5 separation (absolute age and adjusted for body weight on day of acquisition) was delayed in parental and
6 retained F₁ males of the 3500 ppm group. When adjusted for PND 30 body weight, preputial separation
7 was delayed in retained but not parental F₁ males from the 3500 ppm group. Body weights on day of
8 vaginal opening were lower in F₁ females from the 3500 ppm group. Day of vaginal opening was
9 accelerated in the 3500 ppm group if adjusted for PND 21 body weight, but not body weight on the day of
10 acquisition. Due to the lack of effect when adjusted for body weight on day of acquisition, the study
11 authors did not consider effects on vaginal opening to be treatment related.

12
13 Dose-related organ weight changes in F₁ weanlings that were considered to be treatment-related by study
14 authors included decreased absolute and relative (to body or brain weight) spleen and paired testes
15 weights at 3500 ppm. Treatment-related absolute organ weight changes in F₂ weanlings included
16 decreased weights of spleen, paired testes, and seminal vesicles with coagulating glands in the 3500 ppm
17 group. Changes in organ weights relative to body weight in F₂ weanlings included decreased spleen
18 weight in males and females and increased relative left kidney weight in 3500 ppm males. Treatment-
19 related changes in organ weight relative to brain weight in F₂ weanlings were decreased spleen weight in
20 both sexes and decreased paired testes weight at 3500 ppm and seminal vesicles with coagulating glands
21 at 300 and 3500 ppm. Other organ weight effects (e.g, affecting epididymides, thymus, brain, ovaries,
22 and/or uterus with cervix and vagina weights) were not considered to be dose-related due to lack of dose-
23 response relationships or no consistent effects across generations. The study authors reported no gross
24 findings in F₁ or F₂ weanlings. **[Although not clear because the number of animals examined for
25 gross testicular effects was not reported in Tables 23 and 49 of the study, it appeared that the
26 incidence of undescended bilateral testes may have been increased in F₁ and F₂ weanling males of
27 the 3500 ppm group.]** The incidence of hepatic cytoplasm alteration (clear hepatocellular cytoplasm,
28 slightly more basophilic cytoplasm, and/or minute vacuoles) was apparently increased in F₁ males from
29 the 300 and 3500 ppm groups and F₁ females and F₂ males from the 3500 ppm group. The incidence of
30 seminiferous tubule hypoplasia was increased in F₁ and F₂ weanlings from the 3500 ppm group. **[Another
31 histopathological finding that appeared to be possibly increased in weanlings from the 3500 ppm
32 group was unilateral hydronephrosis in F₁ males. It did not appear that histopathological data were
33 statistically analyzed.]**

34
35 The study authors identified bisphenol A NOELs of 30 ppm (~5 mg/kg bw/day) for systemic effects and
36 300 ppm (~50 mg/kg bw/day) for developmental toxicity. **[The lowest benchmark doses were obtained
37 from F₁ body weight data on PND 21: BMD₁₀ 548 mg/kg bw/day, BMDL₁₀ 267 mg/kg bw/day,
38 BMD_{1SD} 580 mg/kg bw/day, BMDL_{1SD} 370 mg/kg bw/day.]**

39
40 **Strengths/Weaknesses:** Strengths include the large number and range of doses examined, the rigor with
41 which the study was performed, the large sample size in each group, the number of additional animals
42 per litter that were retained and examined, the use of a concurrent estrogenic positive control group, and
43 the thoroughness of the histologic evaluation. Weaknesses might include that brain biochemistry and
44 other CNS metrics were not examined, and that statistics was not performed on some histopathology
45 findings.

46
47 **Utility (Adequacy) for CERHR Evaluation Process:** This exceptional study is very useful for the
48 evaluation process, and will carry significant weight in the evaluation of structural, histogenic, and
49 fertility endpoints.

3.0 Developmental Toxicity

3.2.8 Mouse—parenteral exposure postnatally with or without prenatal exposure

3.2.8.1 Female reproductive endpoints

Suzuki et al. (377), supported by Japanese Ministry of Education, Culture, Sports, Sciences, and Technology, the Special Coordination Funds of Science and Technology Agency of the Japanese Government, and the Japanese Ministry of Health, Labor, and Welfare, conducted a study to examine the effects of bisphenol A exposure on the reproductive system of the female mouse. Two sets of studies were conducted, one with prenatal exposure, and one with postnatal exposure. In both studies, ICR/Jcl strain mice were a fed commercial diet (CE-2, CLEA, Tokyo, Japan). **[No information was provided about bedding or caging materials.]** Bisphenol A **[purity not reported]** was administered by sc injection in sesame oil vehicle. For histological examinations, organs were fixed in Bouin solution. Parametric data were analyzed by ANOVA, with post hoc Student *t*-test or Welch *t*-test. Data expressed as proportions were analyzed by Fisher exact probability test. For exposures occurring in the prenatal period, the litter was maintained as the statistical unit by obtaining each mouse from a different litter.

In the prenatal exposure study, mice were administered bisphenol A by sc injection at 0 (vehicle), 10, or 100 mg/kg bw/day on GD 10–18 (day of vaginal plug = GD 0). Other groups of mice were treated with diethylstilbestrol at 0.0067–67 µg/kg bw/day during the same period. **[Numbers of dams treated were not specified.]** On GD 19, fetuses were removed by cesarean section, weighed, adjusted to 7 pups/litter **[numbers for each sex not indicated]**, and fostered to untreated mothers. Pups were weaned at 22 days of age. Some pups were ovariectomized at 30 days of age, and some were killed at 30 or 40 days of age for histological examination of reproductive organs, polyovular follicle numbers, corpora lutea numbers, and mitotic index in uterine and vaginal cells. In the remaining pups, vaginal smears were examined from 41 to 70 days of age. Fertility was then assessed by mating the mice with untreated males (2 or 3 females/male). Offspring were counted and sexed. The authors stated that 2 or 3 pups/litter were used in each analysis. Data tables list the sample size as 8–11/group/time period for the bisphenol A and control groups.

Bisphenol A treatment did not affect the histology of the uterus or vagina in ovariectomized mice. The study authors stated there was no evidence of increased mitogenicity compared to controls in uterine cells of intact or ovariectomized mice exposed to bisphenol A. **[Figure 3 of the study indicated a higher mitotic index in epithelial cells of ovariectomized mice of the high-dose group.]** Mitotic indices were significantly lower in stromal cells of intact mice of both dose groups and in glandular cells of the low-dose group. There was no increase in mitogenicity of vaginal cells compared to the control group; in intact mice, the mitotic index was lower than control values in vaginal epithelial cells of the high-dose group and stromal cells of the low-dose group. Number of vaginal epithelial layers was increased in both bisphenol A dose groups of intact mice compared to control mice. No effect was reported for uterine or vaginal epithelial stratification. There were no effects on numbers of polyovular follicles. **[Data were not shown by study authors.]** The number of mice with corpora lutea at 30 days of age was significantly reduced in the low-dose group (4 of 9 mice in low dose group compared to 7 of 9 mice in control group). Estrous cyclicity was not affected by bisphenol A treatment. In mating studies, bisphenol A exposure did not affect the number of mice giving birth, number of fetuses/litter, or sex ratio. Several effects were observed in mice prenatally exposed to diethylstilbestrol, and most of the effects occurred at the high dose of 67 µg/kg bw/day. In the high-dose diethylstilbestrol group, there were changes in vaginal and uterine histology, increases in mitotic indices in vaginal and uterine cells of ovariectomized animals, vaginal stratification and increased layers of epithelial cells in ovariectomized animals, disrupted estrous cycles, and complete infertility. The number of mice with corpora lutea at 30 days was decreased at the two highest diethylstilbestrol doses (≥ 6.7 at µg/kg bw/day).

In the postnatal exposure experiment, female mice (1.5 g bw) were sc injected with bisphenol A at 0.015 or 0.150 mg/pup/day or diethylstilbestrol at 0.3 or 3 µg/pup/day for 5 days, beginning on the day of birth.

3.0 Developmental Toxicity

1 **[The number of animals treated was not stated. Based on body weights provided by authors,**
2 **bisphenol A doses were estimated at 10 and 100 mg/kg bw/day; diethylstilbestrol doses were**
3 **estimated at 200 and 2000 µg/kg bw/day.]** Two-thirds of mice were ovariectomized at 30 days of age
4 and then killed at 30, 40, or 90 days of age for histological examination of reproductive organs. Numbers
5 of polyovular follicles were determined at 30 days of age, and number of corpora lutea were counted at 30
6 and 90 days of age. Estrous cyclicity was monitored in the remaining mice at 61 to 90 days of age. The
7 90-day-old mice were sc injected with 5 mg/kg bw colchicine and killed 5 hours later. Mitotic rates of
8 uterine and vaginal cells were determined, and histological examinations of reproductive organs were
9 conducted. Sample sizes were 6–17/group/time period in analyses conducted in mice exposed postnatally.

10
11 Vaginal stratification was observed at 40 days of age in 4 of 7 ovariectomized mice of the high-dose
12 bisphenol A group, which was higher than in the control. The incidence of vaginal stratification in 90-
13 day-old ovariectomized mice of the high-dose group (4 of 10) did not attain statistical significance
14 compared to control. In ovariectomized mice, significant increases in the mitotic rate compared to
15 controls were observed in uterine stromal cells and vaginal epithelial cells at the high dose. The number
16 of vaginal epithelial layers was also increased in the high-dose bisphenol A group (~4 layers in treated
17 group compared to 3.5 layers in control group). There were no significant changes in estrous cycles or
18 number of mice with corpora lutea. In 30-day-old mice of the high-dose group, significant increases were
19 observed in the number of mice with polyovular follicles (15 of 17 in exposed group compared to 6 of 15
20 in control group) and the numbers of polyovular follicles/mouse (mean ± SE: 0.8 ± 0.2 in the exposed
21 group and 0.2 ± 0.1 in control group); polyovular follicles contained 2 oocytes in the control and
22 bisphenol A groups. Effects observed in mice treated with both doses of diethylstilbestrol included
23 increased stratification of vaginal cells in ovariectomized mice at 40 and 90 days of age, increased mitotic
24 rates of vaginal and uterine cells in ovariectomized mice, disrupted estrous cycles, and increased
25 polyovular follicles. The study authors concluded that high doses of bisphenol A induce ovary-
26 independent vaginal stratification and polyovular follicles when administered during postnatal but not
27 prenatal development.

28
29 **Strengths/Weaknesses:** The use of diethylstilbestrol as a positive control is a strength. The use of
30 relatively high doses by sc injection is a weakness. Few effects (histology, mitotic activity) were found in
31 the prenatal study but additional effects (e.g., polyovular follicles) were found in the in postnatal study..

32
33 **Utility (Adequacy) for CERHR Evaluation Process:** This study is useful in the evaluation.

34
35 **Nikaido et al. (378)**, supported by the Japanese Ministry of Health, Labor, and Welfare, examined the
36 effects of bisphenol A exposure on the development of the reproductive system in female mice. Mice
37 used in this study were housed in polyisopentene cages with white pine chip bedding. The mice were fed
38 a low-phytoestrogen diet (NIH-07 PLD; Oriental Yeast Co.) and provided water in polycarbonate bottles
39 with rubber stoppers. At 15 days of age, 17–24 female CD-1 mice/group were sc injected with DMSO
40 vehicle, 10 mg/kg bw/day bisphenol A (≥99% purity), or 10 µg/kg bw/day diethylstilbestrol for 4 days.
41 Additional groups were dosed with other compounds, but those results will not be discussed. **[No**
42 **information was provided on the numbers of litters represented.]** Mice were weaned at 21 days of
43 age. Body weights were measured weekly. Day of vaginal opening was determined and estrous cyclicity
44 was assessed over 21-day periods beginning at 5, 9, and 21 weeks of age. Six mice/group/time period
45 were killed and necropsied at 4, 8, 12, and 24 weeks of age. **[In contrast to the Materials and Methods**
46 **section, there was no mention of animals killed at 12 weeks of age in the abstract or results section**
47 **of the study.]** Ovaries, uteri, vaginas, and inguinal mammary glands were fixed in 10% neutral buffered
48 formalin. Histopathological analyses were conducted of the ovary, uterus, and vagina. Mammary glands
49 were examined as whole-mount preparations. It appears that all endpoints were assessed in every mouse.
50 Statistical analyses included homogeneity of variance analysis and ANOVA or Kruskal-Wallis test. If

3.0 Developmental Toxicity

1 statistical significance was obtained, data were further analyzed by Fisher protected least significant
2 difference test.

3
4 Exposure to bisphenol A resulted in no effects on body weight gain, age of vaginal opening, estrous
5 cyclicity, histopathological changes in the uterus or vagina, or growth or development of the mammary
6 gland. At 4 weeks of age, 33% of mice in the control group, 83% of mice in the bisphenol A group, and
7 100% of mice in the diethylstilbestrol group lacked corpora lutea. **[It appears that the study authors
8 considered the lack of corpora lutea to be normal based on the age of mice.]** No effects on corpora
9 lutea numbers or numbers of polyovular follicles were observed at later ages. Mice treated with
10 diethylstilbestrol experienced accelerated vaginal opening and increased time in estrus. In their
11 conclusion, the study authors reiterated the lack of effects in the bisphenol A group.

12
13 **Strengths/Weaknesses:** The use of diethylstilbestrol as a positive control is a strength, but the lack of
14 information on sample size or results later in life detracted from the utility of the study.

15
16 **Utility (Adequacy) for CERHR Evaluation Process:** This paper is not useful in the evaluation.

17
18 **Markey et al. (379)**, supported by NIH, examined the effects of perinatal bisphenol A exposure on
19 reproductive development in mice. CD-1 mice were fed Purina rodent chow that tested “negligible for
20 estrogenicity in the E-SCREEN assay.” Cages and bedding tested negative for estrogenicity in the E-
21 SCREEN assay. Tap water was supplied in glass bottles. From GD 9 (GD 1 = day of vaginal plug)
22 through PND 4, 6–10 mice/group were exposed to bisphenol A **[purity not reported]** at 0 (DMSO
23 vehicle), 0.000025, or 0.000250 mg/kg bw/day through a sc pump. Offspring were culled to 10/litter on
24 PND 7 and weaned on PND 20. One pup/litter from 6–10 litters/treatment group was killed on the day of
25 proestrus at 3 months of age. The uterus and vagina were weighed and subjected to morphometric
26 analysis. The uterus was also assessed for cell proliferation by bromodeoxyuridine (BrdU) incorporation,
27 apoptosis by TUNEL method, and expression of ER α and progesterone receptor by an immunostaining
28 procedure. Data that were normally distributed and showed homogeneity of variance were analyzed by
29 ANOVA and least significant difference test. Other data were analyzed by Kruskal-Wallis and Mann-
30 Whitney *U* test.

31
32 Statistically significant findings are summarized in Table 95. Significant effects observed in 3-month-old
33 offspring exposed to the high dose included decreased absolute and relative (to body weight) vaginal
34 weight, decreased volume of uterine lamina propria, and increased percentage of proliferating uterine
35 glandular epithelial cells. In mice of both dose groups, there were significant increases in expression of
36 ER α and progesterone receptor in uterine luminal epithelial cells; levels of both receptors were also
37 increased in the subepithelial stroma. No treatment effects were observed for apoptosis in uterine luminal
38 and glandular epithelial cells. No treatment effects were observed for vaginal morphometry or cell
39 proliferation. The study authors concluded that environmentally relevant doses of bisphenol A affect the
40 development of the genital tract at the gross and cellular level in the female offspring of mice exposed
41 during pregnancy.

1 **Table 95. Uterine and Vaginal Effects in Mice Exposed Perinatally to Bisphenol A**

Endpoint	Dose, mg/kg bw/day					
	0.000025	0.000250	BMD ₁₀	BMDL ₁₀	BMD _{1SD}	BMDL _{1SD}
Relative vaginal weight	↔	↓26%	0.00011	0.000075	0.00022	0.000129
Absolute volume of uterine lamina propria	↔	↓30%	0.000098	0.000059	0.00025	0.00014
Uterine epithelium						
Glandular cells incorporating BrdU, % ^a	↔	↑3-fold				
Luminal cells expressing ER α , % ^a	↑4.5-fold	↑5.3-fold				
Luminal cells expressing progesterone receptor, % ^a	↑13.6-fold	↑12.5-fold				

↑,↓ Statistically significant increase, decrease compared to controls; ↔ no significant effect compared to controls.

^aBenchmark doses not calculated because number of mice examined in each dose group was not reported.

From Markey et al. (379).

2
3 **Strengths/Weaknesses:** The use of sc pumps to deliver low doses of bisphenol A from GD 9 to PND 4 is
4 a strength. The authors found strong effects (higher in high dose group) on uterine epithelium mitosis and
5 receptor activity for 17 β -estradiol and progesterone.

6
7 **Utility (Adequacy) for CERHR Evaluation Process:** This study is very useful.

8
9 **Muñoz-de-Toro et al. (363)**, supported by NIH and National University of Litoral (Argentina), examined
10 the effect of perinatal bisphenol exposure on mammary gland development in mice. Food, caging, and
11 bedding material were reported to test negligible for estrogenicity in the E-SCREEN. Water was provided
12 in glass bottles. CD-1 mice (n = 6–10/group) were implanted with osmotic pumps designed to deliver
13 bisphenol A at 0 (DMSO vehicle), 0.000025, or 0.000250 mg/kg bw/day from GD 9 (GD 1 = day of
14 vaginal plug) through PND 3 (not defined). Offspring were culled to 10 pups/litter on PND 7. One female
15 offspring/litter, from 6–10 litters/group, was killed on PND 20 and 30 and at 4 months of age. The 4-
16 month-old mice were killed on proestrus. Another group of mice [number not specified] was killed on
17 the first proestrus. Mammary glands were collected for evaluation of mammary structures at 20 and 30
18 days and 4 months of age and day of first proestrus. Mammary glands were also collected from 30-day-
19 old mice for analysis of DNA synthesis by BrdU incorporation, expression of ER α and progesterone
20 receptor using immunohistochemistry techniques, apoptosis by TUNEL method, and *Wnt4* mRNA by
21 RT-PCR. Plasma 17 β -estradiol levels were measured in mice killed at first proestrus. In an experiment to
22 monitor response to 17 β -estradiol, one pup/litter (n = 10/group) was ovariectomized at 25 days of age and
23 implanted with a sc pump supplying vehicle or 0.5 μ g 17 β -estradiol/kg bw/day on PND 25–35. Mice
24 were killed following 17 β -estradiol treatment for examination of mammary structures. Statistical analyses
25 included ANOVA and Dunn post hoc test. If the data were not normally distributed, statistical analyses
26 were done by Kruskal-Wallis and Mann-Whitney test.

27
28 Statistically significant findings are summarized in Table 96. In 30-day-old mice, bisphenol A exposure
29 increased numbers of terminal end buds at both doses and area of terminal end buds at the high dose.
30 Percentages of apoptotic cells were decreased on PND 30 in mice from both bisphenol A dose groups.
31 The percentage of stromal cells undergoing proliferation on PND 30 was reduced in the high-dose

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1 bisphenol A group. The number of epithelial cells expressing progesterone receptors was increased in
 2 both dose groups on PND 30, but there were no treatment-related changes in ER α receptor expression.
 3 Clusters of progesterone receptors were often observed in the ductal epithelium of bisphenol A-treated
 4 mice. Slopes of the correlation between age of first proestrus and mammary length were significantly
 5 reduced in the high-dose group, suggesting slower ductal invasion of stroma. There were no significant
 6 differences in plasma 17 β -estradiol levels in mice killed at first proestrus. Trends for increasing
 7 expression of mRNA for *Wnt4*, a mediator of lateral branching downstream from progesterone receptors,
 8 did not attain statistical significance. The number of lateral branches in mammary gland at 4 months of
 9 age was significantly increased at the low but not the high dose. In mice exposed to the high dose of
 10 bisphenol A during perinatal development and 17 β -estradiol during postnatal development compared to
 11 mice who were exposed to 17 β -estradiol but not bisphenol A, there were increases in numbers, area, and
 12 size of terminal end buds, terminal end bud numbers/ductal area, and terminal end bud area/ductal area.
 13 The study authors concluded that “. . . perinatal exposure to environmentally relevant [bisphenol A] doses
 14 results in persistent alterations in mammary gland morphogenesis.”
 15

16 **Table 96. Effects of Bisphenol A on Mammary Glands of Mice Exposed During Prenatal and**
 17 **Postnatal Development**

Endpoint	Dose, mg/kg bw/day	
	0.000025	0.000250
PND 30		
No. terminal end buds/ductal area	“↑”47% (<i>P</i> = 0.054)	↑58%
Area terminal end buds/ductal area	↔	↑1581%
Apoptotic cells, % ^a	↓85%	↓70%
Stromal cells incorporating BrdU, %	↔	↓50%
Epithelial cells expressing progesterone receptors ^b	↑14%	↑14%
Correlation between age of first estrous and mammary length	↔	↓
Length of ductal tree in mice that had first proestrus at 34 days of age or later	↔	↓56%
Number of lateral mammary gland branches at 4 months of age.	↑	↔
No. terminal end buds following postnatal estradiol exposure ^b	↔	↑82%
Terminal end bud area following postnatal estradiol exposure ^b	↔	↑95%
No. terminal end buds/ductal area following postnatal estradiol exposure ^b	↔	↑102%
Terminal end bud area/ductal area following postnatal estradiol exposure ^b	↔	↑114%

↑,↓ Statistically significant increase, decrease compared to controls; ↔ no significant effect compared to controls. “↑” Increase identified by authors but not statistically significant.

^aValues were estimated from a graph by CERHR.

^bValues were statistically significant compared to animals that were not exposed to bisphenol A but were exposed to 17 β -estradiol in the postnatal period.

From Muñoz-de-Toro (363).

18
 19 **Strengths/Weaknesses:** This study was a good followup on the study of Markey et al. (379) and tested
 20 the same doses using a similar schedule for effects on mammary tissue. The study found significant
 21 effects, especially in high dose mice. The study used relevant doses with long-term perinatal exposure.
 22

23 **Utility (Adequacy) for CERHR Evaluation Process:** This study is very useful.
 24

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3.2.8.2 Male reproductive endpoints

Nakahashi et al.(380), supported by the Japanese Ministry of Education, Science, Sports, and Culture, examined the effect of neonatal bisphenol A exposure on adult sperm count in mice. On the first 5 days of life, 10–15 neonatal SHN mice/group were injected [route not indicated] with sesame oil/DMSO vehicle or with bisphenol A [purity not reported] in sesame oil at 0.0005 or 0.050 mg/day. [Assuming a neonatal mouse weights 2 g, the mice received doses of 0.25 and 25 mg/kg bw/day]. A group of 12 mice received 0.050 mg/day bisphenol A in sesame oil in combination with 100 IU retinol acetate in DMSO vehicle. In a second exposure protocol, pregnant mice were fed a vitamin A-deficient diet (Low vitamin A diet; Clea Japan) from 3 days prior to gestation to PND 5. After PND 5, the dams were fed commercial diet (CE-7, Clea Japan). On the first 5 days of life, their pups (n = 7–9/group) were injected with bisphenol A at 0 (sesame oil) or 0.0005 mg/day. Male offspring from both studies were weaned at 20 days of age and fed the CE-7 diet. Mice were killed at 14 weeks of age and epididymal sperm counts were obtained. [No information was provided about caging and bedding materials. Numbers of litter represented were not indicated. Procedures for statistical analyses were not discussed.]

A 35% reduction in sperm counts was observed in mice from the 0.050 mg/day group compared to the control group. A significant reduction in sperm counts was not observed in the group co-treated with 0.050 mg/day bisphenol A and retinol acetate. Administration of a vitamin A-deficient diet to dams had no effect on sperm counts in their offspring, but sperm counts were reduced in mice born to mothers fed a vitamin A-deficient diet and injected with 0.0005 mg/day bisphenol A in the neonatal period. The study authors concluded that vitamin protects infants from the effects of environmental xenoestrogens.

Strengths/Weaknesses: The bisphenol A doses are difficult to calculate but probably ~0.25 and 25 mg/kg. The lack of husbandry and statistical information is a weakness.

Utility (Adequacy) for CERHR Evaluation Process: This paper is slightly useful. Although the study is weak, it suggests that vitamin A may alleviate the effect of bisphenol A. This finding may be worth following up but is not important at this time.

Aikawa et al. (381), supported by the Japanese Ministry of Education, Science, Sports, and Culture, examined the effects of neonatal bisphenol A exposure on sperm endpoints in adult mice. Unless otherwise specified, dams were fed CE-7 and CA-1 (Clea Japan Inc). [No information was provided about caging or bedding materials.] In the first experiment, SHN mice were sc injected with bisphenol A, bisphenol A plus retinol acetate, or vehicle for 5 days beginning on the day of birth. Doses of each compound were 0.5 or 50 µg/day bisphenol A [purity not reported] (n = 10–14/group), 50 µg bisphenol A plus 100 IU retinol acetate/day (n = 5), and vehicle control (sesame oil for bisphenol A and or DMSO for retinol acetate; n = 11). [Assuming a neonatal mouse weighs 2 g, these bisphenol A doses would be 0.25 and 25 mg/kg bw/day.] In another group, pregnant mice were fed a low vitamin A diet from 3 days prior to gestation to PND 5 and were fed a normal vitamin A-containing diet (CE-7 and CA-1) beginning on the 6th day following parturition [number/group not stated]. Pups born to those dams (n = 7–8/group) were sc injected with 0.5 µg/day bisphenol A or vehicle for 5 days, beginning on the day of birth. In all groups, mice were weaned at 3 weeks of age, individually housed at 8 weeks of age, and killed at 10 weeks of age. Sperm were collected for analysis of motility and abnormalities. In pups not born to vitamin A-deprived dams, testes were fixed in formalin for histopathological evaluation. Data were analyzed by ANOVA and Fisher least significant difference test.

Sperm motility was significantly reduced in mice injected with 50 µg/day bisphenol A (~25 vs. 50% in controls) but was not affected in mice exposed to 50 µg/day bisphenol A plus retinol acetate. Sperm motility was not affected in mice born to mothers fed a normal diet and exposed to 0.5 µg/day bisphenol. Compared to the vehicle control group born to mothers fed a normal diet, the mice born to mothers fed a vitamin A-deficient diet and injected with 0.5 µg/day bisphenol A had significant reductions in sperm

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1 motility [**~19 compared to 50% in vehicle controls**]. Sperm motility was also reduced in the mice born
2 to mothers fed a vitamin A-deficient diet but not exposed to bisphenol A. In groups born to mothers fed a
3 vitamin A-deficient diet, there were no differences in sperm motility following exposure to vehicle or
4 bisphenol A. Percentage abnormal sperm was 6.8% in the vehicle control group and was significantly
5 increased in mice exposed to 0.5 µg/day bisphenol A [**~45%**], 50 µg/day bisphenol A (78.2%), 50 µg/day
6 bisphenol A plus retinol acetate (27.8%), vehicle following birth to vitamin A-deficient mothers [**~45%**],
7 or bisphenol A following birth to vitamin A-deficient mother [**~70%**]. No histopathological alterations
8 were reported in testes of mice exposed to 0.5 or 50 µg/day bisphenol A or 50 µg/day bisphenol A plus
9 retinol acetate. The study authors concluded that neonatal exposure to a relatively large dose of bisphenol
10 A damages sperm motility and morphology, effects that are inhibited by vitamin A and enhanced by
11 vitamin A-deficient diets.

12
13 In a second experiment, 3 pups/group were sc injected with 20 µg 17β-estradiol/day, 20 µg 17β-estradiol
14 plus 100 IU acetate retinol acetate/day, 50 µg bisphenol A/day, or vehicle (sesame oil for bisphenol A and
15 17β-estradiol or DMSO for retinol acetate) for 5 days beginning on the day of birth. Mice were killed at
16 18 days of age. Testis, efferent duct, epididymis, and vas deferens were fixed in formalin and analyzed for
17 ERα using an immunohistochemical method. Data were analyzed by ANOVA and Fisher least significant
18 difference test. Bisphenol A exposure had no effect on ERα expression in male reproductive organs.
19 Exposure to 17β-estradiol increased the numbers of ER-positive cells in vas deferens epithelium, but there
20 was no increase when mice were treated with acetate retinol in addition to 17β-estradiol. The study
21 authors concluded that the lack of effect of bisphenol A may be due to its weak estrogenic activity.

22
23 **Strengths/Weaknesses:** This study provided follow-up information to that of Nakahashi et al.(380). The
24 use of 17β-estradiol as a positive control in the testis histology study is a strength; however, PND 18 is
25 prepubertal in mouse and thus this not the optimum time to look for histological changes. Sperm motility
26 studies were done at 10 weeks and a bisphenol A effect was found.

27
28 **Utility (Adequacy) for CERHR Evaluation Process:** This study is slightly useful.

29
30 **Toyama and Yuasa (334)**, supported in part by the Japanese Ministry of Environment and Ministry of
31 Education, Science, Sports, and Culture, examined the effects of neonatal bisphenol A [**purity not**
32 **reported**] exposure on spermatogenesis during puberty and adulthood in rats and mice. [**No information**
33 **was provided about chow or bedding and caging materials. The rat data are reported in Section**
34 **3.2.4.**] ICR mice were sc injected with bisphenol A in a DMSO and olive oil vehicle on PND 1, 3, 5, 7, 9,
35 and 11 (PND 0 = day of birth). Bisphenol A doses were 0.0001, 0.001, 0.005, and 0.010 mg/kg bw in
36 mice. Additional animals were treated with 17β-estradiol and estradiol benzoate. Animals were killed
37 weekly at 2–10 weeks of age and some pups were also killed at 24 and 31 days of age. There were 5
38 animals/dose/time point in bisphenol groups A groups and apparently 3–4 vehicle control mice. Testes
39 were examined by light and electron microscopy. Males from each experimental group (a total of 12
40 mice) were mated with 2 females [**numbers tested in each dose group not reported**]. A total of 12
41 mouse dams were allowed to complete pregnancy. [**It does not appear that any statistical analyses**
42 **were conducted.**]

43
44 In mature spermatids of 7-week-old mice in the vehicle control group, incidences of deformed acrosome,
45 deformed nucleus, and abnormal ectoplasmic specialization were below 0.3%. In 7-week-old mice treated
46 with ≥0.001 mg/kg bw bisphenol A, the incidence of deformed acrosome was >50–60%, the incidence of
47 deformed nucleus was >40%, and the incidence of abnormal ectoplasmic specialization was >60–70%.
48 [**Data were not shown for individual dose levels.**] Similar effects were observed in the groups treated
49 with 17β-estradiol and estradiol benzoate. No effects were reported at other ages. [**Data were not shown**
50 **by study authors.**] The blood-testis barrier remained intact based on histologic observations. All tested
51 males from the bisphenol A group were fertile, and sex ratio, litter sizes, and pup weights were reported

3.0 Developmental Toxicity

1 to be normal. **[No results were shown for individual dose levels. Fertility data were presented in**
2 **Table 4 and 5 of the study, but it is not clear which dose level(s) were represented.]** The study
3 authors concluded that bisphenol A acts as an estrogen and induces transient changes in the male
4 reproductive system of rodents that resolve in adulthood.

5
6 **Strengths/Weaknesses:** This study appears to have been well performed and documented. The strengths
7 include the use of multiple doses of bisphenol A and the use of both rats and mice, allowing interspecies
8 comparisons. Weaknesses include selective data presentation and failure to examine sperm morphology in
9 the fertile 15 week old animals to determine whether the changes in sperm maturation seen at earlier time
10 points had resolved or whether the animals were fertile in the face of such abnormalities.

11
12 **Utility (adequacy) for CERHR Evaluation Process:** This study is suitable for evaluation and shows
13 that high perinatal doses of bisphenol A result in toxicity notable in rats but not in mice given the same
14 dose of agent.

15 16 3.2.9 Sheep

17 **Evans et al. (382)**, supported by the British Council, Irish Health Research Board, and the Royal Society,
18 examined the effects of bisphenol A exposure on gonadotropin secretion on prepubertal female lambs.
19 **[No information was provided about feed or composition of bedding or caging materials.]** Starting at
20 3 weeks of age, female Poll Dorset lambs were weighed weekly, and blood samples were collected 2
21 times/week for measurement of LH and FSH levels. At 4 weeks of age, lambs were randomly assigned to
22 treatment groups according to body weight. From 4 to 11 weeks of age, 6 lambs/group received biweekly
23 im injections with the 10:1 corn oil/alcohol vehicle, 3.5 mg/kg bw bisphenol A **[purity not reported]**,
24 0.175 mg/kg bw diethylstilbestrol **[listed as 0.0175 in the legend for Figure 1 of the study]**, or 3.5
25 mg/kg bw octylphenol. Because of limited information about environmental bisphenol A levels, lambs in
26 the bisphenol A treatment groups were given the same dose as for octylphenol to allow for comparison.
27 Lambs were ovariectomized at nine weeks of age. **[The text of the methods sections reported**
28 **ovariectomy at the beginning of treatment, but that statement appears to be an error since it is not**
29 **indicated elsewhere in the paper.]** On the last day of treatment, blood was collected every 15 minutes
30 for 6 hours to assess pulsatile LH secretion. All lambs were then killed. Adrenal glands, kidneys, and
31 ovaries were weighed. Uteri were examined as discussed in Morrison et al. (383). Data were analyzed by
32 ANOVA, Dunnet multiple comparison post hoc test, regression analysis, Munro algorithm, and paired *t*-
33 tests.

34
35 Compared to the control group, the bisphenol A group did not experience significant changes in body,
36 kidney, adrenal, or ovarian weights. **[No data were shown for body, kidney, and ovarian weights in**
37 **the control versus bisphenol A group.]** Uteri from the bisphenol A group were reported to be visually
38 larger, but no uterine weights were provided. Over the 7-week treatment period, bisphenol A did not
39 significantly affect blood LH or FSH levels compared to controls. Compared to controls, the bisphenol A
40 group experienced significant decreases **[% change compared to controls]** in concentration **[48%]**,
41 amplitude **[77%]**, and frequency **[66%]** of pulsatile LH secretion. Octylphenol did not have any effect on
42 the endpoints examined. Diethylstilbestrol treatment resulted in decreased blood levels of LH and FSH
43 over the treatment period, including the period following ovariectomy. Concentration, amplitude, and
44 frequency of pulsatile LH secretion were also lower in the diethylstilbestrol group, with a greater
45 magnitude of effect compared to bisphenol A. The study authors concluded that the bisphenol A dose
46 tested can inhibit LH secretion in lambs.

47
48 **Strengths/Weaknesses:** The unique animal model and the use of LH pulsatile response are uncommon but
49 interesting. The high dose level is a weakness. The study found no measurable organ effects but
50 inhibition of LH pulses by bisphenol A and 17 β -estradiol.

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1 **Utility (Adequacy) for CERHR Evaluation Process:** This study is useful in the evaluation process.

2
3 **Morrison et al. (383)**, supported by the Wellcome Trust, Dr. Ferranti, and the Irish Health Research
4 Board, examined the effects of bisphenol A exposure on the lamb uterus. **[No information was provided**
5 **on feed or composition of bedding or caging materials.]** At 4 weeks of age, female Poll Dorsett lambs
6 were randomly assigned to treatment groups according to body weight. Beginning at 4 weeks of age and
7 continuing for 7 weeks, 6 lambs/group received biweekly im injections with the 10:1 corn oil:alcohol
8 vehicle, 3.5 mg/kg bw bisphenol A **[purity not reported]**, 0.175 mg/kg bw diethylstilbestrol, or 3.5
9 mg/kg bw octylphenol. Lambs were ovariectomized during the fifth week of exposure. Throughout the
10 study, blood was collected for measurement of gonadotropin levels and the results of those analyses were
11 reported in the study by Evans et al. (382). Lambs were killed following 7 weeks of exposure. Uteri and
12 cervixes were fixed in Bouin solution for histopathological examination, morphometric measurement, and
13 immunohistochemical detection of ER α and ER β . Statistical analyses included ANOVA with Fisher
14 protected least significant difference.

15
16 Significant effects observed with bisphenol A treatment **[% change compared to controls]** were
17 increased uterine/cervical tract weight **[87%]**, endometrial area **[154%]**, and endometrial/myometrial
18 ratio **[65%]**. Qualitative histopathological observations in uteri from bisphenol A-treated lambs included
19 endometrial edema, decreased endometrial gland density compared to controls, and crowding of cells in
20 the uterine epithelium, which contained substantial amounts of eosinophilic, non-vacuolated cytoplasm.
21 In contrast to uteri from control lambs, mononuclear cell exocytosis was not a common observation in
22 uteri from the bisphenol A group. The cervical epithelium was keratinized in the bisphenol A group.
23 Qualitative analyses revealed that diffuse intracellular staining for ER α and ER β in the uterine
24 subepithelium was most pronounced in the bisphenol A and diethylstilbestrol groups. Similar to animals
25 treated with bisphenol A, the diethylstilbestrol group had increased uterine weight, keratinized cervical
26 epithelium, changes in uterine histology, and keratinized cervical epithelium, but there was no change in
27 endometrial/myometrial ratios. No changes were observed following exposure to octylphenol. The study
28 authors concluded that bisphenol A exposure altered the uterocervical environment of lambs.

29
30 **Strengths/Weaknesses:** The follow-up of the study of Evans et al. (382) found effects on the uteri and
31 endometrial area.

32
33 **Utility (Adequacy) for CERHR Evaluation Process:** This study is useful in combination with that of
34 Evans et al. (382)

36 3.2.10 Submammalian species

38 3.2.10.1 Invertebrates

39 **Hill et al. (384)** supported by the Council on Undergraduate Research and the Association for Biological
40 Laboratory Education, examined the effects of bisphenol A on the development of 2 freshwater sponge
41 species. (*Heteromyenia* sp. and *Eunapius fragilis*). Sponge gemmules were incubated in tissue culture
42 wells containing bisphenol A at 0, 0.16, 16, 80, or 160 ppm **[mg/L]**. The control group was incubated in
43 the spring water vehicle. There were 5 replicates/treatment. Nonylphenol and ethylbenzene were also
44 examined. Growth was measured on days 3, 6, and 9. Because growth patterns were similar at all 3
45 evaluation periods, statistical analyses were conducted only for day 6 data. Data were analyzed by
46 ANOVA and Tukey multiple comparison test. In both species, abnormal development or malformation of
47 the water vascular system was observed at a bisphenol A dose of 16 ppm and germination was completely
48 inhibited at 80 and 160 ppm. Significantly reduced growth rates were observed in *Heteromyenia* sp. at
49 160 ppm. Similar effects were observed with nonylphenol and ethylbenzene. The study authors stated that
50 sponges may prove useful for examining endocrine-disrupting compounds.

3.0 Developmental Toxicity

1 **Strengths/Weaknesses:** This study used a unique model with a focus on the aquatic system.

2
3 **Utility (Adequacy) for CERHR Evaluation Process:** This study may have utility for environmental
4 assessment, but its utility for human risk assessment is not clear.

5
6 **Roepke et al. (385)**, supported by the National Oceanic and Atmospheric Administration, examined the
7 effects of bisphenol A exposure on development of two species of sea urchin, *Strongylocentrotus*
8 *purpuratus* and *Lytechinus anamesus*. In dose-response studies, sea urchin embryos were incubated from
9 1 to 96 hours postfertilization in media containing bisphenol A at 0, 250, 500, 750, or 1000 µg/L.
10 Development toxicity was assessed at 96 hours by examining larvae at the pluteus stage. The larvae were
11 categorized as normal, delayed, abnormal, elongated, or hatched. Data were obtained in 3 replicates.
12 Results were reported to be similar for the 2 species, and unless otherwise indicated, data were shown for
13 *S. purpuratus*. In additional studies, sea urchin embryos were incubated in bisphenol A at 0–500 µg/L
14 with and without addition of tamoxifen or bisphenol A at 0–750 µg/L with and without the addition of ICI
15 182,780. Data were analyzed by ANOVA followed by Tukey-Kramer test or Tukey or Student-Newman-
16 Keuls tests for pair-wise multiple comparison. An EC₅₀ of 226.6 µg/L (lower limit: 121.6, upper limit:
17 323.5 µg/L) was estimated for developmental toxicity associated with bisphenol A exposure. Based on
18 EC₅₀ values, 17β-estradiol was ~15 times more potent than bisphenol A. Tamoxifen inhibited
19 developmental toxicity, and ICI 182,780 enhanced the developmental toxicity induced by bisphenol A;
20 similar results were obtained for 17β-estradiol. The study authors concluded that bisphenol A induced
21 developmental toxicity in sea urchins through a tamoxifen-sensitive mechanism at levels exceeding
22 environmentally relevant concentrations.

23
24 **Strengths/Weaknesses:** The use of 2 species and multiple concentrations are strengths.

25
26 **Utility (Adequacy) for CERHR Evaluation Process:** This study is useful in aquatic (salt water)
27 systems. Its utility for human risk assessment is not clear.

28
29 **Andersen et al. (386)**, supported by the Danish Strategic Environmental Research Program, evaluated the
30 effects of bisphenol A on female sexual maturation in the zooplanktonic crustacean *Acartia tonsa*. Eggs
31 were grown in the presence of the algal food source for the organism after exposure of the algae to
32 bisphenol A (>99% purity) for 3 hours to promote sorption by the algae of the test chemical. The treated
33 algae were added to *Acartia tonsa* eggs to give nominal bisphenol A concentrations of 0.2, 2, and 20
34 µg/L. **[Actual concentrations were not reported. An untreated or vehicle-treated control appears to**
35 **have been used.]** 17β-Estradiol 23 µg/L was used as a positive control, and 2,3-dichlorophenol 13.6 µg/L
36 was used as a negative control. On the eighth day of incubation, 10–25 juvenile *Acartia tonsa*/group were
37 transformed to an egg-collection apparatus, in which exposure to treated algae continued. Eggs were
38 collected daily and counted until day 12, at which time a stable adult level of egg production was
39 established. Egg production by group was compared using Student *t*-test. **[A repeated-measures test**
40 **appears not to have been used.]** A significant increase in egg production was shown on day 10 in
41 animals treated with bisphenol A 20 µg/L and 17β-estradiol 23 µg/L compared to control. The authors
42 concluded that bisphenol A accelerated female reproductive maturation in *Acartia tonsa* and that the
43 effect appeared to be estrogenic.

44
45 **Strengths/Weaknesses:** Strengths are the use of multiple exposure levels, the inventive method of
46 feeding bisphenol A to the test organisms, and the use of 17β-estradiol as a positive control.

47
48 **Utility (Adequacy) for CERHR Evaluation Process:** This study may have utility for environmental
49 assessment, but its utility for human risk assessment is not clear.

3.0 Developmental Toxicity

1 **Watts et al. (387)**, supported by the European Union, examined development and reproduction in 2
2 generations of nonbiting midges (*Chironomus riparius*) exposed to bisphenol A. The study began with
3 incubation of 4 egg ropes/group in media containing vehicle, bisphenol A, or ethinyl estradiol
4 **[apparently at the same concentrations described below]**. Twenty 1st-instar larvae from the appropriate
5 media were added to each exposure jar containing dechlorinated water and sediment spiked with
6 bisphenol A at concentrations of 0 (ethanol vehicle control and dechlorinated tap water control), <0.010,
7 0.078, 0.55, 77, 750, or 10,400 µg/L. Four replicate jars were prepared for each dose level.
8 Concentrations in sediment were verified. Numbers and sexes of adults emerging from each replicate jar
9 were determined. Egg ropes produced by the first generation were counted and placed in media
10 containing test solutions or vehicle controls. Four egg ropes/group were selected and used to reseed the
11 sediments with the second generation of larvae. Adults emerging from the second generation were
12 counted. Statistical significance was determined by ANOVA. In the first generation, adult emergence was
13 delayed in females from the <0.010, 0.55, and 77 µg/L bisphenol A groups but was not affected in males.
14 Males were reported to emerge significantly earlier than females. In the second generation, emergence of
15 males and female adults was significantly delayed at ≥0.078 µg/L bisphenol A. At concentrations of
16 0.010–750 µg/L, there were no significant differences in the percentage of adults emerging in either
17 generation. No second-generation adults emerged in the group exposed to 10,400 µg/L. There were no
18 effects on sex ratio. Exposure to bisphenol A did not significantly affect the number of eggs produced by
19 the first generation. In contrast to bisphenol A, exposure to ethinyl estradiol accelerated adult emergence.
20 The study authors concluded that the endpoints evaluated indicated general sediment toxicity but were not
21 useful for detecting estrogenic effects.

22
23 **Strengths/Weaknesses:** The wide range of exposure levels and the use of ethinyl estradiol as a positive
24 control are strengths.

25
26 **Utility (Adequacy) for CERHR Evaluation Process:** This study may have utility for environmental
27 assessment, but its utility for human risk assessment is not clear.

28
29 **Watts et al. (388)**, supported by the European Union, examined the effects of bisphenol A exposure on
30 moulting and mouthpart deformities in nonbiting midge (*Chironomus riparius*) larvae. Four egg-
31 ropes/group were incubated in media containing bisphenol A at 0 (ethanol vehicle or dechlorinated water
32 group), 0.010, 0.1, 1, 10, 100, or 1000 µg/L. Concentrations of bisphenol A were verified in the 1000
33 µg/L group. Upon hatching, exposures were continued in 10 larvae/group. Endpoints examined included
34 survival, time of moulting to successive instars, wet weight 2 days after moulting to fourth instar, and
35 mouthpart morphology in fourth-instar head capsules. Statistical analyses included ANOVA, Tukey-
36 Kramer multiple comparison test, and Kruskal-Wallis test. **[Effects were similar in ethanol and water
37 controls.]** Moulting was delayed and larval weights were significantly decreased in the 1000 µg/L
38 bisphenol A group. Deformities of the mentum were significantly increased in the range of 0.010–1 µg/L
39 bisphenol A. The effects of ethinyl estradiol were also examined, and the study authors noted similar
40 patterns of malformations, with greater incidence following exposure to ethinyl estradiol than bisphenol
41 A. The study authors concluded that exposure to bisphenol A delayed moulting and increased mouth part
42 deformities at concentrations that were at opposite ends of the exposure range.

43
44 **Strengths/Weaknesses:** This study is similar in its strengths to that of Watts et al. (387).

45
46 **Utility (Adequacy) for CERHR Evaluation Process:** This study may have utility for environmental
47 assessment, but its utility for human risk assessment is not clear.

48 3.2.10.2 Frog

49
50 **Iwamuro et al. (389)**, support not indicated, conducted a series of studies to examine the effects of
51 bisphenol A exposure on development of the frog *Xenopus laevis*. In a study to assess survival and

3.0 Developmental Toxicity

1 morphological abnormalities, 60–100 stage 7 embryos/group were exposed to bisphenol A at 0 (ethanol
2 vehicle), 10, 20, 25, 30, 50, or 100 μM [0, 2.3, 4.6, 5.7, 6.8, 11, or 23 mg/L]. Siblings were randomly
3 distributed among different treatment groups. Survival was assessed at 48, 96, and 120 hours. At least 3
4 embryos/group were examined for malformations at 5–7 days following fertilization. Data were analyzed
5 by chi-squared test. Survival of embryos was significantly reduced following exposure to ≥ 25 μM [5.7
6 mg/L] bisphenol A for 96 or 120 hours. Complete mortality was observed at concentrations ≥ 50 μM [11
7 mg/L]. The study authors calculated a median LD_{50} for survival of 21 μM [4.8 mg/L]. The malformation
8 rate was reported for the 10 and 25 μM [2.3 and 4.6 mg/L] group, and significant increases in
9 malformations occurred in the 25 μM [4.6 mg/L] group. The types of malformations were reported as
10 scoliosis, swollen head, and shortened distance between eyes. The effects of 17 β -estradiol were also
11 examined. An increase in malformations was observed with exposure to 10 μM 17 β -estradiol, but there
12 was no effect on survival.

13
14 In a second study, metamorphosis was observed in 10–12 tadpoles (stage 52) placed in solutions
15 containing 10 or 25 μM [2.3 or 5.7 mg/L] bisphenol A with and without the addition of 0.1 μM thyroxine
16 for 21 days. Expression of thyroid hormone receptor- β gene was measured by RT-PCR in 3 regions
17 (head, trunk, and tail) of tadpoles that were exposed to 10 or 100 μM [2.3 or 23 mg/L] bisphenol A with
18 and without the addition of 0.1 μM triiodothyronine or thyroxine. Negative controls were exposed to
19 ethanol/DMSO vehicle. Metamorphosis data were analyzed by Duncan new multiple range test.
20 Bisphenol A significantly inhibited both spontaneous and thyroxine-induced metamorphosis. All
21 concentrations of bisphenol A reduced expression of thyroid hormone receptor- β hormone and inhibited
22 increases in thyroxine- and triiodothyronine-induced expression.

23
24 In a third study, tails were removed from 4 tadpoles/group and cultured for 4 days in media containing 10
25 or 100 μM [2.3 or 23 mg/L] bisphenol A with and without the addition of 0.1 μM triiodothyronine.
26 Negative controls were exposed to ethanol/DMSO vehicles. Data were analyzed by Duncan new multiple
27 range test. Growth of the tails was measured over a 4-day period. Neither bisphenol A dose significantly
28 affected tail growth. Both bisphenol A doses blocked tail shortening that was induced by triiodothyronine.
29 The study authors concluded that high doses of bisphenol A adversely affect development of *Xenopus*
30 *laevis* embryos and larvae.

31
32 **Strengths/Weakness:** The wide range of exposure levels is a strength.

33
34 **Utility (Adequacy) for CERHR Evaluation Process:** This report may be useful in ecotoxicology
35 assessments but is slightly useful for the CERHR evaluation.

36
37 **Oka et al. (390)**, support not indicated, examined the effects of bisphenol A exposure on development of
38 the frog *Xenopus laevis*. Embryos were exposed to the ethanol vehicle or 10–100 μM [2.3–23 mg/L]
39 bisphenol A from developmental stage 6 until the early tadpole stage (late stage 10). Embryos were
40 harvested at stages 19, 23, 33/34, and 40 and prepared for histological examination to determine the
41 presence of apoptotic cells. Apoptosis was also assessed using a TUNEL staining method. Ten embryos
42 were killed at the tail bud stage (stage 35/36, 37/38, and 40), and genomic DNA was isolated and
43 examined by electrophoreses to determine if 180 base pair ladders indicative of apoptosis were present.
44 [No information was provided on the number of individual doses examined or the number of
45 embryos exposed/dose. No quantitative data were presented by authors, and it does not appear that
46 data were statistically analyzed.] Embryos exposed to 40–100 μM [9.1–23 mg/L] bisphenol A died
47 during the gastrula stage. Developmental abnormalities were observed in embryos exposed to 20 μM [4.6
48 mg/L] bisphenol A. The abnormalities included open neural tubes at stage 19, morphological defects at
49 stages 23 and 33/34, and crooked vertebrate, swollen abdomen, and malformed head at stage 40.
50 Malformations persisted following stage 40, and death occurred during the tadpole stage. In stage 33/34
51 and 40 embryos of the 20 μM [4.6 mg/L] group, apoptotic cells were observed in the prosencephalon,

3.0 Developmental Toxicity

1 mesencephalon, rhombencephalon, and spinal cord. Apoptosis was confirmed using the TUNEL staining
2 method. Using the DNA ladder method, it was found that apoptosis also occurred at stages 35/36, 37/38,
3 and 40. The authors briefly stated that they tested stage 10, 19, or 23 embryos and found normal
4 development following bisphenol A exposure. **[No additional details were provided.]** The effects of
5 17 β -estradiol were also examined. Malformations were observed in embryos exposed to 10 μ M 17 β -
6 estradiol, but apoptotic cells were not observed in the nervous system. A very brief description was
7 provided of a study in which embryos were simultaneously exposed to 20 μ M **[4.6 mg/L]** bisphenol A
8 and 1–10 μ M 17 β -estradiol. Co-exposure with 17 β -estradiol did not inhibit bisphenol A-induced
9 apoptosis. The study authors concluded that bisphenol A induced malformations and apoptosis in
10 *Xenopus laevis* at concentrations exceeding environmental levels and that the effects did not appear to
11 occur through an estrogenic mechanism.

12
13 **Strengths/Weaknesses:** The use of 17 β -estradiol exposure to suggest a non-estrogenic mechanism of
14 bisphenol A toxicity is a strength. The omission of some important details and the high concentrations are
15 weaknesses.

16
17 **Utility (Adequacy) for CERHR Evaluation Method:** This study is slightly useful.

18
19 **Sone et al. (391)**, supported by the Japanese Ministry of Environment and Ministry of Education, Culture,
20 Sports, Science, and Technology, examined the effects of bisphenol A exposure on the development of
21 *Xenopus laevis* embryos. Three different sets of experiments were conducted. Data were analyzed by
22 ANOVA followed by Fisher protected least significant difference test. From 3 to 96 hours following
23 fertilization, embryos were exposed to bisphenol A at 1, 2.5, 5, 10, 15, 20, 25, or 30 μ M (0.3, 0.6, 1.1,
24 2.3, 3.4, 4.6, 5.7, or 6.8 mg/L). Each exposure was replicated 3 times. Negative control groups consisted
25 of the ethanol vehicle, medium alone, or dilution medium. Rates of normal embryo development were
26 equivalent in the 3 different negative control groups. In groups exposed to ≥ 20 μ M bisphenol A, there
27 was a significant decrease in normal embryos and a non-significant increase in mortality rate.
28 Teratogenicity was characterized by short body length, microcephaly, flexure, edema, and abnormal gut
29 coiling. Increases in embryo abnormalities were also observed following exposure to ≥ 10 μ M 17 β -
30 estradiol or nonylphenol.

31
32 To determine sensitive stages, embryos were exposed to control media or 20 μ M **[4.6 mg/L]** bisphenol A
33 for 45–48-hour periods ranging from 3 to 48 hours post fertilization, 12–60 hours post-fertilization, 24–72
34 hours post-fertilization, 36–84 hours post-fertilization, or 48–96 hours post-fertilization. Body length,
35 gross malformations, and distance between eyes were measured at 96 hours following exposure. **[The**
36 **methods section indicated that 59–71 embryos were examined in the bisphenol A group for each**
37 **time period of exposure. However, a figure in the study reported the sample size as 3/time period.]**
38 During the period of 3–48 hours following fertilization, statistically significant effects in the bisphenol A
39 group included decreased body length and increased incidences of microcephaly, flexure, edema, and
40 abnormal gut coiling. No increases in abnormal effects were observed following exposure at later time
41 periods. Abnormalities were observed following exposure to 17 β -estradiol or nonylphenol at early or late
42 stages.

43
44 In the third part of the study, embryos were exposed to 20 μ M **[4.6 mg/L]** bisphenol A from 3 to 96 hours
45 following fertilization. RNA was isolated from whole embryos and subjected to analysis by cDNA
46 microarray. Results obtained in microarray analyses were confirmed by PCR analysis. The sample size
47 was reported as 2. The microarray analysis revealed 179 up-regulated and 103 down-regulated genes
48 following exposure of embryos to bisphenol A. The study authors identified 27 genes in which expression
49 was changed following exposure to bisphenol A, nonylphenol, or 17 β -estradiol. The identified genes
50 included: *KNP-1a*, *CmaB*, *XIRG*, α -skeletal tropomyosin, apelin, cyclin G1, *Ube213*, *HGF*, toponin C2,
51 ribosomal protein L9, and *Rattus norvegicus* similar to *CG10042-PA*. The other genes were not identified.

3.0 Developmental Toxicity

1 The study authors concluded that these findings might provide clues to deciphering mechanisms of
2 teratogenic effects associated with bisphenol A and the other compounds examined in this study.

3
4 **Strengths/Weaknesses:** The inclusion of 17 β -estradiol as a comparator was a strength and the high
5 bisphenol A concentration is a weakness.

6
7 **Utility (Adequacy) for CERHR Evaluation Process:** This paper is slightly useful.

8
9 **Pickford et al. (392)**, supported by the Bisphenol A Global Industry group, the Society of the Plastics
10 Industry, the Bisphenol A Sector Group of the European Chemical Industry Council, and the Japan
11 Chemical Industry Association, examined the effects of bisphenol A exposure on development of frog
12 gonads. Beginning at stage 43/45 (~2 days post-hatching, 4 days post-fertilization, exposure day 0) and
13 continuing through stage 66, *Xenopus laevis* larvae were exposed to bisphenol A at nominal
14 concentrations of 0 (water control), 1.0, 2.3, 10, 23, 100, or 500 $\mu\text{g/L}$ in a flow-through test system.
15 Actual concentrations were verified as 0.83, 2.1, 9.5, 23.8, 100, and 497 $\mu\text{g/L}$. A positive control group
16 was exposed to 2.7 $\mu\text{g/L}$ 17 β -estradiol. There were 4 replicate test vessels/dose, with each containing 40
17 larvae (i.e., 160 larvae/test condition). Larvae were observed daily for mortality, behavior, and
18 appearance. Growth and development were assessed on all larvae of a replicate tank on exposure days 32
19 and 62 (36 and 68 [66?] days post fertilization). Froglets were killed and observed at completion of
20 metamorphosis (stage 66). Total length was measured, sex was determined, and testes and ovaries were
21 assessed for abnormalities such as asymmetry, complete absence, presence of melanocytes, irregular
22 shape, segmentation or fragmentation, vacuoles, and ambiguous sexual morphology. Data were analyzed
23 by Fisher exact test, ANOVA, Wilcoxon rank sum test, *G* test, and chi-squared test. Following exposure
24 to bisphenol A, there were no significant differences in survival, distribution of developmental stages on
25 day 32 or 62, time to completion of metamorphosis (stage 66), or length of stage 66 froglets. Bisphenol A
26 exposure did not affect sex ratio or abnormalities in testis or ovary [**data were not shown by authors for**
27 **testis and ovary effects**]. In contrast, exposure to 17 β -estradiol resulted in an increase in ratio of females
28 to males and testicular and ovarian abnormalities. The study authors identified a no observed effect
29 concentration of 500 $\mu\text{g/L}$ for bisphenol A.

30
31 **Strengths/Weaknesses:** The use of a wide range of exposure levels is a strength, but the incomplete data
32 presentation with missing organ weight data and the lack of histological evaluations are weaknesses.

33
34 **Utility (Adequacy) for CERHR Evaluation Process:** This paper is slightly useful.

35
36 **Levy et al. (393)**, supported by the Ministry of Environment and Traffic of Baden-Württemberg,
37 evaluated the effect of bisphenol A on gonad development in *Xenopus laevis* tadpoles. Tadpoles ($n =$
38 40/group) were exposed beginning at stages 42/43 to ethanol vehicle or to bisphenol A (>99% purity) or
39 17 β -estradiol, both at concentrations of 10^{-8} or 10^{-7} M [**bisphenol A concentrations 2.3 and 23 $\mu\text{g/L}$.**
40 **Actual concentrations were 90–105% of target concentrations after addition of bisphenol A to the**
41 **media but decreased to low levels by the end of the 48-hour period between media changes.**] After
42 completion of metamorphosis, froglets were killed for examination of gonads. Tadpoles not completing
43 metamorphosis were killed after 120 days of chemical exposure for examination of gonads. In a second
44 experiment, bisphenol A concentrations were 10^{-8} , 10^{-7} , or 10^{-6} M [**2.3, 23, or 228 $\mu\text{g/L}$**] and the 17 β -
45 estradiol positive control used a concentration of 10^{-7} M. In a third experiment, 50 tadpoles/group were
46 treated for 2 weeks with ethanol vehicle, bisphenol A 10^{-7} M [**23 $\mu\text{g/L}$**], or 10^{-7} 17 β -estradiol after which
47 whole-body homogenates were used for extraction of RNA and determination of *ER* by RT-PCR.
48 Statistical analyses were performed with Kruskal-Wallis *H* test followed by Mann-Whitney *U* test. The
49 gonadal sex of control animals was 56% male and 44% female. 17 β -Estradiol treatment increased the
50 female ratio to 81% at 10^{-7} M and 84% at 10^{-8} M. Bisphenol A treatment resulted in a significant increase

3.0 Developmental Toxicity

1 in females (69%) at 10^{-7} M [23 µg/L]. At 10^{-8} M bisphenol A, there were 65% females, which did not
2 reach statistical significance. In the second experiment, a significant increase in females was seen after
3 treatment with 10^{-7} M [23 µg/L] (70%, compared to 48% in controls and 96% with 17β-estradiol
4 treatment). There was no significant effect of bisphenol A at 10^{-8} M [2.3 µg/L] (51% female) or 10^{-6} M
5 [228 µg/L] (53% female). Bisphenol A and 17β-estradiol both resulted in increased *ER* mRNA. The
6 authors concluded that bisphenol A affects the sexual development of *Xenopus laevis*, probably through
7 an estrogenic mechanism.

8
9 **Strengths/Weaknesses:** The measurement of bisphenol A in the media is a strength, but its lack of
10 stability is a weakness.

11
12 **Utility (Adequacy) for CERHR Evaluation Process:** This study is slightly useful

13
14 **Yang et al. (394)**, supported by the Chinese Ministry of Science and Technology, examined the effects of
15 bisphenol A exposure in black-spotted pond frog tadpoles. Thirty tadpoles/tank were exposed in duplicate
16 to bisphenol A ($\geq 95\%$ purity) at concentrations of 0, 0 (+DMSO vehicle), 2, 20, or 200 µg/L [ppb] for up
17 to 60 days. Tadpoles were also exposed to mixtures containing bisphenol A + nonylphenol at 2 + 2, 20 +
18 20, or 200 + 200 µg/L. Additional tadpoles were exposed to mixtures containing the same bisphenol
19 A/nonylphenol mixtures in addition to *p,p'*-DDE 2 + 2 + 0.5, 20 + 20 + 5, or 200 + 200 + 50 µg/L. Five
20 tadpoles/tank were pooled at 15, 30, 45, and 60 days. The tadpoles were homogenized for measurement of
21 testosterone and thyroxine levels by radioimmunoassay. Alkaline-labile phosphate was measured as a
22 biomarker for vitellogenin. Data were analyzed by ANOVA.

23
24 Malformations of tail flexure were observed in 10% of tadpoles exposed to 200 µg/L bisphenol for 45
25 days, and similar rates of malformation (13.3%) were observed in the mixtures containing 200 µg/L
26 bisphenol A. A “decrease” (not statistically significant) in thyroxine levels was observed following 60
27 days of exposure to all bisphenol A doses (≥ 2 µg/L). “Increases” (not statistically significant) in
28 testosterone levels were reported with all bisphenol A doses at 30 days of exposure. *p,p'*-DDE at ≥ 5 µg/L
29 inhibited increases in testosterone level observed with mixtures of bisphenol A and nonylphenol [not
30 statistically analyzed]. “Increases” (not statistically significant) in alkaline-labile phosphate levels were
31 reported following 30 or more days of exposure to all bisphenol A doses. In animals exposed to bisphenol
32 A and nonylphenol in combination compared to either compound alone, alkaline-labile phosphate levels
33 were increased at 15 days of exposure but decreased at 60 days of exposure [not statistically analyzed].
34 *p,p'*-DDE inhibited the increase in alkaline-labile phosphate levels induced by the bisphenol A +
35 nonylphenol mixture on day 15 of exposure [not statistically analyzed].

36
37 **Strengths/Weaknesses:** The lack of attention to statistical analysis is a weakness and makes the authors’
38 conclusions unreliable.

39
40 **Utility (Adequacy) for CERHR Evaluation Process:** This study is not useful in the evaluation.

41 3.2.10.3 Fish

42
43 **Kishida et al. (395)**, supported by the National Science Foundation and USEPA, included bisphenol A in
44 a study to test the utility of changes in CYP450 aromatase mRNA expression as a marker of xenoestrogen
45 effects in the CNS of zebrafish (*Danio rerio*). Fish embryos were incubated in solutions containing
46 bisphenol A at 0 (DMSO vehicle), 0.01, 0.1, or 10 µM [0, 2.3, 23, or 228 µg/L] from 2 to 48 hours post-
47 fertilization. Expression of the CYP450 aromatase gene was determined in 50 embryos/treatment group
48 using an RT-PCR/Southern blot technique. [There was no mention of statistical analyses of data.] The
49 Southern blot analysis revealed a ~3-fold increase in the band intensity of CYP450 aromatase at the high
50 concentration (10 µM) of bisphenol A. The potency of bisphenol A was determined to be lower than
51 those of 17β-estradiol and diethylstilbestrol, which induced ~3–4-fold increases in band intensity at

3.0 Developmental Toxicity

1 concentrations up to 3 orders of magnitude lower than bisphenol A. In additional experiments with
2 exposure to bisphenol at 2–48 hours post-fertilization, embryo mortality was increased by exposure to 10
3 and 20 μM [228 and 457 $\mu\text{g/L}$] bisphenol A and malformations (curved tails) were increased by exposure
4 to 20 μM . The effects were similar to those observed with 17 β -estradiol, but bisphenol A was less potent.
5 [Very few protocol details were provided, and no data were shown by study authors for mortality
6 and malformation endpoints.] The study authors concluded that bisphenol A could act as a
7 developmental neurotoxicant by upregulating CYP450 aromatase expression but that further studies were
8 needed to determine if there are changes in neural estrogen biosynthesis or CNS development.
9

10 **Strengths/Weaknesses:** A weakness of this paper for the current evaluation is the lack of morphometric
11 data. The significance of the observed change in aromatase is not clear.
12

13 **Utility (Adequacy) for CERHR Evaluation Process:** This study is not useful in the evaluation.
14

15 **Segner et al. (145)**, supported by the European Commission, examined estrogenicity responses and in
16 vivo life cycle effects in zebrafish exposed to bisphenol A. Estrogenicity studies are discussed in Section
17 2. One hundred fertilized eggs/vessel were exposed to bisphenol A (98% purity) at 0, 94, 188, 375, 750,
18 or 1500 $\mu\text{g/L}$ under semistatic conditions. Exposures were continued until fish became sexually mature.
19 The numbers of fish/vessel were adjusted to 50 following 42 days of exposure and 30 following 75–78
20 days of exposure. Two replicates were examined. Bisphenol A concentrations were confirmed by GC/MS.
21 Endpoints evaluated included survival, behavior, growth, time to first spawning, egg production, and
22 fertilization success (percent fertilized eggs/vessel/day). Statistical analyses included ANOVA and
23 William test. EC_{50} values were calculated by probit analysis and analyzed by Kruskal-Wallis and Mann-
24 Whitney U tests. 17 β -Estradiol, ethinyl estradiol, and 4-tert-octylphenol were also examined using similar
25 protocols. The authors only discussed results for reproductive success because they stated that it was the
26 most consistent and reproducible effect following exposure of the fish to estrogenic substances. An EC_{50}
27 value of 6140 nM [1.4 mg/L] bisphenol A was obtained for fertilization success, and the study authors
28 stated that the value exceeded concentrations typically found in the environment. Bisphenol A had a
29 relative potency of 0.0000006 compared to 17 β -estradiol and was 45 times less potent than 4-tert-octyl-
30 phenol. The study authors concluded that the in vivo potency of the compounds was overestimated by in
31 vitro estrogenicity assays (described in Section 2).
32

33 **Strengths/Weaknesses:** This study was well-performed.
34

35 **Utility (Adequacy) for CERHR Evaluation Process:** This study is useful in showing a lack of effect on
36 fertilization at environmentally relevant concentrations of bisphenol A.
37

38 **Metcalfe et al. (166)**, supported by the Environmental Science and Technology Alliance Canada, the
39 Natural Sciences and Engineering Research Council of Canada, and Health Canada, exposed medaka
40 (*Oryzias latipes*) from 1 day after hatching until 85–110 days after hatching to bisphenol A at 0, 10, 50,
41 100, or 200 $\mu\text{g/L}$ ($n = 60$ fish/treatment). Over the 48 hours between media change, actual concentrations
42 were a mean 59.6% of nominal concentrations. Fish were killed and embedded in paraffin for section.
43 Gonads were evaluated to determine the sex of the fish and whether testes contained ova, an intersex
44 condition. Length and weight of the animals and sex ratio were not altered by treatment [statistical
45 methods not reported]. There were 2 instances of intersex gonads in males exposed to bisphenol A 10
46 $\mu\text{g/L}$ and no instances at higher concentrations. Histologic changes in testes including a reduction in germ
47 cells were noted at 50 $\mu\text{g/L}$ and higher. At 200 $\mu\text{g/L}$, oogenesis in females was more advanced than in
48 controls.
49

50 **Strengths/Weaknesses:** Strengths of this study are the step-sectioning of gonads and the use of several
51 positive control estrogens, which worked as expected.

3.0 Developmental Toxicity

1
2 **Utility (Adequacy) for CERHR Evaluation Process:** This study is useful in showing a LOEL for
3 bisphenol A of 10 µg/L.
4

5 **Yokota et al. (396)**, supported by the Japanese Environment Agency, exposed medaka (*Oryzias latipes*)
6 to bisphenol A (>99% purity) at 0, 3.2, 16, 80, 400, or 2000 µg/L from fertilization until 60 days after
7 hatching (n = 60/treatment). Actual bisphenol A concentrations were generally within 3% of nominal
8 concentrations prior to hatching. After hatching, the lower 2 concentrations were ~70–80% of nominal
9 and the higher concentrations were ~90% of nominal. Fish were assessed for survival, time to hatching,
10 and growth. Sixty days after hatching, 19 or 20 fish/treatment were killed and sectioned for examination
11 of the gonads using hematoxylin and eosin staining of fixed specimens. Statistical analysis was performed
12 using ANOVA and nonlinear regression. Hatchability was >90% in all treatment groups. Time to hatch
13 and mortality were not affected by treatment, although there was a non-concentration dependent delay in
14 hatching at 13 µg/L. Body length and weight 60 days after hatching were negatively correlated with
15 bisphenol A concentration, and length and weight at 2000 µg/L were significantly lower than control
16 values on pair-wise comparison. Based on external appearance and gonad examination, there were more
17 females than males at 400 µg/L and there were no males at 2000 µg/L. Control sex ratio was 2:1
18 (male:female). There were 6 fish with intersex gonads among the 19 examined in the 2000 µg/L group.
19 The authors concluded that bisphenol A adversely affects the early life stage of medaka with alteration of
20 sexual differentiation.
21

22 **Strengths/Weaknesses:** This study was well performed.
23

24 **Utility (Adequacy) for CERHR Evaluation Process:** This paper is useful in showing a LOEL of 13
25 µg/L, which is consistent with the results of Metcalfe et al. (166).
26

27 **Pastva et al. (397)** support not indicated, examined the effects of bisphenol A exposure on development
28 of medaka (*Oryzias latipes*). In a study examining abnormalities in embryos, 5 eggs were placed in
29 individual vials containing bisphenol A at 0, 20, or 200 µg/L. There were 5 vials/exposure concentration,
30 for a total of 25 embryos/group. The exposure period began 5 hours following fertilization and was
31 continued for 9 days. Embryos were examined for malformations daily by observing them through the
32 clear protective membrane of the egg. The severity of malformations was scored and severity indices
33 were determined. In a second study examining mortality, newly hatched larvae were exposed for 96 hours
34 to a method control solution, ethanol vehicle control solution, or 200 µg/L bisphenol A. Ten larvae were
35 added to each jar, and there were 3 replicates/test solution (i.e., 30 larvae /concentration). Data were
36 analyzed by *t*-test. The malformation severity index was significantly increased at 5–8 days following
37 fertilization in embryos exposed to 200 µg/L bisphenol A, but the severity index did differ significantly
38 from the control value on day 9. Abnormalities consisted of pericardial edema, hemorrhage, and
39 hemostasis. Larval mortality was not affected by exposure to 200 µg/L bisphenol A. The study authors
40 concluded that exposure to environmentally relevant concentrations of bisphenol A resulted in embryonic
41 deformities in medaka, but that the embryos were able to repair the abnormalities prior to hatching.
42

43 **Strengths/Weaknesses:** This study using medaka is similar in design to the FETAX assay, which uses
44 *Xenopus*. These types of assays have not been demonstrated to have relevance for human risk assessment.
45

46 **Utility (Adequacy) for CERHR Evaluation Process:** This paper is not useful in the evaluation.
47

48 **Lee et al. (398)**, supported by Jeonnam Regional Environment Technology Development Center, exposed
49 51-day-old Korean rockfish (*Sebastes schlegeli*) fry to bisphenol A in feed at 0, 0.05, 0.5, 5, 50, and 100
50 mg/kg diet for 29 days [**purity of bisphenol A and stability in feed not indicated**]. At the end of the
51 experiment, gonads were removed and sex determined by light microscopy of stained sections. There was

3.0 Developmental Toxicity

1 no effect of bisphenol A on sex ratio compared to controls. [The data presentation and statistical
2 analysis are unclear: the number of female fish and number of male fish in each dose group are
3 presented as averages with an unspecified error and analyzed by Student *t*-test. Whole numbers
4 would have been expected with chi-squared analysis.] The authors concluded that there was no
5 estrogenic effect of bisphenol A on sex differentiation in the Korean rockfish.

6
7 **Strengths/Weaknesses:** The use of a positive control, which worked as expected, is a strength of this
8 study. The inadequate presentation of data and statistical analysis is a weakness.

9
10 **Utility (Adequacy) for CERHR Evaluation Process:** This paper is not useful in the evaluation.

11
12 **Honkanen et al. (399)**, supported by the Finnish Graduate School of Environmental Science and
13 Technology and the Academy of Finland, examined the effects of bisphenol A exposure on yolk-sac fry
14 of landlocked salmon. Ten 8-day-old fry/beaker were exposed to bisphenol A at concentrations of 0, 10,
15 100, or 1000 µg/L for 42 days. The ethanol vehicle and pure tap water were used as negative controls.
16 There were 3–4 replicates/dose. One fry/beaker was photographed and killed following 6 days of
17 exposure. After 6 weeks of exposure, all remaining fry were blotted and weighed. Three fry/beaker were
18 photographed and 3 fry/beaker were examined histologically. Statistical analyses included ANOVA and
19 Tukey test. Effects observed in fry exposed to the highest bisphenol A concentration included: yolk sac
20 edema and hemorrhaging around gill arches and the front part of the yolk sac at 6 days of exposure;
21 phlegmatic behavior (lack of activity during siphoning to renew solutions) on the 8th day of exposure; and
22 darkening of color at 17 days of exposure. No increases in mortality were observed. At the end of the
23 exposure period, wet weights were increased in fry exposed to the highest concentration, and the study
24 authors stated that the effect was due to fluid accumulation. In fry exposed to the mid and high
25 concentration of bisphenol A, strongly stained fragments were observed in nuclei and storage substances
26 in liver were decreased. No abnormalities were observed in histological examinations of heart, kidney,
27 and thyroid gland. The study authors concluded that bisphenol A induced toxicity in fry at concentrations
28 rarely found in the environment.

29
30 **Strengths/Weaknesses:** The range of concentrations used in this study is a strength.

31
32 **Utility (Adequacy) for CERHR Evaluation:** The finding of an effect only at a high concentration of
33 bisphenol A may have importance for environmental assessments but is not of utility in the current
34 evaluation process.

35 36 3.2.10.4 Reptile and bird

37 **Stoker et al. (400)**, supported by the Argentine National Agency for the Promotion of Science and
38 Technology and Argentina Ministry of Health, examined the effects of in ovo bisphenol A exposure on
39 sexual development of the crocodylian reptile *Caiman latirostris*. A preliminary experiment was
40 conducted to determine the effects of temperature on sex determination, and it was established that
41 incubation at 30°C resulted in production of females while incubation at 33°C resulted in the production
42 of males. In the main experiment, eggs were collected from 5 nests in Argentina. Half the eggs were
43 incubated at 30°C and the other half at 33°C. Care was taken to avoid exposing eggs to putative sources of
44 estrogens such as spray paint, plastic, and nesting materials. At each incubation temperature, eggs from
45 each nest were equally distributed among treatment groups. Twenty days following collection, 1
46 egg/nest/incubation temperature was opened for stage determination. At developmental stage 20,
47 bisphenol A was applied topically to the eggshell at concentrations of 1.4 or 140 ppm (0.09 or 9 mg/egg).
48 Other eggs were treated with 0.014 or 1.4 ppm 17β-estradiol. Control eggs were left untreated or exposed
49 to the ethanol vehicle. Hatchlings were weighed and measured at birth. At 10 days of age, 4
50 animals/group/incubation temperature were killed for determination of sex by examination of internal
51 genitalia. Sex determination was confirmed by histological evaluation of organs, which were fixed in 10%

3.0 Developmental Toxicity

1 buffered formalin. Morphometric analysis of seminiferous tubules was also conducted in 10-day-old
2 animals. The remaining animals (6–11/group/incubation temperature) were raised until 6 months of age,
3 at which time they were killed, measured, and sexed by examination of external genitalia. Evaluators
4 were blinded to treatment conditions. Statistical analyses included Kruskal-Wallis ANOVA and Mann-
5 Whitney *U* test.

6
7 At 33°C, there was 100% sex reversal in the high-dose bisphenol A and high-dose 17β-estradiol groups at
8 10 days and 6 months of age. Whereas 100% of control and low-dose animals in the 33°C group were
9 male, 100% of animals in the high-dose bisphenol A and 17β-estradiol group were female. Although there
10 was no sex reversal in the low-dose bisphenol A or 17β-estradiol groups incubated at 33°C, morphometric
11 evaluations at 10 days of age revealed significantly increased perimeter of seminiferous tubules, which
12 had empty lumens. There were no significant effects reported for bisphenol A following incubation at
13 30°C. The study authors concluded that bisphenol A induced estrogenic effects in caiman as evidenced by
14 reversed gonadal sex and disrupted gonadal histoarchitecture.

15
16 **Strengths/Weaknesses:** This study appears to have been well performed and the use of a positive control
17 is a strength. A weakness is the expression of exposure level in terms of total egg weight, which precludes
18 easy comparison to human exposure levels.

19
20 **Utility (Adequacy) for CERHR Evaluation Process:** This study has limited utility in the evaluation.

21
22 **Berg et al. (401)**, supported by the Foundation for Strategic Environmental Research and the Swedish
23 Council for Forestry and Agricultural Research, examined the effects of bisphenol A exposure on
24 development of sex organs in quail and chicken embryos. The effects of tetrabromobisphenol A were also
25 examined but will not be discussed. Bisphenol A (99.4% purity) was injected into yolk of Japanese quail
26 eggs on the third day of incubation and into chicken (domestic fowl) eggs on the fourth day of incubation
27 at doses of 0 (propylene glycol vehicle), 67, and 200 µg/g egg. Eggs were also injected with
28 diethylstilbestrol at doses of 2, 20, and 200 ng/g egg. Two days before the anticipated hatching date,
29 embryos were examined for mortality (32–43 quail embryos and 34–91 chicken embryos/group
30 examined) and müllerian duct abnormality or testicular histopathology (8–15 quail embryos/group and 7–
31 30 chicken embryos/group examined). Testes were fixed in 4% formalin. Data were analyzed by Fisher
32 exact probability test.

33
34 Exposure to bisphenol A did not increase mortality in quail embryos. Incidence of females with abnormal
35 müllerian ducts was increased in quail embryos exposed to the high bisphenol A dose but the incidence of
36 ovotestis in males was not increased by bisphenol A exposure. Mortality of chicken embryos was
37 increased following exposure to both bisphenol A dose levels. The incidence of male chicken embryos
38 with ovotestis was increased at the high dose of bisphenol A but there was no effect on females with
39 abnormal müllerian ducts. Effects observed in one or more diethylstilbestrol groups included increased
40 incidence of females with abnormal müllerian ducts in quail embryos and males with ovotestis in quail
41 and chicken embryos. Based on study findings, the study authors concluded that bisphenol A can cause
42 estrogen-like malformations in reproductive organs of birds.

43
44 **Strengths/Weaknesses:** The detailed evaluation of genital tract morphology is a strength, but the
45 expression of exposure level in µg per g egg makes it difficult to compare to human exposure levels.

46
47 **Utility (Adequacy) for CERHR Evaluation Process:** This paper is of limited utility.

48
49 **Halldin et al. (402)**, supported by the European Union and numerous Swedish agencies, examined the
50 effect of in ovo exposure to bisphenol A on sexual behavior of male Japanese quail. On day 3 of
51 incubation, Bisphenol A was injected into the yolk of quail eggs at doses of 67 or 200 µg/g egg. After

3.0 Developmental Toxicity

1 hatching, male and female chicks were housed together. Males were individually housed at 7 weeks of
2 age and examined for sexual behavior at 9 weeks of age. Behavior with a sexually receptive female was
3 evaluated by observing actions such as neck grab, mount attempt, mounts, and cloacal contact movement.
4 Testing was conducted for 2 minutes/day over 5 consecutive days. At the completion of testing, testis
5 weight was measured, gonado-somatic index was determined, and plasma testosterone levels were
6 measured. **[No information was provided for the number of eggs injected, use of a negative control,
7 the number of birds tested, methods of testosterone measurement, or statistical analysis conducted.]**
8 No effects of bisphenol A exposure were reported for any of the effects examined including sexual
9 behavior, testicular weight, gonado-somatic index, or plasma testosterone levels. **[No data were shown.]**
10 Sexual behavior was reportedly affected at an ethinyl estradiol dose of 0.006 µg/g egg and
11 diethylstilbestrol doses of 0.019 and 0.057 µg/g egg. The study authors concluded that bisphenol A was
12 not shown to affect any of the endpoints examined in Japanese quail, which were demonstrated to be a
13 well suited model for studying effects of estrogenic compounds.

14
15 **Strengths/Weaknesses:** The use of 2 positive controls and the attention to sexual behavior are strengths.
16 Weaknesses are the expression of exposure level in µg per g egg, making it difficult to compare to human
17 exposure levels, the lack of detail in the reporting of methods and results, and the lack of apparent
18 statistical analysis.

19
20 **Utility (Adequacy) for CERHR Evaluation Process:** This paper is of limited utility.

21
22 **Panzica et al. (403)**, supported by the University of Torino and Region Piemonte, conducted a study that
23 intended to examine the effects of in ovo bisphenol A exposure on the vasotocin system and sexual
24 behavior of Japanese quail. In 2 sets of experiments, quail eggs were injected with bisphenol A at 50, 100,
25 or 200 µg/egg following 3 days of incubation. Exposure to bisphenol A resulted in a dramatic decrease in
26 the number of live chicks hatching (8–11% versus 55–60% in controls). Chicks that hatched survived less
27 than a week. Dissection of non-hatched embryos indicated that development was blocked immediately
28 following injection in most embryos. A high rate of malformations was observed in chicks that died
29 following hatching. **[No further information was presented for methods, and no data were presented
30 for individual doses.]**

31
32 **Strengths/Weaknesses:** Weaknesses are the expression of exposure level in µg per g egg, making it
33 difficult to compare to human exposure levels, and the lack of data presentation.

34
35 **Utility (Adequacy) for CERHR Evaluation Process:** This paper is not useful in the evaluation.

36
37 **Furuya et al. (404)**, supported by the Japanese Ministry of Education, Science, Sports, and Culture,
38 examined the effects of bisphenol A exposure on growth of testes and combs of male chickens. Beginning
39 at 2 weeks of age, male white Leghorn chicks were orally dosed weekly with corn oil vehicle (n = 5) or
40 200 mg bisphenol A (n = 12). **[The specific method of oral dosing was not reported. It is assumed
41 that birds were dosed until they were killed.]** Chickens were killed at 16 weeks of age. Combs and
42 testes were weighed. Testes were fixed in 4% paraformaldehyde and examined histologically. **[Statistical
43 methods were not discussed, and the levels of statistical significance were not reported.]** Bisphenol
44 treatment did not affect body weight, but comb and testis weight were significantly lower in the chickens
45 exposed to bisphenol A. Spermatogenesis was disturbed in the chickens of the bisphenol A group, as
46 observed by small seminiferous lumen and scarcity of spermatids and mature sperm. Diameter of
47 seminiferous tubules and incidence of seminiferous tubules with mature sperm were significantly lower in
48 the bisphenol A group. The study authors concluded that bisphenol A might disturb the growth of comb
49 and testes in male chickens, possibly through an endocrine mechanism.

3.0 Developmental Toxicity

1 **Strengths/Weaknesses:** The study of male puberty in chickens is a strength. Weaknesses are the use of a
2 single dose level and the lack of information on dosing and statistical analysis. The paper would have
3 been strengthened by measurement of hormone levels.
4

5 **Utility (Adequacy) for CERHR Evaluation Process:** This paper is of limited utility.
6

7 **Furuya et al. (405)**, supported by the Japanese Ministry of Education, Science, Sports, and Culture,
8 examined the effects of bisphenol A exposure on development of male chicks. Beginning at 2 weeks of
9 age, male white Leghorn chicks were orally dosed every 2 days with bisphenol A at 0 (alcohol/corn oil
10 vehicle) 0.002, 0.020, 0.200, 2, or 200 mg/kg bw. The high-dose level was considered to be a positive
11 control based on previous observations in the laboratory. **[No information was provided about the
12 specific method of oral dosing, number of birds treated, purity of bisphenol A, or the type of feed or
13 caging and bedding materials used. It was implied but not clearly stated that exposures were
14 continued until the birds were killed.]** The birds were killed at 5, 10, 15, 20, and 25 weeks of age. The
15 comb, wattle, and testes were weighed. Part of the testicular tissue was used to isolate mRNA for
16 evaluation of *ERα* and aromatase expression by RT-PCR. Additional testicular tissue was fixed in 10%
17 buffered formalin for histopathology analysis and assessment of spermatogenesis by using
18 immunohistochemistry techniques to measure proliferating cell nuclear antigen levels. **[Methods for
19 statistical analyses were not reported.]**
20

21 Although responses were not dose-related, significant decreases in weight (doses at which effects were
22 observed) were reported for comb and wattle at 10 weeks of age (≥ 0.002 mg/kg bw), testis at 10 weeks of
23 age (200 mg/kg bw), comb and testis at 15 weeks of age (≥ 0.020 mg/kg bw), wattle at 15 weeks of age (\geq
24 0.2 mg/kg bw), comb at 20 weeks of age (≥ 0.200 mg/kg bw), testis at 20 weeks of age (200 mg/kg bw),
25 and comb and testis at 25 weeks of age (200 mg/kg bw). There were no effects on body weight.
26 Histopathological observations in testis (doses at which effects were observed) included significant and
27 dose-related reductions in the number of spermatogonia at 5 weeks of age (≥ 2 mg/kg bw) and number of
28 spermatogonia, spermatocytes, and spermatids at 10–25 weeks of age (≥ 0.02 mg/kg bw, except for
29 decreases in spermatocytes at 10 weeks of age, which occurred at ≥ 0.200 mg/kg bw). Seminiferous tubule
30 diameter was significantly reduced at all ages in groups exposed to ≥ 0.020 mg/kg bw. Significant and
31 dose-related reductions in testicular proliferating cell nuclear antigen levels were observed at ≥ 0.200
32 mg/kg bw at 10 weeks of age and ≥ 0.020 mg/kg bw at 15–25 weeks of age. *ERα* mRNA was significantly
33 increased according to dose (doses at which effects were observed) at 10 weeks of age (≥ 0.020 mg/kg
34 bw), 15 and 20 weeks of age (≥ 0.200 mg/kg bw/day), and 25 weeks of age (200 mg/kg bw). Significant
35 and dose-related increases were also observed for aromatase mRNA expression (doses at which effects
36 were observed) at 5 weeks of age (≥ 0.002 mg/kg bw), 10 weeks of age (0.200 mg/kg bw), and 15 weeks
37 of age (200 mg/kg bw). The study authors concluded that exposure to bisphenol A at environmentally
38 relevant levels may affect male chicken phenotypes and result in unbalanced gene expression in the testis.
39

40 **Strengths/Weaknesses:** This paper is a more detailed follow-up of the previous paper by these authors
41 (404), and replication of these results is a strength. Additional strengths are the use of multiple exposure
42 levels and the oral route of administration. The lack of information on statistical methods is a weakness.
43

44 **Utility (Adequacy) for CERHR Evaluation Process:** This paper is useful in demonstrating an effect on
45 spermatogenesis in chickens at the 20 μ g/kg bw dose level.
46

47 3.2.11 *In vitro*

48 **Takai et al. (406)**, supported by the Japanese Ministry of Education, Science, and Culture, the Ministry
49 of Health and Welfare, and the Science and Technology Agency, examined the effects of *in vitro*
50 bisphenol A exposure on preimplantation mouse embryos. Two-cell embryos were obtained from B6C3F₁
51 mice and incubated for 48 hours in media containing bisphenol A at concentrations ranging from 100 pM

3.0 Developmental Toxicity

1 to 100 μM [**23 ng/L to 23 mg/L**]. A negative control group was exposed to the ethanol vehicle and the
2 effects of tamoxifen were also tested. Cell numbers were counted, and trophoblast spreading was
3 evaluated in blastocysts. Statistical analyses included chi-squared, Fisher post hoc, and Student *t*-tests.
4 The number of embryos or samples/group ranged from 14 to 400 for each endpoint evaluated. Significant
5 effects observed with bisphenol A exposure (percent change vs. control) included increased rate of
6 development from 2- to 8-cell embryos following 24 hours exposure to 3 nM [**0.68 $\mu\text{g/L}$**] (94% vs. 88%),
7 increased development to the blastocyst stage following 48 hours exposure to 1 and 3 nM [**0.23 and 0.68**
8 **$\mu\text{g/L}$**] (69% in both dose groups vs. 58.7%), and decreased development to the blastocyst stage following
9 48 hours exposure to 100 μM [**23 mg/L**] bisphenol A (31.2 vs. 58.7%). No effects were observed at
10 concentrations between 10 nM and 10 μM [**23 $\mu\text{g/L}$ and 2.3 mg/L**] bisphenol A. **[Data were not shown**
11 **by study authors.]** Addition of 100 nM tamoxifen to cultures decreased development to the blastocyst
12 stage at 1 and 3 nM [**0.23 and 0.68 $\mu\text{g/L}$**] bisphenol A and increased development to blastocyst stage at
13 100 μM [**23 mg/L**] bisphenol A. Trophoblast spreading was increased in blastocysts exposed to 100 μM
14 [**23 mg/L**] bisphenol A. Bisphenol A exposure did not affect morphology of or cell numbers in
15 blastocysts. The study authors concluded that environmentally relevant concentrations of bisphenol A
16 may affect early embryonic development through the ER and may also affect subsequent development.
17

18 **Strengths/Weaknesses:** The wide range of bisphenol A concentrations is a strength. The postulated
19 involvement of the ER in bisphenol A activity could have been more convincingly demonstrated with a
20 positive control such as 17 β -estradiol and with a more specific estrogen antagonist than tamoxifen. The
21 use of serum-free and phenol red-free media is an appropriate way to avoid estrogenic contamination but
22 is an artificial environment compared to the estrogen-rich milieu in which preimplantation embryos
23 normally develop.
24

25 **Utility (Adequacy) for CERHR Evaluation Process:** This study provides some mechanistic information
26 but is not useful in the evaluation process.
27

28 **Takai et al. (407)**, supported by the Japanese Ministry of Education, Science, Sports, and Culture, the
29 Ministry of Health and Welfare, and the National Institute for Environmental Studies, examined the
30 effects of in vitro preimplantation exposure of mice to bisphenol A. Two-cell embryos were obtained
31 from B6C3F₁ mice and incubated for 48 hours in media containing bisphenol A at 0 (ethanol vehicle), 1
32 nM [**0.23 $\mu\text{g/L}$**] or 100 μM [**23 mg/L**]. Embryos were assessed for number developing to the blastocyst
33 stage, and then blastocysts were transferred to uterine horns of pseudopregnant mice (7/mouse). The
34 dams were allowed to deliver and nurse the litters until weaning on PND 21 (day of birth not defined).
35 Pups were randomly culled to maintain litter sizes at no more than 6. Body weight of pups was measured
36 at birth and at weaning. Litters and pups were considered the experimental unit for statistical analyses.
37 Statistical analyses included chi-squared and Fischer protected least significant difference tests. The
38 number of embryos developing to the blastocyst stage was significantly increased by exposure to
39 bisphenol A at 1 nM [**0.23 $\mu\text{g/L}$**] but decreased by exposure to 100 μM [**23 mg/L**] (72.2 and 33.3% at
40 each respective concentration versus 62.1% in controls). Developing embryos appeared morphologically
41 normal and there were no significant differences in the numbers of cells. Birth weight, number of
42 pups/litter, and sex ratio were not affected by treatment. At weaning, pups in both dose groups weighed
43 more than controls (34–39% greater) and the effect was significant on a litter and pup basis. The study
44 authors concluded that bisphenol A may affect early embryonic and postnatal development at low,
45 environmentally relevant concentrations.
46

47 **Strengths/Weaknesses:** This study was cleverly designed as a follow-up to the previous study and
48 appears to show that a low concentration of bisphenol A stimulates early embryo development while a
49 high concentration inhibits early embryo development. The use of serum-free and phenol red-free media
50 is an appropriate way to avoid estrogenic contamination but is an artificial environment compared to the
51 estrogen-rich milieu in which preimplantation embryos normally develop. The trophic effects of

3.0 Developmental Toxicity

1 bisphenol A at low concentration may have been compensating for the estrogen deprivation of the control
2 culture. It would have been interesting to compare physiologic concentrations of 17 β -estradiol to the
3 control culture conditions.

4
5 **Utility (Adequacy) for CERHR Evaluation Process:** This study did not evaluate the effect of
6 exogenous bisphenol A under physiologic conditions. It is not useful in the evaluation.

7
8 **Li et al. (408)**, support not indicated, examined the effect of in vitro bisphenol A exposure on
9 postimplantation mouse and rat embryos. A limited amount of information was available for the study,
10 which was published in Chinese, but included an abstract and data tables presented in English. GD 8.5
11 mouse embryos and GD 9.5 rat embryos were cultured for 48 hours in media containing bisphenol A at 0,
12 40, 60, 80, or 100 mg/L. Exposure of rat embryos to bisphenol A concentrations \geq 60 mg/L resulted in
13 reduced crown-rump length and yolk sac diameter and affected yolk sac circulation and morphologic
14 differentiation of the nervous system, heart, and forelimbs. Additional effects observed in rats at \geq 80
15 mg/L included reductions in head length, number of somites, and flexion and changes in morphologic
16 differentiation of the otic and optic system and tail. Exposure of mouse embryos to \geq 60 mg/L bisphenol A
17 resulted in reductions in flexion, yolk sac diameter, and yolk sac circulation and changes in morphologic
18 differentiation of the olfactory system and branchial arches. In mouse embryos exposed to \geq 80 mg/L
19 bisphenol A, there were reductions in head and crown-rump length and number of somites and changes in
20 morphologic differentiation of the visual system, heart, brain, auditory system, and fore- and hindlimb
21 buds. The study authors concluded that high concentrations of bisphenol A are toxic to rat and mouse
22 embryos in vitro.

23
24 **Strengths/Weaknesses:** The use of excessively high concentrations of bisphenol A is a weakness.

25
26 **Utility (Adequacy) for CERHR Evaluation Process:** This paper is not useful in the evaluation.

27
28 **Monsees et al. (409)**, supported by the Federal Environmental Agency of Germany, examined the effects
29 of bisphenol A exposure on rat Sertoli cell cultures. Sertoli cell cultures were prepared using testes from
30 18–21-day-old Sprague Dawley rats. The cultures were exposed for 24 hours to bisphenol A or ethinyl
31 estradiol at 0 or 10–50 μ M [**2.3–11 mg/L**]. The effects of pesticides and heavy metals were also examined
32 but will not be discussed. Endpoints assessed following the incubation period included viability by
33 measurement of mitochondrial enzyme activity and lactate and inhibin B production. There were 8
34 replicates/experiment, and the experiment was repeated 3 times. Data were analyzed by Student *t*-test or
35 unpaired Mann-Whitney test. Exposure of cells to bisphenol A resulted in increased lactate production
36 (up to 30%) at \sim 25 μ M [**5.7 mg/L**] bisphenol A and increased inhibin B production at \sim 10 μ M [**2.3 mg/L**]
37 and greater. There was no effect on cell viability following exposure to bisphenol A. Effects of ethinyl
38 estradiol included increased mitochondrial dehydrogenase activity and a biphasic effect on inhibin B
39 production, with an increase at \sim 10 μ M and decreases at higher doses. The study authors concluded that
40 secretion of lactate and inhibin B by Sertoli cells appeared to be sensitive markers for exploring possible
41 Sertoli cell toxicants.

42
43 **Strengths/Weaknesses:** The use of high concentrations of bisphenol A is a weakness. It is not clear how
44 the increased lactate and inhibin B production would correlate with reproductive capacity.

45
46 **Utility (Adequacy) for CERHR Evaluation Process:** This study is not useful in the evaluation.

47
48 **Iida et al. (410)**, supported by an unnamed grantor and by Takeda Science Foundation, examined the
49 effects of in vitro bisphenol A exposure on cultured rat Sertoli cells. The cell cultures were prepared using
50 testes of 18-day-old rats and were exposed for up to 48 hours to bisphenol A at concentrations ranging
51 from 50 to 100 μ M [**11–23 mg/L**]. Control cells were incubated in the DMSO-containing media.

3.0 Developmental Toxicity

1 Morphology was examined by phase-contrast microscopy, and viability was assessed using the CellTiter
2 96 system in cells exposed to 0, 50, 100, 150, 200, and 300 μM [0, 11, 23, 34, 46, and 68 mg/L].
3 Immunochemistry analyses were conducted to detect transferrin and caspase-3 and apoptosis was
4 assessed using a TUNEL method in cells exposed to 0, 100, and 200 μM [0, 23, and 46 mg/L] bisphenol
5 A for 48 hours. A fluorescence staining technique was used to examine actin structure in cells incubated
6 with 200 μM [46 mg/L] bisphenol A. Experiments were performed in triplicate and repeated at least 3
7 times. Data were analyzed by ANOVA.

8
9 Bisphenol A concentrations of $\geq 150 \mu\text{M}$ [34 mg/L] increased detachment of Sertoli cells from substrate
10 and reduced viability. In a time-response study, cell viability was reduced following exposure to 200 μM
11 [46 mg/L] bisphenol A for ≥ 12 hours. Transferrin secretion by Sertoli cells was decreased following
12 incubation with bisphenol A [apparently at $\geq 100 \mu\text{M}$ (23 mg/L); statistical significance not indicated].
13 Following incubation with 200 μM [46 mg/L] bisphenol A, observations included solitary cells with a
14 cortical ring of actin filaments and underdeveloped stress fibers, cells with membrane blebs consisting of
15 protruding actin filaments, and round cells with a disorganized actin cytoskeleton and chromatin
16 condensation. The study authors indicated that the observations were consistent with apoptosis.
17 Expression of capsase-3 was observed in the round Sertoli cells. Capsase-3-positive cells were rarely
18 observed in control cells, but were observed at incidences of $< 1\%$ in the 100 μM [23 mg/L] group and
19 $\sim 9\%$ in the 200 μM group. Further examinations revealed that most and possibly all of TUNEL-positive
20 cells were stained with the caspase-3 antibody. The study authors concluded that decreased viability of
21 Sertoli cells was most likely due to apoptosis and not necrosis.

22
23 **Strengths/Weaknesses:** The evaluation of multiple endpoints is a strength; however, the concentrations
24 of bisphenol A were much higher than are likely to be achieved with human exposures.

25
26 **Utility (Adequacy) for CERHR Evaluation Process:** This paper is not useful for the evaluation.

27
28 **Miyatake et al. (374)**, supported by the Japanese Ministry of Health, Labor, and Welfare, and the
29 Ministry of Education, Culture, Sports, Science, and Technology, conducted a series of studies to
30 examine the effect of bisphenol A exposure on cultures of mouse neuron/glia cells and astrocytes. Cell
31 cultures were obtained from midbrains of ICR mice on PND 1. Statistical analyses included ANOVA
32 followed by Student *t*-test.

33
34 In the first 2 studies, astrocyte and neuron/glia cultures were incubated for 24 hours in media containing
35 bisphenol A or 17 β -estradiol at 0 or 10 fM to 1 μM [bisphenol A concentrations of 2.3 pg/L–0.23
36 mg/L] for 24 hours, and intensity of glial fibrillary acidic protein immunoreactivity was measured. In
37 astrocyte cultures activation of cells, as determined by stellate morphology and significantly increased
38 glial fibrillary acidic protein, occurred with exposure to bisphenol A at 100 fM [23 pg/L], 1 pM [0.23
39 ng/L], 10 pM [2.3 ng/L], 10 nM [2.3 $\mu\text{g/L}$], 100 nM [23 $\mu\text{g/L}$], and 1 μM [0.23 m/L], but the effect was
40 not observed in cells exposed to bisphenol A at 10 fM [2.3 pg/L], 100 pM [23 ng/L], or 1 nM [0.23
41 $\mu\text{g/L}$]. In neuron/glia cultures, a significant increase in glia fibrillary acidic protein was observed at
42 bisphenol A concentrations of 100 fM [23 pg/L], 1 pM [0.23 ng/L], 10 pM [2.3 ng/L], 100 nM [23
43 ng/L], and 1 μM [0.23 mg/L], but not at bisphenol A concentrations of 10 fM [2.3 pg/L], 100 pM [23
44 ng/L], 1 nM [0.23 $\mu\text{g/L}$] or 10 nM [2.3 $\mu\text{g/L}$]. Increases in glial fibrillary acidic protein immunoreactivity
45 were not observed in astrocyte or neuron/glia cultures following treatment with 17 β -estradiol. The study
46 authors concluded that exposure of cell cultures to bisphenol A results in biphasic activation of astrocytes.

47
48 In a third study, the role of steroid hormone receptors in bisphenol A-induced astrocyte activation was
49 examined. Astrocyte and neuron/glia cell cultures were pretreated with an ER antagonist (ICI 182,780),
50 an ER agonist/antagonist (tamoxifen), a progesterone receptor antagonist (mifepristone), or an androgen
51 receptor antagonist (flutamide) for 24 hours. The cultures were then incubated with bisphenol A at 0, 1

3.0 Developmental Toxicity

1 pM [0.23 ng/L], or 1 μM [0.23 mg/L], with and without the receptor ligands, for another 24 hours. None
2 of the ligands attenuated astrocyte activation, and the study authors concluded that bisphenol A-induced
3 activation of astrocytes was not mediated by estrogen, progesterone, or androgen receptors.

4
5 In a fourth study, mouse midbrain astrocyte or neuron cultures were incubated for 24 hours in media
6 containing bisphenol A at 0, 1 pM [0.23 ng/L], 1 nM [0.23 μg/L], or 1 μM [0.23 mg/L]. A fluorescent
7 technique was used to measure calcium levels following treatment of cells with 1–100 μM dopamine. In
8 astrocyte and neuron cultures, dopamine-induced increases in intracellular calcium were enhanced
9 following pretreatment with bisphenol A at 1 pM [0.23 ng/L], but not at 1 nM [0.23 μg/L] or 1 μM [0.23
10 mg/L]. In neuron cells, pretreatment with 1 μM [0.23 μg/L] bisphenol A suppressed dopamine-induced
11 increases in intracellular calcium. The study authors concluded that in vitro bisphenol A exposure results
12 in altered dopamine responsiveness in astrocytes and neurons.

13
14 In a fifth study, neuron/glia cultures were incubated in media containing bisphenol A or 17β-estradiol at 1
15 pM, 1 nM, or 1 μM for 24 hours [bisphenol A concentrations of 0.23 ng/L, 0.23 μg/L, and 0.23 mg/L].
16 An immunohistochemistry technique was used to identify apoptotic cells by the presence of caspase-3.
17 Treatment with 1 μM [0.23 μg/L] bisphenol A activated caspase-3 in neurons. No increase in caspase 3
18 was observed following exposure to cells to 17β-estradiol. The study authors concluded that high in vitro
19 exposures to bisphenol A may result in toxicity to neurons.

20
21 **Strengths/Weaknesses:** The use of multiple concentrations of bisphenol A over a wide range. the
22 evaluation of multiple endpoints, and the comparison to known receptor ligands are strengths.

23
24 **Utility (Adequacy) for CERHR Evaluation Process:** This paper is interesting in suggesting a non-
25 hormonal mechanism of bisphenol A activity. Although the paper contains suggestive mechanistic
26 information, it is not useful for the evaluation.

27 28 3.3 Utility of Developmental Toxicity Data

29 30 3.3.1 Human

31 There are no human data on developmental effects of bisphenol A.

32 33 3.3.2 Experimental animals

34 There are 21 studies in which bisphenol A was given at a single dose level to rats and 6 studies in which
35 bisphenol A was given at a single dose level to mice. These studies explored various aspects of bisphenol
36 A developmental effects but are not useful in establishing dose-response relationships. The lowest dose
37 level evaluated in these studies was 0.0024 mg/kg bw/day in rats (306) and 0.002 mg/kg bw/day in mice
38 (353). There are 23 rat and 29 mouse studies in which bisphenol A was given at multiple dose levels.
39 These studies included oral and subcutaneous administration routes; due to pharmacokinetic
40 considerations, studies using the oral route are of greater utility in estimating human risk.

41 42 3.4 Summary of Developmental Toxicity Data

43 44 3.4.1 Human

45 There are no human data on developmental effects of bisphenol A.

46 47 3.4.2 Experimental animal

48 Studies considered by Expert Panel members to be of utility in evaluating developmental toxicity in rats
49 are summarized in Table 97 (multiple dose-level studies) and Table 98 (single dose level studies). Studies
50 considered by Expert Panel members to be of utility in evaluating developmental toxicity in mice are

3.0 Developmental Toxicity

1 summarized in Table 99 (multiple dose-level studies) and Table 100 (single dose level studies). The
2 discussion of developmental toxicity is arranged according to general endpoints evaluated.

3.4.2.1 General developmental toxicity (growth, survival, malformations)

3.4.2.1.1 Oral

7 Prenatal studies with oral dosing of rats consistently demonstrated no malformations at doses up to 1000
8 mg/kg bw/day (273, 276). Reduced fetal survival and body weights at birth or during the postnatal period
9 were reported in studies with oral exposures occurring throughout the entire gestation and/or lactation
10 periods (276, 293, 411). LOAELs for decreased numbers of live fetuses or pups ranged from 475 to 1000
11 mg/kg bw/day (276, 293, 411). LOAELs for decreased pup body weight at birth were observed at 300–
12 1000 mg/kg bw/day (276, 293, 411). The LOAEL for reduced body weight during the postnatal period
13 was 475 mg/kg bw/day (293, 411). In studies that were less rigorously designed or reported, similar
14 findings were reported for pup body weights during the lactation period (305, 315); 1 study reported a
15 lower LOAEL for postnatal body weight effects (4–40 mg/kg bw/day). The data were not sufficiently
16 reported to allow benchmark dose estimates.

17
18 No increase in teratogenicity was observed in mice with oral bisphenol A doses of ≤ 1250 mg/kg bw/day
19 (273). Prenatal developmental toxicity reported for mice included increased resorptions (LOAEL 1250
20 mg/kg bw/day) and decreased fetal body weight (LOAEL 1250 mg/kg bw/day) (273). Decreased body
21 weight during the postnatal period was also reported in offspring of mouse dams exposed to bisphenol A
22 during the entire gestation and lactation period (LOAEL 600 mg/kg bw/day), but the effect was not
23 observed in a second generation exposed according to the same protocol (376). An increase in hepatic
24 cytoplasmic variation at weaning was also observed in offspring of mouse dams exposed during gestation
25 and lactation (LOAEL 50–600 mg/kg bw/day) (376). A single dose level study with gestational exposure
26 in mice reported increased lactational body weight gain and decreased postnatal pup survival at 0.0024
27 mg/kg bw/day (345).

3.4.2.1.2 Parenteral

30 Similar to studies with oral dosing of dams, sc dosing of rat pups during the lactation period resulted in
31 lower body weights than controls (333); the LOAEL was observed at 427 mg/kg bw/day.

33 In studies in which mouse dams were parenterally dosed during gestation, effects on offspring weight
34 occurred at lower doses than in oral exposure studies, but no evidence of a dose-response relationship was
35 observed in many of those studies. In the mouse parenteral exposure studies, decreased fetal or birth
36 weight were observed at doses of 0.00025–5 mg/kg bw/day (364, 365, 367, 368) and decreased body
37 weight at weaning was reported at 0.002 mg/kg bw/day (364). In contrast, another mouse study with
38 gestational exposure reported an increase in offspring body weight gain at a dose of 0.5 mg/kg bw/day
39 (366). Decreased pup viability on PND 4 following gestational exposure of the dam to 0.00025 mg/kg
40 bw/day was reported in one mouse study (365).

3.4.2.2 Reproductive development

3.4.2.2.1 Oral

45 Delays in vaginal opening were observed in offspring of rat dams receiving high oral doses of bisphenol
46 A on GD 6–15 or during the entire gestational and lactational period (278, 293, 411). LOAELs for
47 delayed vaginal opening were reported at 50–475 mg/kg bw/day. When rat dams were dosed with
48 bisphenol A at ≤ 384 mg/kg bw/day beginning on GD 11 or later, no delays in vaginal opening were
49 observed (297, 305). No delays in vaginal opening were observed with doses of bisphenol A ≤ 1.2 mg/kg
50 bw/day administered to dams during gestation or lactation (217, 292, 293, 311, 411).

3.0 Developmental Toxicity

1 One study reported estrous cycle alterations in offspring of rats given 1.2 mg/kg bw/day bisphenol A in
2 drinking water from GD 6 through the lactation period (217). Estrous cycles alterations were not reported
3 in other rat oral exposure studies covering a wide range of dose (<1–475 mg/kg bw/day) administered
4 during all or part of the gestational or lactational periods (292, 293, 297, 305, 311, 411).

5
6 Studies suggest that preputial separation is delayed following oral administration of high bisphenol A
7 doses (LOAELs 47.5–475) to male rat offspring in the post weaning period (293, 307, 411). No effects on
8 preputial separation were observed when treatment of rat dams with high doses (50–384 mg/kg bw/day)
9 ended during the gestation or lactation period (278, 305). Oral doses of bisphenol A \leq 1 mg/kg bw/day
10 also had no effect on preputial separation (292, 293, 411).

11
12 Effects on rat sperm parameters were inconsistent. Decreased sperm count and daily sperm production
13 was reported in offspring of dams exposed during gestation (LOAEL 50 mg/kg bw/day for sperm count/g
14 testis, LOAEL 50 mg/kg bw/day for daily sperm count/g testis) (278). A single dose level study reported
15 decreased numbers of rats undergoing spermatogenesis following postweaning exposure of males to 100
16 mg/kg bw/day (307). In contrast, no consistent effects on sperm parameters were observed in rats
17 following exposures with up to 475 mg/kg bw/day during the prenatal, lactational, and post-weaning
18 periods (293, 411). Other rat studies with gestational and lactational doses ranging from <1 to 4 mg/kg
19 bw/day also reported no effects on sperm parameters (292, 294, 311). Testicular histopathology was only
20 reported in a single dose level study at a bisphenol A dose of 100 mg/kg bw/day administered in the post-
21 weaning period (307).

22
23 Although some sporadic effects were reported for anogenital distance in male and female rats, study
24 authors concluded that the endpoint was not affected by prenatal, lactational, and/or post-weaning
25 exposure to bisphenol A (217, 278, 292, 293, 305, 311, 411).

26
27 In oral dosing studies, no effects on rat prostate weight were observed with bisphenol A doses of <1–475
28 mg/kg bw/day administered during the gestational, lactational, and/or post-weaning periods (278, 293,
29 294, 297, 305, 311, 411).

30
31 There were some indications that bisphenol A exposure may affect serum LH levels in male rats after
32 exposure to \leq 1.2 mg/kg bw/day administered during gestational or postnatal periods, but the biological
33 significance of the effect was uncertain because of questions regarding exposure characterization, lack of
34 dose response relationships, and reproducibility of the effect (217, 306). A study utilizing single and
35 multiple dose levels suggested possible alterations in testosterone levels following bisphenol A exposure
36 of 0.0024 mg/kg bw/day during the prenatal or postnatal period (306).

37
38 Exposure of mice to bisphenol A during pre- and postnatal development delayed preputial separation
39 (LOAEL 600 mg/kg bw/day)(376). Effects reported for anogenital distance were inconsistent. A single
40 dose study reported an increase in anogenital distance in mice at 0.050 mg/kg bw/day (347). A second
41 study with a wide dose range (0.003–600 mg/kg bw/day) reported no consistent or dose-related effects on
42 anogenital distance (376).

43
44 One group of investigators reported decreased sperm production efficiency (LOAEL 0.020 mg/kg
45 bw/day) (341) and increased prostate weight at 0.002 but not 0.020 mg/kg bw/day (205, 341) in offspring
46 of mouse dams exposed during pregnancy. Those prostate effects were consistent with findings in single
47 dose level studies with gestational exposure of mice, however, it is noted that the studies had differing
48 periods of exposure and ages of evaluation. One of these studies demonstrated increased prostate weight
49 at 0.050 mg/kg bw/day (347). Another study demonstrated increased numbers of prostate ducts and
50 proliferating cell nuclear antigen staining in dorsolateral prostate and increased prostate duct volume in
51 dorsolateral and ventral prostate at 0.010 mg/kg bw/day (351). However, no effects on prostate or sperm

3.0 Developmental Toxicity

1 production was observed in more robust studies with multiple dose levels and larger group sizes. Two
2 mouse studies (342, 343) that attempted to replicate earlier findings on prostate weight and sperm
3 production (205, 341) reported no increase in prostate weight or decreases in sperm production, efficiency
4 of sperm production, and/or sperm concentration at doses ≤ 0.2 mg/kg bw/day. A third mouse study with
5 exposures occurring during gestation, lactation, and post-lactational periods also reported no effects on
6 prostate weight, daily sperm production, or efficiency of daily sperm production at doses of 0.003–600
7 mg/kg bw/day (376). A fourth mouse study demonstrated no effect on sperm density following low-dose
8 exposure (≤ 0.200 mg/kg bw/day) during gestation or the post weaning period (369).

9
10 Seminiferous tubule hypoplasia in mouse weanlings was reported following exposure during pre- and
11 postnatal development (LOAEL 50–600 mg/kg bw/day; BMD₁₀ 283–591 mg/kg bw/day) but the effect
12 was not observed in mice examined in adulthood (376). The findings were similar to those in studies
13 reporting no testicular histopathology or lesions in reproductive organs following prenatal or postnatal
14 exposure to bisphenol A at ≤ 0.2 mg/kg bw/day (342, 369). Changes in testicular expression of *ER β* and
15 *ER α* mRNA were reported with post-weaning exposure of mice to 17.5 mg/kg bw/day bisphenol A (370).

16
17 Following exposure of mice during pre- and postnatal development; no effect on age of vaginal opening,
18 estrous cyclicity, or numbers of ovarian primordial follicles were observed at doses ranging from 0.003–
19 600 mg/kg bw/day (376). No effect on age of vaginal opening was reported but there was a shortened
20 period between vaginal opening and first estrus following gestational exposure to 0.0024 mg/kg bw/day
21 in a single dose level study (345).

22 23 3.4.2.2.2 Parenteral exposure

24 In rat sc dosing studies, vaginal opening was delayed with exposure of offspring to bisphenol A on PND
25 0–9 (LOAEL 105 mg/kg bw/day) (333), but no delay was observed in a single dose level study (300
26 mg/kg bw/day) in which rats were exposed for a shorter period (PND 1–5) (325). Decreased numbers of
27 rats with normal estrous cycles were observed with sc dosing of pups during the lactation period (LOAEL
28 427 mg/kg bw/day) (333). Effects reported for female reproductive organs following direct postnatal sc
29 dosing of rats included increased numbers of females with cleft clitoris (LOAEL 105 mg/kg bw/day),
30 increased numbers with polycystic ovaries (LOAEL 105 mg/kg bw/day), and decreased numbers with
31 corpora lutea, numbers of corpora lutea, and corpora lutea area (most sensitive effect level: LOAEL ≤ 105
32 mg/kg bw/day) (333).

33
34 No effects on preputial separation were observed following postnatal sc dosing of male rats with
35 bisphenol A at ≤ 1 or 300 mg/kg bw/day in a single and multiple dose level study (335). Results for the
36 prostate were inconsistent following direct exposure in the postnatal period. In contrast to oral dosing
37 studies, some biochemical and ultrastructural effects (e.g., changes in area of androgen receptor positive
38 cells and prostatic acid phosphate-positive cells and a slight increase in secretory granules and slight
39 decrease in microvilli) were observed with direct postnatal sc dosing of rats with bisphenol A at ≤ 0.025
40 mg/kg bw/day (286, 332). However, concerns regarding the prostate studies were noted by the Expert
41 Panel. In 1 study (286), the use of pure DMSO as a vehicle and lack of dose-response relationships were
42 noted. Insufficient reporting of data and no evaluation past the developmental stage were concerns noted
43 for the second study (332). Other multiple or single dose level rat studies conducted with doses ≤ 1 mg/kg
44 bw/day reported no effects on prostate morphology or weight (335, 336). In a single dose level study (50
45 mg/kg bw/day) there was no histological evidence of prostate inflammation, but increases were observed
46 for lateral prostate weight and focal luminal polymorphonuclear cellular infiltrate (326).

47
48 In studies in which rats were injected with single dose levels during the lactational period, reduced height
49 of efferent duct epithelium on PND 18 and 25 was observed with exposure to 37 mg/kg bw (324) and
50 advanced testicular lumen formation and changes in Sertoli cell volume occurred at 100 mg/kg bw/day
51 (327). Other single dose-level rat studies reported no histopathological alterations at ≤ 20 mg/kg bw (329)

3.0 Developmental Toxicity

1 or effects on Leydig cells at ≤ 100 mg/kg bw (330). No effect on sperm count, motility, or morphology
2 was reported at ≤ 1 mg/kg bw/day administered during the postnatal period (335). A postnatal rat
3 exposure study that provided no information for individual doses reported increases in deformed
4 acrosome, deformed nucleus, and abnormal ectoplasmic specialization in sperm at ≥ 0.010 mg/kg bw/day
5 (334).

6
7 Areolas and nipple were increased in male and female rats sc injected with 400 mg/kg bw/day bisphenol
8 A on GD 11–20 (246, 288). Serum prolactin levels were increased in male and female rats sc injected
9 with ≥ 20 mg/kg bw/day on PND 1–5, but dose response was questionable (331). A single dose level
10 study also demonstrated an increase in serum prolactin during treatment of male rats with 50 mg/kg bw
11 bisphenol A but the effect did not persist to adulthood (326). It appears that plasma testosterone may be
12 increased following treatment with ≥ 20 mg/kg bw/day (330) but not with doses ≤ 20 mg/kg bw/day (329,
13 335).

14
15 Studies with neonatal parenteral exposures in mice reported decreased sperm counts at 25 mg/kg bw/day
16 (380, 381). No reduction in sperm count was reported following gestational exposure to ≤ 5 mg/kg bw/day
17 (368). Increases in deformed acrosome, deformed nucleus, and abnormal ectoplasmic specialization in
18 sperm were reported following lactation exposure of mice to ≥ 0.001 mg/kg bw/day (334). No evidence of
19 testicular histopathology was observed following injection of mouse neonates with ≤ 25 mg/kg bw/day
20 (381). Anogenital distance was increased in 60-day old male mice that had been exposed to ≥ 0.002 mg/kg
21 bw/day, but the effect was not observed at weaning (364).

22
23 Effects of parenteral gestational exposure of mice on vaginal opening were inconsistent. At doses
24 between 0.020 and 2.5 mg/kg bw/day, either no effect or accelerated vaginal opening was reported (361,
25 365). A delay in vaginal opening was reported following gestational exposure of mice to bisphenol A
26 0.00025 mg/kg bw/day (365). Decreased age at first estrus was reported following gestational exposure of
27 mice to bisphenol A 0.020 mg/kg bw/day bisphenol A (364). Anogenital distance was increased in
28 weanling female mice that had been exposed to bisphenol A ≥ 0.002 mg/kg bw/day, but no dose-response
29 relationship was observed and the effect was not present at 60 days of age (364).

30
31 Changes in estrous cyclicity in mice were reported following gestational exposure to ≥ 0.002 mg/kg
32 bw/day (361, 364, 366), but the effects were not always dose-related. Another mouse study reported no
33 effect on estrous cyclicity at ≤ 100 mg/kg bw/day administered during the neonatal period (377). The
34 number of 4-week-old mice with no corpora lutea and with vaginal cornification was increased following
35 gestational exposure to ≥ 0.5 mg/kg bw/day (366). A decrease in the number of ovariectomized mice with
36 corpora lutea was observed following gestational exposure to 10 mg/kg bw/day and increases in
37 polyovular follicles were observed following neonatal exposure to 100 mg/kg bw/day (377). Increased
38 fluid-filled ovarian bursae were reported following gestational exposure to ≥ 0.025 mg/kg bw/day (361).

39
40 Increases in vaginal epithelial layers were reported in ovariectomized mice that had been exposed to
41 bisphenol A ≥ 10 mg/kg bw/day during gestation or 100 mg/kg bw/day during the neonatal period (377).
42 Increased mitotic rate in uterine stromal and vaginal epithelial cells was also increased following neonatal
43 mouse exposure to 100 mg/kg bw/day (377). Decreased uterine lamina propria volume, increased BrdU
44 incorporation by uterine epithelial cells, and increased expression of progesterone receptor (not dose
45 related) and ER α by uterine epithelial cells was observed in mice at a bisphenol A dose of ≤ 0.000250
46 mg/kg bw/day (379).

47
48 In mouse studies examining the effects of parenteral gestational exposure on the mammary gland,
49 changes in the development of mammary structures, BrdU incorporation, and progesterone receptor
50 expression by mammary epithelial cells were observed at a bisphenol A dose of ≥ 0.000025 (360, 361,

3.0 Developmental Toxicity

1 363), but the results were not always dose-related and there was no consistency of response at different
2 evaluation time periods.

3
4 Effects reported in lambs receiving biweekly injections with 3.5 mg/kg bw bisphenol A from 4 to 11
5 weeks of age were decreased concentration, amplitude, and frequency of pulsatile LH secretion; increased
6 uterine/cervical tract weight, endometrial area, and endometrial/myometrial ratio; endometrial edema;
7 decreased endometrial gland density; crowding of cells in the uterine epithelium; substantial amounts of
8 eosinophilic, non-vacuolated cytoplasm in uterine epithelium; and keratinized cervical epithelium (382,
9 383).

10 3.4.2.3 Nervous system endpoints

11 3.4.2.3.1 Oral

12
13 No effects on SDN-POA volume were observed in offspring of rats orally exposed to bisphenol A doses
14 ranging from ≤ 0.3 to 384 mg/kg bw/day during the gestation and lactation period (297, 305, 311). One of
15 the studies demonstrated reduced sexual dimorphic differences in locus ceruleus volume at ≥ 0.030 mg/kg
16 bw/day (311), but some uncertainty regarding the dose was noted. Another low-dose study demonstrated
17 changes in various parameters associated with somatostatin and receptor subtype 3 or GABA in both
18 immature and adult rats that had been exposed to ≥ 0.040 mg/kg bw/day during gestation and
19 lactation (313). Single dose level rat studies demonstrated reduced sexually dimorphic difference in
20 corticotropin-releasing hormone neurons in anterior stria terminalis at 2.5 mg/kg bw/day (284) and
21 changes in expression of ER α and tyrosine hydroxylase by cells of the anteroventral periventricular
22 nucleus of males and females at 100 mg/kg bw/day (338).

23
24
25 Two multiple dose-level studies and a single dose level study with exposure occurring during prenatal
26 and/or postnatal periods demonstrated changes in sexually dimorphic behaviors (e.g., activity, rearing,
27 test performance) of rats at doses ≤ 0.1 mg/kg bw (285, 311, 322). One study demonstrated changes in
28 avoidance test performance and grooming in adult male rats exposed to 4 but not ≥ 40 mg/kg bw/day
29 during part of the gestation and lactation periods (315). An additional single and multiple dose level study
30 also demonstrated changes in learning test performance by males or females following prenatal and/or
31 postnatal exposure to a bisphenol A dose ≤ 0.25 mg/kg bw/day (322). No changes in sexual behavior were
32 reported for female rats exposed to 0.3–320 mg/kg bw/day or males exposed to ≤ 0.3 mg/kg bw/day
33 during the gestation and/or lactation period (297, 311).

34
35 A number of studies were conducted in which rats were orally dosed with bisphenol A at 0.040 mg/kg bw
36 throughout the entire mating and gestation period and/or 0.400 mg/kg bw/day from GD 14 through PND
37 6 (317-321). Numerous effects were observed under both dosing conditions. Based on behaviors in hole-
38 board and elevated maze tests, the study authors concluded that anxiety and motivation to explore were
39 reduced in both treated males and females, but there was no clear masculinization of female behavior
40 (317). In a study examining sexual performance, the study authors concluded that bisphenol A slightly
41 intensified female behavior and slightly reduced male performance in a limited number of parameters
42 while having no effects on other male parameters (318). Based on performance in sexual behavior and
43 intruder testing, the study authors concluded that bisphenol A potentiated female behavior and
44 depotentiated male behaviors. Based on findings of a social interaction study, the study authors concluded
45 that 2 factors of female behavior were masculinized by treatment, play with females and sociosexual
46 exploration (319). In a second study examining social interactions, the study authors noted that although
47 some aspects of female behavior were defeminized, bisphenol A did not clearly induce masculinization of
48 female behavior. Changes in open-field behavior and impulsivity by both males and females were
49 reported in another study (320).

3.0 Developmental Toxicity

1 Behavioral effects have been reported in mice after exposure to ≥ 0.002 mg/kg bw/day bisphenol A during
2 gestation and/or lactation including increased aggression at 8 but not 12 or 16 weeks of age (358) and
3 increased morphine preference (374). A single dose level study reported decreased *d*-amphetamine
4 preference in female mice following gestational exposure to 0.010 mg/kg bw/day (359).

5 6 *3.4.2.3.2 Parenteral*

7 One single dose level rat study demonstrated no effect on SDN-POA volume following neonatal exposure
8 to 300 mg/kg bw/day (325). Mice that had been parenterally exposed to 0.010 mg/kg bw/day bisphenol A
9 during gestation demonstrated changes in maternal behavior during lactation (e.g., time nursing, spent in
10 nest, and grooming) (352).

11 12 *3.4.2.4 Other endpoints*

13 Limited studies suggest no effect on thyroid function following oral exposure of rat dams during the
14 gestation and lactation periods. A non-dose related increase in serum thyroxine levels was only observed
15 on 1 of 4 evaluation periods during postnatal development at ≥ 1 mg/kg bw/day orally (308). No effects on
16 thyroxine or thyroid stimulating hormone-induced increases in thyroxine levels were observed with
17 exposure to ≤ 40 mg/kg bw/day during pre- and postnatal development (302).

18
19 Following oral exposure of mice to bisphenol A during gestation, changes were observed for mRNA
20 expression of arylhydrocarbon receptors, receptor repressor, or nuclear translocator and retinoic acid and
21 retinoid X receptors in brain, testes, and/or ovary at 0.00002–20 mg/kg bw/day, but many responses were
22 not dose-related (353-355). The study authors suggested those changes as possible mechanisms for
23 bisphenol A-induced toxicity.

24
25 Increases in immune response were observed following gestational exposure of mice to bisphenol A at
26 oral doses ≥ 0.3 mg/kg bw/day (356).

27
28 A summary of LH and testosterone effects observed in humans and in bisphenol A-exposed experimental
29 animals is included in Section 4.4.
30

3.0 Developmental Toxicity

- 1 **Questions for the Expert Panel**
- 2 Are human data sufficient for an evaluation of the developmental toxicity of bisphenol A following
- 3 prenatal exposure?
- 4 If so, what are the relevant exposure conditions and endpoints?
- 5 Are human data sufficient for an evaluation of developmental toxicity of bisphenol A following
- 6 exposure of children?
- 7 If so, what are the relevant exposure conditions and endpoints?
- 8 Are experimental animal data sufficient for an evaluation of the developmental toxicity of bisphenol A
- 9 following prenatal or lactational exposure or other exposures of immature animals?
- 10 If so, what are the relevant experimental animal models, exposure conditions, and endpoints?
- 11 If the experimental animal data are sufficient for an evaluation, are the data assumed relevant, relevant,
- 12 or not relevant?
- 13
- 14 Note: The definitions of the term sufficient and the terms assumed relevant, relevant, and not relevant
- 15 are in the CERHR guidelines at <http://cerhr.niehs.nih.gov/news/guidelines.html>.
- 16

3.0 Developmental Toxicity

Table 97. Summary of Developmental Toxicity in Multiple Dose Rat Studies

Strain, route	Endpoint	Dose, mg/kg bw/day						Reference
		NOAEL	LOAEL	BMD ₁₀	BMDL ₁₀	BMD _{1SD}	BMDL _{1SD}	
CD, gavage GD 6–15	Implantation sites, resorptions, body weight, viability, sex ratio, and malformations.	≥640						Morrissey et al. (273)
Sprague Dawley, gavage GD 1–20	↓ Live fetuses/litter	300	1000	929	348	982	713	Kim et al. (276)
	↓ Male body weight	100	300	456	339	694	497	
	↓ Female body weight	300	1000	439	328	682	490	
	↓ Ossification	300	1000					
Wistar-derived Alderley Park, gavage GD 6–21	Delayed vaginal opening	0.1	50	68	51	35	16	Tinwell et al. (278)
	↓ Sperm count/testis	0.1	50	55	30	57	31	
	↓ Sperm count/g testis	0.1	50	81	41	68	34	
	↓ Daily sperm count/testis	0.1	50	56	31	59	31	
	↓ Daily sperm count/g testis	0.1	50	83	42	70	34	
	Anogenital distance, first day of estrus, or age of preputial separation, or prostate weight.	≥50 (high dose)						
	↑ Relative area of prostate vimentin-positive cells		≤0.025 (low dose) ^a					
	↓ Relative area of prostate α-smooth muscle actin positive cells		0.025					
	↓ Area of androgen receptor-positive cells in prostate periductal stroma		0.025 ^a					
↓ Area of prostatic acid phosphate-positive cells in epithelial cells		0.025 ^a						
CD, gavage, 2-generations exposure including pre- and postnatal development periods	Prenatal or postnatal growth or survival, developmental landmarks, anogenital distance, age of puberty, fertility, estrous cyclicity, or sperm counts.	≥0.2 (high dose)						Ema et al. (292)

3.0 Developmental Toxicity

Strain, route	Endpoint	Dose, mg/kg bw/day						Reference
		NOAEL	LOAEL	BMD ₁₀	BMDL ₁₀	BMD _{1SD}	BMDL _{1SD}	
Sprague Dawley, dietary multiple generations with exposure during pre-and post natal development	Live F1 pups/litter	47.5	475	268	192	559	394	(293, 411)
	Live F2 pups/litter	47.5	475	422	152	459	294	
	Live F3 pups/litter	47.5	475	236	174	376	286	
	F1 body weight, PND 4	47.5	475	406	283	561	400	
	F1, F2, or F2 body weight, PND 7	47.5	475	217–328	183–257	265–410	218–313	
	F1, F2, or F2 body weight, PND 14	47.5	475	183–243	163–209	177–227	153–191	
	F1, F2, or F2 body weight, PND 21	47.5	475	208–252	166–226	223–267	175–220	
	↑ Age at F1 vaginal opening	47.5	475	394	343	206	176	
	↑ Age at F2 vaginal opening	47.5	475	404	336	277	228	
	↑ Age at F3 vaginal opening	47.5	475	471	401	396	203	
	↑ Age at F1 preputial separation	4.75	47.5	466	411	188	163	
	↑ Age at F2 preputial separation	47.5	475	300	255	241	203	
	↑ Age at F3 preputial separation	47.5	475	547	473	222	189	
Mating, fertility, pregnancy, or gestational indices; precoital interval, postimplantation loss, estrous cyclicity, and reproductive organ histopathology; sperm count, morphology or motility; anogenital distance in males or females; areolas/nipples in males.	≥475 (high dose)							
Han-Wistar, drinking water from prior to mating through gestation and lactation	Male reproductive organ weights, sperm production, testicular histopathology.	≥0.775–4.022 (high dose)						Cagen et al. (294)
Sprague Dawley, drinking water from GD 6 through	↓ Normal estrous cycles	0.1	1.2					Rubin et al. (217)
	↓ Serum LH level	0.1	1.2	0.94	0.48	1.6	0.78	

3.0 Developmental Toxicity

Strain, route	Endpoint	Dose, mg/kg bw/day						Reference
		NOAEL	LOAEL	BMD ₁₀	BMDL ₁₀	BMD _{1SD}	BMDL _{1SD}	
lactation period	Day of vaginal opening, or anogenital distance	≥1.2 (high dose)						
Sprague Dawley rat, dietary from GD15 to PND10	Male pup weight, PND 2	49–80 ^b	232–384 ^b	232–378 ^b	106–173 ^b	226–369 ^b	80–130 ^b	Takagi et al. (305)
	Female pup weight, PND 2	49–80 ^b	232–384 ^b	234–381 ^b	127–207 ^b	229–374 ^b	104–169 ^b	
	Male pup weight, PND 2–10	49–80 ^b	232–384 ^b	130–213 ^b	73–119 ^b	156–254 ^b	87–141 ^b	
	Female pup weight, PND 2–10	49–80 ^b	232–384 ^b	148–242 ^b	88–144 ^b	228–373 ^b	80–131 ^b	
	Anogenital distance, vaginal opening, preputial separation, or estrous cyclicity, prostate weight, SDN-POA volume.	≥232–384 ^b (high dose)						
Long Evans, gavage PND 21 to 35	↓ Serum 17β-estradiol		0.0024 (low dose) ^{a,c}					Akingbemi et al. (306)
	↓ Serum LH and testosterone		0.0024 (low dose) ^{a,c}					
CD, gavage GD 6 through PND 20	Thyroxine or thyroid stimulating hormone-induced increases in thyroxine levels.	≥40 (high dose)						Kobayashi et al. (302)
Sprague Dawley, gavage GD 11 through PND 20	Volume of SDN-POA, age or weight at vaginal opening or first estrous, estrous cyclicity, mean lordosis intensity, prostate weight, or histopathology in ventral prostate, ovary, or uterus.	≥320 (high dose)						Kwon et al. (297)
Wistar, [apparently through drinking water during the entire gestation and lactation period]	↓ In sexually dimorphic differences in activity (e.g. more movement, rearing and time in center by control females than males)		≤0.030 (low dose)					Kubo et al. (311)

3.0 Developmental Toxicity

Strain, route	Endpoint	Dose, mg/kg bw/day				Reference	
		NOAEL	LOAEL	BMD ₁₀	BMDL ₁₀		BMD _{1SD}
	Change in sexually dimorphic differences in locus ceruleus volume (i.e., larger in control females than males but larger in treated males than females)		≤0.030 (low dose)				
	Anogenital distance, day of testicular descent or vaginal opening, male or female sexual behavior, ventral prostate weight, sperm count or motility, estrous cycles, histopathology in testis or ovary, or SDN-POA volume.	≥0.3 (high dose)					
Sprague Dawley, orally by pipette before mating through gestation and lactation period	Changes in various parameters associated with somatostatin receptor subtype 3 or GABA		≤0.040 (low dose)				Facciollo et al. (313)
F344/N, oral GD 10 through PND 20	↓ Male postnatal body weight ↓ Female postnatal body weight	4	40 ≤4 (low dose)				Negishi et al. (315)
	↑ Immobility by females ↓ Response in avoidance testing and ↑ grooming by adult males	4	40 ^{a,c} 4 ^{a,c,e}				
F344, gavage of pups PND 1–14 with	↓ Water maze performance (i.e. time spent in escape quadrant) by females ↓ In sexually dimorphic differences in water maze performance (i.e. better acquisition by control males than female)	0.1	0.25 ≤0.1 (low dose) ^c				Carr et al. (322)
Sprague Dawley dietary GD 6 through lactation	↑ Serum thyroxine levels only on PND 15		≤1 (low dose)				Zoeller et al. (308)

3.0 Developmental Toxicity

Strain, route	Endpoint	Dose, mg/kg bw/day						Reference
		NOAEL	LOAEL	BMD ₁₀	BMDL ₁₀	BMD _{1SD}	BMDL _{1SD}	
Wistar, sc GD 8 to 23	↑ RC3/neurogranin mRNA expression in males or 0.040		≤1 (low dose)					Ramos et al.(286)
	↑ Relative area of prostate vimentin-positive cells		≤0.025 (low dose) ^a					
	↓ Relative area of prostate α-smooth muscle actin positive cells		0.025					
	↓ Area of androgen receptor-positive cells in prostate periductal stroma		0.025 ^a					
	↓ Area of prostatic acid phosphate-positive cells in epithelial cells		0.025 ^a					
Sprague Dawley, sc GD 11–20	↑ Prominent nipples and areolas in males and females	50	400					Naciff et al. (246); Naciff et al. (288)
Fischer 344 pups sc PND 1–5	↑ Serum prolactin in males and females		≤20 (low dose) ^a					Khurana et al. (331)
F344 pups sc injected PND 1–21	↑ In secretory granules and ↓ in microvilli on glandular epithelium of ventral prostate	0.004	0.020					Fukumori et al. (332)
Sprague Dawley pups sc PND 0–9	↓ Body weight in lactation period	105	427	286	200	233	156	Kato et al. (333)
	↑ Age of vaginal opening	26	105	345	267	159	116	
	↓ No. with normal estrous cycles	105	427	81	28			
	↑ No. with cleft clitoris	26	105	299	failed			
	↓ Ovary weight	105	427	85	59	140	93	
	↓ Uterus, wet weight	105	427	66	55	128	96	
	↓ Uterus, blotted weight	105	427	273	128	318	168	
	↓ Uterine fluid weight	26	105	42	34	139	104	

3.0 Developmental Toxicity

Strain, route	Endpoint	Dose, mg/kg bw/day						Reference
		NOAEL	LOAEL	BMD ₁₀	BMDL ₁₀	BMD _{1SD}	BMDL _{1SD}	
	↑ No. with polycystic ovaries		≤105 (lowest dose examined)	81	24			
	↓ No. corpora lutea	105	427	238	90			
	↓ No. with corpora lutea	105	427	65	38	137	83	
	↓ Corpora lutea area		≤105 (lowest dose examined)	42	37	84	66	
Wistar pups sc PND 1, 3, 5, 7, 9, and 11	↑ Deformed acrosome, deformed nucleus, and abnormal ectoplasmic specialization in sperm	0.001	0.010 ^d					Toyama and Yuasa (334)
Sprague Dawley pups, sc PND 0 to 9	Age of preputial separation; copulation rate; fertility; sperm count, motility, or morphology; serum testosterone level, histopathology of testis; or prostate weight	1 (high dose)						Kato et al. (335)

^aThere was little-to-no evidence of a dose-response relationship.

^bEstimated doses in dams during gestation and lactation

^cNo effects were observed at one or more higher dose levels

^dNo information available for individual doses

^eEffects were observed at some but not all periods of evaluation

3.0 Developmental Toxicity Data

1 **Table 98. Summary of Developmental Toxicity in Single Dose-Level Rat Studies**

Species, strain, route	Dose, mg/kg bw/day	Significant developmental findings	Reference
Rat, Wistar, drinking water	2.5, apparently during gestation and lactation	↓ Sexually dimorphic difference in corticotropin-releasing hormone neurons in anterior stria terminalis	Funabashi et al. (284)
Rat, Wistar, drinking water	0.015, from GD 13 to PND 0	↓ Sexually dimorphic differences for rearing and struggling during swim test; ↑immobility of males during swim test; ↑ diving by females in swim test (leading to sexual dimorphic behavior not observed in controls)	Fujimoto et. al. (285)
Rat, Wistar, dietary	1000–1600, prior to mating through gestation and lactation periods.	Variable effects on body weight	Takashima et al. (299)
Rat, Long Evans, gavage	0.0024, GD 12 through PND 21	↑ Body weight; ↓ paired testes and seminal vesicles weight; ↓ testicular testosterone level; and ↓ basal and LH-induced ex vivo testosterone production	Akingbemi et al. (306)
Rat, Long Evans, gavage	0.0024, PND 21–90	↑ Serum LH level; ↓ weight of seminal vesicles; ↓ testicular testosterone level; and ↓ basal and LH-induced ex vivo testosterone production.	Akingbemi et al. (306)
Rat, Sprague Dawley, gavage	100, PND 23–53	↑ Age of preputial separation; ↑ kidney and thyroid weights; ↓ liver weight; ↓ cortical thickness of the kidney; ↑ hydronephrosis; ↑ multinucleated giant cells in seminiferous tubules; ↓ no. undergoing spermatogenesis	Tan et al. (307)
Rat, Sprague Dawley, oral by pipette	0.040 during pregnancy and lactation (through cross-fostering treated offspring were indirectly exposed only during gestation or lactation)	In response to formalin injection: ↓ paw jerking with postnatal exposure; ↑ paw flexion with prenatal exposure	Aloisi et al. (314)
Rat, F344/N, gavage	0.1 from GD 3 to PND 20	↓ Correct avoidance responses and ↑ failure of avoidance in active avoidance testing; no ↑ in locomotion following trans-2-phenylcyclopropylamine hydrochloride challenge in males	Negishi et al. (316)

3.0 Developmental Toxicity Data

Species, strain, route	Dose, mg/kg bw/day	Significant developmental findings	Reference
Rat, Sprague Dawley, orally by pipette	0.040 from 10 days prior to conception until weaning of pups	↓ Head dipping and crosses by females in wholeboard test; ↓ time in center and ↑ time grooming by females in elevated maze test; ↑ open arm entry and time in open arms, and ↓ stretched posture by males in elevated maze test.	Farabollini et al. (317)
Rat, Sprague Dawley, orally by pipette	0.400 from GD 14 through 6 days following delivery of pups	↓ Head dipping by males and females in wholeboard test; ↓ time in center, open arm entries, and entries by females in elevated maze test; ↑ open/total entries and ↓ stretched posture by males in elevated maze test	Farabollini et al. (317)
Rats, Sprague Dawley orally by pipette	0.040 mg/kg bw/day from mating through weaning of pups (through cross-fostering treated offspring were indirectly exposed only during gestation or lactation).	↑ defensive behavior, ↓ ambivalent behavior, and ↑ defensive/agonistic behaviors by prenatally exposed males in intruder testing; ↓ exit latency in diestrus and proestrus and ↑ lordosis frequency in proestrus in sexual behavior testing of pooled group of gestationally and lactationally exposed females; ↑ intromissions in postnatally exposed males and ↑ intromission latency and genital sniffing in prenatally exposed males	Farabollini et al. (318)
Rats, Sprague Dawley orally by pipette	0.040 from 10 days prior to conception until weaning of pups	In pooled results for all ages: ↑ play with females by females; ↓ sociosexual exploration by males; ↑ social interest by males In results for PND 35: ↑ social interest by males and females; ↓ low-intensity mating elements by females; ↓ sociosexual exploration by males	Dessi-Fulgheri et al. (319)
	0.400 from GD 14 through PND 6	In pooled results for all ages: ↓ low-intensity mating elements in males and females; ↓ sociosexual exploration in males and females; ↓ social interest in males and females; In results for PND 35: ↓ low-intensity mating elements; ↓ sociosexual exploration by males	Dessi-Fulgheri et al. (319)

3.0 Developmental Toxicity Data

Species, strain, route	Dose, mg/kg bw/day	Significant developmental findings	Reference
Rat, Sprague Dawley, oral by pipette	0.040 from mating through gestation periods.	↓ Time spent in novel area by females and ↑ activity in novel area by males and females; changes in impulsive behaviors [not clear due to a possible error in study figure labeling]; ↓ rearing and crossing behaviors in response to d-amphetamine by males	Adriani et al. (320)
Rat, Sprague Dawley, oral by pipette	0.040 during gestation and lactation	↑ Social and non-social exploration, ↓ play with males, and ↓ grooming by female rats.	Porrini et al. (321)
Rat pup, Wistar, sc	37 from PND 2–12	Transient changes: ↓ testis weight at 35 days of age and ↓ epithelial cell height in the efferent ducts at 18 and 25 days of age	Fisher et al. (324)
Rat pup, Sprague Dawley, sc	300 from PND 1 to 5	No effects on age of vaginal opening or preputial separation, copulation or fertility indices, sexual behavior of males, histopathological alterations in males, or female reproductive organs, or effects on SDN-POA. [Panel noted possible ↑ number of apically located nuclei in prostate, but a definitive conclusion could not be made based on 1 photograph]	Nagao et al. (325)
Rat, Wistar, sc	50 from PND 22 to 32	↑ Serum prolactin levels on PND 29 but not PND 120; ↑ lateral but not ventral prostate weight; ↑ focal luminal polymorphonuclear cellular infiltrate in prostate	Stoker et al. (326)
Rat pup, Wistar, sc	100 from PND 2 to 12.	No histological evidence of prostate inflammation Advanced testicular lumen formation, ↑testis weight, ↑Sertoli cell volume/testis, ↑ spermatocyte nuclear volume/unit Sertoli cell, and ↑ plasma FSH on PND 18; ↑ plasma FSH on PND 25; ↑ testicular weight in adulthood	Atanassova et al. (327)
Rat pup, Wistar, sc	20–100, from PND 2 to 12	No effects on gross structure of or ERβ, ERα, androgen or progesterone receptor proteins in the seminal vesicle	Williams et al. (328)

3.0 Developmental Toxicity Data

Species, strain, route	Dose, mg/kg bw/day	Significant developmental findings	Reference
Rat pup, Wistar, sc	4–20 on PND 2, 4, 6, 8, 10, and 12	No effects on plasma testosterone, rete testis luminal area, efferent duct luminal area, efferent duct epithelial cell height, or vas deferens epithelial cell height	Rivas et al. (329)
Rat pup, Wistar, sc	20–100 from PND 2 to 12	↑ Plasma testosterone on PND 18. No effects on testis weight, percent Leydig cell nuclear volume/testis, Leydig cell nuclear volume/testis, total Leydig cell volume (nuclear + cytoplasmic volume/testis)	Sharpe et al. (330)
Rat pup, Sprague Dawley, sc	0.010 on PND 1, 3, and 5; half the rats exposed to 17β-estradiol and testosterone in adulthood	In rats with no 17β-estradiol and testosterone exposure in adulthood: no effects on dorsal prostate weight, histopathology alterations, proliferation index, or apoptotic index. In rats with 17β-estradiol and testosterone exposure in adulthood: ↑ incidence and severity of prostatic intraepithelial neoplasia; ↑ proliferation and apoptosis in regions of prostatic intraepithelial neoplasia	Ho et al. (336)
Rat pup, Sprague Dawley, sc	100 from PND 1 to 19	In Anteroventral periventricular nucleus of females: ↓ cells positive for both ERα + tyrosine hydroxylase. In Anteroventral periventricular nucleus of males: ↑ cells positive for tyrosine hydroxylase and ↓ percent cells positive for both tyrosine hydroxylase and ERα	Patisaul et al. (338)

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3.0 Developmental Toxicity Data

1 **Table 99. Summary of Developmental Toxicity in Multiple Dose Mouse Studies**

Strain, route	Endpoint	Dose, mg/kg bw/day				Reference		
		NOAEL	LOAEL	BMD ₁₀	BMDL ₁₀		BMD _{1SD}	BMDL _{1SD}
CD-1, gavage GD 6–15	↑ Resorptions/litter	1000	1250	817	377	1245	1162	Morrissey et al. (273)
	↓ Fetal body weight/litter	1000	1250	1079	785	1249	1024	
CF-1, by pipette GD 11–17	Sperm production efficiency	0.002	0.020	0.011	0.007	0.010	0.007	vom Saal et al. (341) ^a
	↑ Preputial weight and ↓ seminal vesicle and epididymis weight		0.002 ^{b,c}					
CF-1, by pipette GD 11 to 17	↑ Prostate weight		≤0.002 (low dose)					Nagel et al. (205); vom Saal et al. (341)
CF-1, by pipette GD 11 to 17	Prostate, preputial gland, seminal vesicle, or epididymis weight; cauda epididymal sperm concentration, daily sperm production, or efficiency of sperm production; testicular histopathology.	≥0.2 (high dose)						Cagen et al. (342)
CF-1, by pipette GD 11 to 17	Prostate weight and sperm production.	≥0.020 (high dose)						Ashby et al. (343)
ICR, oral GD 6.5–13.5 or 6.5–17.5	↑ mRNA expression for arylhydrocarbon receptor in brain, testis, and ovary.		≤0.00002 (low dose) ^c					Nishizawa et al. (354)
	↑ mRNA expression for retinoic acid α receptor in brain and ovary.		≤0.00002 (low dose) ^c					
	↑ mRNA expression for retinoic acid α receptor in testis.	0.20	20					
	↑ mRNA expression for retinoid X α receptors in brain.		≤0.00002 (low dose) ^c					
	↑ mRNA expression for retinoid X α receptor in testis and ovary.	0.002	0.020 ^c					

3.0 Developmental Toxicity Data

Strain, route	Endpoint	Dose, mg/kg bw/day				Reference		
		NOAEL	LOAEL	BMD ₁₀	BMDL ₁₀		BMD _{1SD}	BMDL _{1SD}
ICR, oral GD 6.5–13.5 or 6.5–17.5	↑ mRNA expression for arylhydrocarbon receptor, arylhydrocarbon receptor repressor, and arylhydrocarbon receptor nuclear translocator in brain, testis, and ovary.		≤0.00002 (low dose) ^c					Nishizawa et al. (355)
DBA/1 J, oral for 18 days beginning on day of mating	↑ Immune response	0.030	0.300					Yoshino et al. (356)
CD-1, oral by pipette GD 11–17	↑ Aggression ↓ Testis weight		≤0.002 (low dose) ^d ≤0.002 (low dose) ^{d,e}					Kawai et al. (358)
C57BL/6N, gavage GD 11–17 or PND 21 to 43	Sperm density or lesions in reproductive organs ↓ Absolute seminal vesicle weight in group exposed during gestation	≥0.200 (high dose)	≤0.002 (low dose) ^{b,c}					Nagao et al. (369)
ICR, drinking water from prior to mating through gestation and lactation	↓ Brain weight ↓ Kidney weight ↓ Testis weight		≤0.0013 (low dose in dams during pregnancy) ^{b,c} 0.0013 (in dams during pregnancy) 0.0013 (low dose in dams during pregnancy) ^b					Kabuto et al. (107)
C57BL/6, drinking water for 8 weeks, beginning at	↓ Expression of <i>ERβ</i> mRNA and ↑ expression of <i>ERα</i>	0.175	17.5					Takao et al. (370)

3.0 Developmental Toxicity Data

Strain, route	Endpoint	Dose, mg/kg bw/day						Reference
		NOAEL	LOAEL	BMD ₁₀	BMDL ₁₀	BMD _{1SD}	BMDL _{1SD}	
weaning ddy, oral form mating through lactation periods	mRNA in testis ↑ Preference for morphine- associated cage compartment		≤0.003 (low dose)					Miyatake et al. (374)
CD-1, dietary, multiple generations with exposure during pre- and postnatal development	↓ F1 body weight on PND 7, 14, and 21	50	600	548–560	267–313	580–617	370–506	Tyl et al. (376)
	↓ F1 male body weight at PND 21 necropsy	50	600	564	313	640	599	
	↓ F1 female body weight at PND 21 necropsy	50	600	387	254	776	598	
	Hepatic cytoplasmic variation, F1 male	5	50	124	92.5			
	Hepatic cytoplasmic variation, F2 male	50	600	224	178			
	Hepatic cytoplasmic variation, F1 female	5	50	333	200			
	Seminiferous tubule hypoplasia, F1 male	50	600	591	406			
	Seminiferous tubule hypoplasia, F2 male	5	50	283	233			
	Age of preputial separation, F1 parental or non-mated males	50	600	727–754	572–576	491–551	364–414	
	Anogenital distance per body weight, F1 male on PND 21	5	50	1373	607	1769	616	
	Postnatal survival; daily sperm production; efficiency of daily sperm production; sperm motility or morphology; estrous cyclicality; numbers of ovarian primordial follicles; mating or fertility indices; or adult prostate weight	≥600 (high dose)						

3.0 Developmental Toxicity Data

Strain, route	Endpoint	Dose, mg/kg bw/day				Reference		
		NOAEL	LOAEL	BMD ₁₀	BMDL ₁₀		BMD _{1SD}	BMDL _{1SD}
CD-1, sc GD 9–20	↓ Bromodeoxyuridine incorporation in mammary epithelium at 10 days of age		≤0.000025 (low dose) ^{d,e}					Markey et al. (360)
	↓ Bromodeoxyuridine incorporation in mammary stromal cells at 1 month of age	0.000025	0.000250 ^d					
	↑ Mammary duct, terminal duct, terminal end bud, and alveolar bud areas; ↑ bromodeoxyuridine incorporation in; stromal cells; and ↑ alveoli containing secretory products.		≤0.000025 (low dose) ^{d,e}					
CD-1 mice, sc, GD 9 through remainder of pregnancy	↑ Estrous cycle disruptions		≤0.025 (low dose) ^b					Markey et al. (361)
	↓ Vaginal weight Fluid-filled ovarian bursae	0.025	0.25 ≤0.025 (low dose)					
	↑ Alveolar buds/lobulo-alveoli in mammary at 6 months of age		≤0.025 \ (low dose) ^d					
	↓ Alveolar buds/lobulo-alveoli in mammary at 9 months of age Vaginal opening	0.25 (high dose)	≤0.025 \ (low dose) ^{b,c,d}					
ICR/Jcl, sc GD 11–17	↓ Female body weight at weaning		≤0.002 (low dose) ^b	0.065	0.017	0.088	0.021	Honma et al. (364)
	↓ Male body weight at birth	0.002	0.020	0.054	0.020	0.031	0.015	
	↑ Anogenital distance of females at weaning		≤0.002 (low dose) ^{b,c}					

3.0 Developmental Toxicity Data

Strain, route	Endpoint	Dose, mg/kg bw/day				Reference		
		NOAEL	LOAEL	BMD ₁₀	BMDL ₁₀		BMD _{1SD}	BMDL _{1SD}
ICR, sc GD 7 to 18	↑ Anogenital distance of males on PND 60		≤0.002 (low dose)	0.035	0.020	0.035	0.020	Iwasaki and Totsukawa (365)
	↓ Age at vaginal opening	0.002	0.020					
	↓ Body weight at vaginal opening		≤0.002 (low dose)					
	↓ Age at 1 st estrus	0.002	0.020					
	↑ Estrous cycle length		≤0.002 (low dose) ^b	0.021	0.007	0.12	0.021	
	↑ Cornified cells		≤0.002 (low dose) ^c	0.17	0.020	0.44	0.021	
	↓ Lymphocytes in vaginal smear		≤0.002 (low dose) ^c	0.26	0.020	0.26	0.020	
	↓ Birth weight		≤0.00025 (low dose) ^c					
	↓ Pup viability on PND 4		≤0.00025 (low dose) ^c					
	↑ Age of vaginal opening		≤0.00025 (low dose) ^c					
	↓ Age of vaginal opening		2.5					
Outbred CD-1 (ICR), sc GD 15–18	↓ Uterine weight following 17β-estradiol exposure		≤0.00025 (low dose) ^c					Nikaido et al. (366)
	↑ Uterine weight following 17β-estradiol exposure		0.025 ^c					
	↑ Body weight gain		≤0.5 (low dose) ^e					
	↓ Age of vaginal opening	0.5	10					
	↑ Estrous cycle length		≤0.5 (low dose)					
ICR, ip every 3 days, beginning on day of mating	↑ No. with no corpora lutea and vaginal cornification at 4 weeks of age		≤0.5 (low dose) ^d					Park et al. (367)
	↓ Fetal body weight	0.5	5					

3.0 Developmental Toxicity Data

Strain, route	Endpoint	Dose, mg/kg bw/day						Reference
		NOAEL	LOAEL	BMD ₁₀	BMDL ₁₀	BMD _{1SD}	BMDL _{1SD}	
ICR, ip every 3 days, beginning on day of mating	↓ Male body weight	0.5	5					Park et al. (368)
	Male or female reproductive organ weights or sperm parameters	5 (high dose)						
ICR/Jcl, sc GD 10–18 (female offspring ovariectomized)	↑ No. of vaginal epithelial layers		≤10 (low dose)					Suzuki et al. (377)
	↓ No. with corpora lutea		≤10 (low dose) ^c					
ICR/Jcl, sc for 5 days beginning at birth (mice later ovariectomized except those used to monitor estrous cycles)	↑ Mitotic rate in uterine stromal cells and vaginal epithelial cells	10	100					Suzuki et al. (377)
	↑ Vaginal epithelial layers	10	100					
	↑ No. with polyovular follicles and no. polyovular follicles/mouse	10	100					
	Estrous cyclicity	100 (high dose)						
CD-1, sc GD 9 through PND 4	↓ Vaginal weight	0.000025	0.000250	0.00011	0.000075	0.00022	0.000129	Markey et al. (379)
	↓ Uterine lamina propria volume	0.000025	0.000250	0.000098	0.000059	0.00025	0.00014	
	↑ Uterine epithelial glandular cells incorporating BrdU	0.000025	0.000250					
	↑ Uterine epithelial luminal cells expressing ER α		≤0.000025 (low dose)					
	↑ Uterine epithelial cells expressing progesterone receptor		≤0.000025 (low dose) ^b					
CD-1, sc GD 9 through PND 3 (not defined)	↑ Numbers of mammary terminal end bud and mammary gland branches		≤0.000025 (low dose)					Muñoz-de-Toro et al. (363)
	↓ Mammary apoptotic cells		≤0.000025 (low dose) ^c					
	↓ BrdU incorporation by mammary stromal cells	0.000025	0.000250					

3.0 Developmental Toxicity Data

Strain, route	Endpoint	Dose, mg/kg bw/day				Reference	
		NOAEL	LOAEL	BMD ₁₀	BMDL ₁₀		BMD _{1SD}
	↑ Mammary epithelial cells expressing progesterone receptors		0.000025 ^b				
	↓ Correlation between age of first estrous and mammary length	0.000025	0.000250				
SHN, injection on first 5 days of life	↓ Sperm count	0.25	25				Nakahashi et al.(380)
SHN, injection on first 5 days of life	↓ Sperm count	0.25	25				Aikawa et al. (381)
	Histopathological alterations in testes	≥25 (high dose)					
ICR, sc PND 1, 3, 5, 7, 9, and 11	↑ Deformed acrosome, deformed nucleus, and abnormal ectoplasmic specialization in sperm	0.0001	0.001				Toyama and Yuasa (334)

^aPanel members had differing opinions regarding the utility of the study.

^bThere was little-to-no evidence of a dose-response relationship.

^cNo or opposite effects were observed at one or more higher dose level.

^dThe effect was observed at some but not other time points of evaluation.

^eMagnitude of response was greater at lower than higher doses.

3.0 Developmental Toxicity Data

1 **Table 100. Summary of Developmental Toxicity in Single Dose-Level Mouse Studies**

Strain, route	Dose, mg/kg bw/day	Significant developmental findings	Reference
CF1, oral	0.0024 on GD 11–17	↑ Body weight at weaning; ↓ postnatal pup survival; ↓ period between vaginal opening and first estrus	Howdeshell et al. (345)
CD-1, oral	0.050 on GD 16–18	No effect on age of vaginal opening ↑ Anogenital distance adjusted for body weight on PND 3, 21, and 60; ↑ prostate weights on PND 3, 21, and 60; ↓ relative (to body weight) epididymis weight in the bisphenol A group on PND 60; ↑ androgen receptor binding on PND 21 and 60	Gupta (347)
CD-1, oral by pipette	0.010 on GD 14–18	↑ No. of prostate ducts and proliferating cell nuclear antigen staining in dorsolateral prostate; ↑ prostate duct volume in dorsolateral and ventral prostate	Timms et al. (351)
CD-1, oral by pipette	0.010 on GD 14–18; offspring mated and dosed with 0 or 0.010 on GD 14–18.	In mice exposed only during gestational development or in adulthood during pregnancy: ↓ time nursing and in nest and ↑ time nest building, resting alone, grooming, and out of nest In mice exposed during both gestational development and in adulthood during pregnancy: ↑ time resting alone	Palanza et al. (352)
ICR, oral	0.002 mg/kg bw/day from 6.5–11.5, 6.5–13.5, 6.5–15.5, and 6.5–17.5 days post coitum	Variable changes in retinoic acid retinoid X receptors α mRNA expression in brain, ovary, and testis, depending on brain region and day of exposure	Nishizawa et al. (353)
CD-1, oral from syringe	0.010 on GD 11–18	↓ Place preference associated with <i>d</i> -amphetamine in females	Laviola et al. (359)

4.0 REPRODUCTIVE TOXICITY DATA

4.1 Human

4.1.1 Female

Takeuchi and Tsutsumi (65), supported by the Japanese Ministry of Education, Science, Sports, and Culture, the Ministry of Health and Welfare, and the Science and Technology Agency, measured bisphenol A in the blood serum of 14 healthy women, 11 healthy men, and 16 women with polycystic ovary syndrome [**diagnostic criteria not discussed**]. The healthy women were evaluated in the mid-follicular phase of the menstrual cycle. Bisphenol A was measured using a competitive ELISA. Serum was also evaluated for total and free testosterone, 17β -estradiol, androstenedione, dehydroepiandrosterone sulfate, LH, FSH, and prolactin. Statistical analysis was by ANOVA with least squares difference test. Correlation coefficients were obtained from a linear regression analysis. Mean \pm SEM bisphenol A serum concentrations (ng/mL) were 0.64 ± 0.10 in normal women, 1.49 ± 0.11 in normal men, and 1.04 ± 0.10 in women with polycystic ovary syndrome. Bisphenol A serum concentrations were significantly correlated with total testosterone ($r = 0.595$) and free testosterone ($r = 0.609$) in all subjects and in all female subjects ($r = 0.559$ for total testosterone and 0.598 for free testosterone). Bisphenol A serum concentrations were not significantly correlated with any other hormone measures. The authors concluded that either bisphenol A stimulates testosterone production or metabolism of bisphenol A is inhibited by testosterone. They further suggested that displacement of sex steroids from sex-hormone binding globulin by bisphenol A might disrupt the estrogen-androgen balance.

Strengths/Weaknesses: Quality assurance for the hormone radioimmunoassays appeared adequate; however, there was no standardization for time of day for the serum samples, which can cause a lot of variation in male testosterone samples. ELISA has not been standardized for human sera, although the authors cited a 0.97 correlation between this assay and the better quality HPLC analysis. Very little information was given on the selection of the comparison group beyond mean age and body-mass index. Very little information was given on recruitment methods, and participation rates/exclusions are unknown. The lack of diagnostic criteria for polycystic ovary syndrome is a weakness. No potential confounders were identified except mean age and body-mass index. Mean values appear to have been similar between groups. The positive correlations between bisphenol A level and total/free testosterone levels in all women and in entire study group were noted, but there was no adjustment for confounders or effect modifiers in these correlation/regression analyses. No information was given on whether the data were normally or lognormally distributed. It is difficult to know whether the differences found are meaningful, even though they are consistent with each other. The study was limited by small numbers in each group; power was not evaluated, and the results should be regarded as descriptive. The authors did not address whether there might be alternative explanations for the observed differences (e.g., differences in exposures).

Utility (Adequacy) for CERHR Evaluation Process: This paper gives some insight for potential mechanisms affecting the levels of bisphenol A in the body. These data could be compared to those found in other studies of humans and laboratory animals to look for consistency. These data also highlight potential pitfalls in exposure classification/definition present in studies of exposure to bisphenol A and possible health outcomes. Bisphenol A appears to be associated with increased testosterone and free testosterone in men and a group of women with polycystic ovary syndrome, but this study's utility is limited by its small size, crude design, crude tests, and inadequate analyses. Although many methodologic flaws limit the study's utility, the results are consistent with those of other surveys. It is interesting that the observation of a non-statistically significant negative correlation between bisphenol A and FSH in the entire study group was also reported by Hanaoka et al.(82).

4.0 Reproductive Toxicity Data

1 **Takeuchi et al. (64)**, supported by the Japanese Ministry of Education, Science, Sports, and Culture, the
2 Ministry of Health, Labor, and Welfare, the National Institute for Environmental Studies, and the Science
3 and Technology Agency, examined relationships between serum sex hormone and bisphenol A
4 concentrations in women with ovarian dysfunction and obesity. Fasting blood samples were collected
5 during the midfollicular phase from 19 non-obese and 7 obese healthy women with normal menstrual
6 cycles. Blood samples were also obtained from 7 women with hyperprolactinemia, 21 patients with
7 hypothalamic amenorrhea, and 13 non-obese and 6 obese patients with polycystic ovary syndrome. **[It**
8 **not known whether any of these subjects were the same as those reported earlier by this group**
9 **(65).]** Mean ages in each group were ~25–29 years. Blood serum was analyzed for bisphenol A levels
10 using an ELISA technique, and total and free testosterone, 17β -estradiol, androstenedione,
11 dehydroepiandrosterone sulfate, LH, FSH, prolactin, and insulin levels were measured using by RIA.
12 Statistical analyses included ANOVA, least significant difference test, and linear regression analysis.
13

14 Compared to non-obese healthy women, concentrations of bisphenol A in serum were significantly higher
15 **[% increase compared to healthy non-obese controls]** in non-obese women with polycystic ovary
16 syndrome **[48%]**, obese women with polycystic ovary syndrome **[65%]**, and obese healthy women
17 **[46%]**. Significant positive correlations were found between bisphenol A level in serum and body mass
18 index ($r = 0.500$) and serum levels of total testosterone ($r = 0.391$), free testosterone ($r = 0.504$),
19 androstenedione ($r = 0.684$), and dehydroepiandrosterone sulfate ($r = 0.514$). The study authors
20 concluded that there is a strong relationship between serum levels of bisphenol A and androgens, possibly
21 due to androgen effects on metabolism of bisphenol A.
22

23 **Strengths/Weaknesses:** Quality assurance for the hormone radioimmunoassays appears adequate. In
24 contrast to the 2002 article by these authors (65), blood draws were time-standardized to 9:00–10:00 AM
25 after overnight fasting. The authors cited a 0.97 correlation between the ELISA, which was not
26 standardized for human sera, and the better quality HPLC analysis, which was not used. It was not clear
27 whether any of the women in this study were also reported by the authors in their 2002 publication. No
28 potential confounders or effect-modifiers were identified except mean age and body-mass index, and
29 neither of these was controlled. There were positive correlations between bisphenol A level and body-
30 mass index, total testosterone, free testosterone, androstenedione, and dehydroepiandrosterone sulfate for
31 all study groups. These correlations are also found (with the exception of total testosterone) in the control
32 (“normal women”) group as well. Many of the hormones were likely to have log distributions but were
33 clearly not transformed prior to analysis. No information was given on whether the data were normally or
34 lognormally distributed, and there was no adjustment for age, body-mass index, and multiple other
35 potential confounders/effect modifiers. It is difficult to know whether the differences found were
36 meaningful, even though they were consistent with each other. Study power was not assessed, and results
37 should be regarded as descriptive.
38

39 **Utility (Adequacy) for CERHR Evaluation Process:** Bisphenol A appeared to be associated with
40 androgens (testosterone, free testosterone, androstenedione, dehydroepiandrosterone sulfate) and
41 conditions that may promote hyperandrogenism (obesity, polycystic ovarian syndrome) in this cross-
42 sectional survey with very small numbers of women in each group. There were some methodologic im-
43 provements (standardization of time of blood draw), and results are relatively consistent with the authors’
44 2002 survey; although, it was not clear whether the same groups of women were being re-studied. The
45 utility of the study was reduced by the remaining methodologic flaws including very small sample size,
46 crude ELISA test for bisphenol A, and inadequate adjustment for confounders/effect modifiers.
47

48 **Sugiura-Ogasawara et al. (67)**, supported by the Japanese Ministry of Health, Labor, and Welfare,
49 conducted a study to determine if there is an association between recurrent miscarriage and bisphenol A
50 levels in blood. The cases in this study were 45 patients with a history of 3 or more (3–11) consecutive
51 first trimester miscarriages. Mean \pm SD age of the cases was 31.6 ± 4.4 . None of the cases had a history

4.0 Reproductive Toxicity Data

1 of live birth. All were seen at a Japanese hospital between August, 2001 and December, 2002. Half of the
2 cases were housewives and half were employed in various occupations. A hysterosalpingography
3 analyses was conducted in cases, and chromosome analyses were conducted for both cases and their
4 partners. Women were excluded from the study if uterine anomalies were observed or chromosomal
5 abnormalities were detected in either partner. Serum bisphenol A levels were determined by ELISA.
6 Immunological endpoints examined included antinuclear antibodies, antiphospholipid antibodies, and
7 natural killer cell activity. Blood testing for hypothyroidism, diabetes mellitus, and hyperprolactinemia
8 was conducted. Blood samples were obtained 5–9 days following ovulation in at least 2 cycles. Blood
9 samples to determine progesterone and prolactin levels were taken at 3 months following the last abortion
10 and prior to the next conception. For subsequent pregnancies, ultrasounds were conducted, and aborted
11 embryos/fetuses were karyotyped. Serum levels of bisphenol A in cases were compared to those of 32
12 healthy non-pregnant hospital employees with no history of live birth, infertility, or miscarriage. Mean \pm
13 SD age of controls was 32.0 ± 4.8 . None were taking oral contraceptives. Like the cases, the controls
14 lived near Nagoya City. Statistical analyses included Welch test, Mann-Whitney test, and Pearson
15 correlation coefficient.

16
17 Bisphenol A levels (mean \pm SD) were reported to be significantly higher in women with recurrent
18 miscarriages (2.59 ± 5.23 ng/mL) compared to healthy controls (0.77 ± 0.38 ng/mL). In the 45 cases,
19 incidences of abnormal conditions were 15.6% for hypothyroidism, 13.3% for antiphospholipid
20 antibodies, 22.2% for antinuclear antibodies, 11.1% for hyperprolactinemia, and 20.5% for luteal phase
21 defect. Serum levels of bisphenol A were significantly higher in patients who tested positive versus
22 negative for antinuclear antibodies (Mean \pm SD 7.382 ± 9.761 vs. 1.222 ± 1.54 ng/mL). Thirty-five of the
23 patients became pregnant and 48.6% had another miscarriage. Serum bisphenol A levels in patients who
24 miscarried were 4.39 ± 8.08 ng/mL, and serum bisphenol A in patients with successful pregnancies were
25 1.22 ± 1.07 ng/mL (not statistically significant). The study authors concluded that exposure to bisphenol
26 A is associated with recurrent miscarriage.

27
28 In a letter to the editor, Berkowitz (412) stated that this study did not support an association between
29 bisphenol A blood levels and recurrent miscarriage. Several limitations were noted for the study. Timing
30 and numbers of blood samples collected were not clearly defined. It was noted that because bisphenol A
31 has a short half life, it would be critical to know if blood samples were obtained in a timeframe relevant to
32 the occurrence of miscarriage. Although differences in serum bisphenol A levels in cases compared to
33 controls achieved statistical significance, it was noted that median levels of bisphenol A in serum were
34 nearly identical in patients with recurring miscarriages (0.71 ng/mL) and controls (0.705). The similarities
35 in median values suggested there were no differences between the two groups, and it was suggested that
36 apparent differences in mean serum levels of bisphenol A were due to a few individuals, as was
37 demonstrated in Figure 1 of the Sugiura-Ogasawara et al. (67) report. It was stated that the Welch test was
38 inappropriate for statistical analyses. It was noted that the 2 evaluation groups could not be considered
39 comparable because of differences in occupation (housewives compared to medical workers) and
40 unknown fertility of controls. Because the controls were not evaluated for factors such as hypothyroidism
41 and systemic lupus erythematosus (associated with antinuclear antibodies), the conditions may have been
42 overrepresented in cases and may have been the cause of the reported differences between the 2 groups.
43 Although it was noted that mean bisphenol A levels were (non-significantly) lower in women who
44 subsequently became pregnant and had a successful pregnancy compared to those who miscarried, it was
45 noted that the median level of bisphenol A was actually higher in women with the successful pregnancies.
46 Lastly it was noted that the ELISA method for measuring bisphenol A levels has not been validated and is
47 subject to inaccuracy due to extensive cross-reactivity.

48
49 In a response to the comments by Berkowitz (412), Sugiura-Ogasawara (413) stated that although
50 measurement of bisphenol A levels at various time points would have been ideal, obtaining samples every
51 day during pregnancy would have been difficult. Sugiura-Ogasawara clarified that bisphenol A values

4.0 Reproductive Toxicity Data

1 were based on a single sample in each individual, but that similar tendencies were observed for a second
2 blood sample. With respect to the use of women with live births as controls, Sugiura-Ogasawara
3 explained that the same blood samples were used for measurements of other environmental compounds,
4 some of which are known to decrease after delivery. It was noted that none of the cases had systemic
5 lupus erythematosus, and that use of controls with hypothyroidism or antinuclear antibodies was not
6 considered important for the study. Superiority of the HPLC method compared to the ELISA method for
7 measuring serum bisphenol A levels was acknowledged, but it was stated that the ELISA method was
8 used because of limited funding. It was reiterated that the study was preliminary and used a small number
9 of volunteers, and that additional studies using a larger sample and more appropriate analytical methods
10 were needed.

11
12 **Strengths/Weaknesses:** The letter from Berkowitz (412) summarizes many of the weaknesses of this
13 study. No quality assurance information was given for the biomarker/hormone measurements. As the
14 Berkowitz letter points out, the ELISA method is not standardized for human sera, the distribution of
15 exposure was not normal, and median values of the two groups were similar. The medians were similar in
16 the two groups, with two women skewing the mean. The comparison group was similar in age and body-
17 mass index to the patients, but occupationally dissimilar, including exposure to a number of potential
18 reproductive toxicants. The controls include more women working as physicians and nurses, suggesting
19 higher education levels and potentially socio-economic status, which could have translated into many
20 unexamined lifestyle and other differences. Fertility status was unknown. No information was given on
21 response rate, so the potential for response bias is unknown. It appears that the investigators included the
22 “controls” only for the comparison of the bisphenol A levels, since no data were presented on their health
23 conditions or subsequent pregnancies. One would expect that women with no pregnancy history would be
24 more likely to be younger than a group of women with up to 11 miscarriages, which was not the case
25 here, reinforcing the probable differences between the 2 groups. With the exception of age and body-mass
26 index, potential confounders and effect modifiers were not effectively managed, and not controlled in
27 analyses. The time between exposure and observation was not appropriate; patients had multiple
28 spontaneous abortions likely for various reasons. The authors’ conclusions require the assumption that
29 bisphenol A measurement levels represent those present during the spontaneous abortion event (this study
30 arose in women being evaluated for levels of DDE and polychlorinated biphenyls, which have longer
31 half-lives than does bisphenol A). Non-normal data were not appropriately transformed for analysis, and
32 means were inappropriately reported instead of medians or transformed means. Welch’s test was used
33 “...to compare bisphenol A levels...because the distribution of the two groups might have differed.”
34 Welch’s test is a *t*-test for groups with unequal variance, not different distributions (both should be
35 normal, which was probably not the case). Study Figure 1 (boxplots) was a nice representation of the data
36 which did not support the authors’ interpretation.

37
38 **Utility (Adequacy) for CERHR Evaluation Process:** Because of the design and analysis flaws in this
39 work, the only conclusion that can be drawn is that for some reason, very small numbers of women
40 (maybe 4 or 5) with repeated spontaneous abortions have elevated levels of bisphenol A and that a few of
41 these also have elevated antinuclear antibody levels. The reasons for those few individuals having
42 elevated levels are not clear.

4.1.2 Male

43
44 **Luconi et al. (414)**, supported by the Italian Public Health Project, examined the effects of in vitro
45 exposure of human spermatozoa to bisphenol A. Semen was collected from normozoospermic men, and
46 spermatozoa were separated. Intracellular calcium was measured using a spectrofluorimetric method in
47 cells treated with 1 μ M bisphenol A, 1 μ M 17 β -estradiol, 10 μ M progesterone, or the same concentrations
48 of bisphenol A in combination with 17 β -estradiol or progesterone. Effects on acrosome reaction were
49 examined using a fluorescent staining method in cells exposed to 1 μ M [0.23 mg/mL] bisphenol A for 2
50 hours, with and without exposure to 10 μ M progesterone. [In the study figures summarizing results,
51

1 **sample numbers in studies involving bisphenol A were listed at 5–11. It is not known if the sample**
 2 **numbers represented total numbers of sperm donors. Very few protocol details were provided in**
 3 **the methods section and many of the limited details presented above were obtained from the results**
 4 **section.]** Data were analyzed by Student *t*-test and 1-way ANOVA. Treatment of spermatozoa with
 5 bisphenol A resulted in a modest influx of calcium, but bisphenol A had no effect on calcium responses
 6 induced by 17 β -estradiol or progesterone. Bisphenol A exposure did not affect basal acrosome reaction or
 7 acrosome reaction induced by progesterone. Results were in contrast to those observed with 17 β -estradiol,
 8 which inhibited the acrosome reaction induced by progesterone. The study authors concluded that
 9 bisphenol A dose not likely interact with 17 β -estradiol or progesterone membrane receptors in human
 10 spermatozoa.

11
 12 **Strengths/Weaknesses:** This paper provides very limited information on the spermatozoa samples, such
 13 as the number of donors and the number of samples per donor. The study used Bisphenol A as a tool to
 14 investigate the structure and behavior of the plasma membrane-bound steroid receptor. While a novel
 15 approach, this study used a concentration of bisphenol A that was lower than some of the 17 β -estradiol
 16 concentrations, despite a significant literature showing that bisphenol A is less efficient at binding the ER
 17 than is 17 β -estradiol itself, and only 1 concentration was examined. Also, while the whole-cell approach
 18 is interesting, there was no other characterization of the structure or intra-cellular connections of the
 19 receptor, and no assessment was made of whether these modest effects altered the function or behavior of
 20 the sperm. The general dearth of information about these receptors and their function limits the use which
 21 can be made of these data.

22
 23 **Utility (Adequacy) for CERHR Evaluation Process:** This paper is not useful in the evaluation process.

24
 25 **Hanaoka et al. (82)**, supported by the Japanese Ministry of Health and Welfare and Ministry of
 26 Education, Science, Sports, and Culture, examined possible relationships between bisphenol A exposure
 27 and hormone levels in male workers. Exposed workers included 42 men in 3 Japanese plants who sprayed
 28 an epoxy hardening agent consisting of a mixture of bisphenol A diglycidyl ether (10–30%), toluene (0–
 29 30%), xylene (0–20%), 2-ethoxyethanol (0–20%), 2-butoxyethanol (0–20%), and methyl isobutyl ketone
 30 (0–30%). The workers were said to wear “protection devices” during spraying. Controls consisted of 42
 31 male assembly workers from the same plants who did not use bisphenol A diglycidyl ether, were within 3
 32 years of age to exposed workers (37 \pm 9 years vs. 38 \pm 10 years), and smoked the same number of
 33 cigarettes/day as exposed workers (21 \pm 7 vs. 21 \pm 6/day). **[Variances were not defined.]** Percentages of
 34 smokers were 86% in both groups, but percentages of alcohol drinkers were significantly lower in the
 35 exposed workers (43%) than in controls (57%). Urine and blood samples were obtained during periodic
 36 health examinations performed in June and July, 1999. Urinary bisphenol A was measured by HPLC, and
 37 urinary organic solvent metabolites were measured by GC or HPLC. Plasma LH, FSH, and free
 38 testosterone levels were measured by immunosolvent assay in a commercial laboratory. Data were log
 39 transformed and compared by paired *t*-test, Pearson correlation coefficient, and chi-squared test.
 40 Adjustments were made by linear regression for age and drinking habits, which were considered possible
 41 confounders.

42
 43 Urinary bisphenol A concentrations were significantly higher in exposed workers (median: 1.06
 44 μ mol/mol creatinine; range: <0.05 pmol to 11.2 μ mol/mol creatinine) than in controls (median: 0.52
 45 μ mol/mol creatinine; range: <0.05 pmol to 11.0 μ mol/mol creatinine). Average difference was reported as
 46 2.5 (95% CI 1.4–4.7; *P* = 0.002). Bisphenol A was not detected in 3 exposed workers and 1 control.
 47 Urinary solvent metabolites were detected more frequently in exposed workers than controls. No
 48 differences in plasma testosterone or LH concentrations were observed between exposed workers and
 49 controls. Plasma FSH concentrations were significantly lower in exposed workers (median: 5.3 mIU/mL;
 50 range: 4.0–8.3 mIU/mL) than in controls (median 7.6 mIU/mL; range 5.4–11.0 mIU/mL; average
 51 difference = 1.3; 95% CI –1.5 to –1.0). A “mild correlation” was reported between urinary bisphenol A

4.0 Reproductive Toxicity Data

1 and FSH ($r = -0.20$, $P = 0.071$) but was not observed for urinary solvent levels. A statistically significant
2 relationship was observed between FSH and bisphenol A following adjustment for alcohol intake ($r =$
3 -0.23 ; $P = 0.045$). The study authors concluded that bisphenol A may be generated endogenously
4 following exposure to bisphenol A diglycidyl ether, and bisphenol A may disrupt gonadotropic hormone
5 secretion in men.

6
7 **Strengths/Weaknesses:** Quality assurance for the hormone radioimmunoassays appeared adequate.
8 Blood draws and urine samples were time standardized between 10 AM and 12 noon. Reference values
9 were given and population values were considered in the discussion. Data were considered lognormal and
10 evaluated nonparametrically or transformed. Use of HPLC for bisphenol A and standard methods for the
11 other urinary metabolites with creatinine-adjustment are strengths. The epoxy sprayer workers were
12 matched to coworkers from other parts of the process. Exposure measurements were compared and
13 discussed in both groups with the understanding that the comparison group was unlikely to be truly
14 unexposed. All selected workers participated in the study. Analyses were adjusted for age and alcohol
15 use, and workers were matched on age (± 3 years) and cigarette use. A plausible ($P = 0.07$) correlation
16 between bisphenol A and decreasing FSH was reported. The authors took care to note that all levels were
17 within the clinical normal range. Correlations between other workplace exposures and hormones were not
18 observed. Blood and urine samples were collected concurrently, but not on the first day of the week.
19 According to Brock et al. (415), urine glucuronides of bisphenol A are a longer-lived (12-48h) biomarker,
20 so the sampling appears to have been appropriate. Statistical methods were appropriate to the study size
21 and distribution of the data. Alcohol use, which varied between the groups, was controlled. Non-normal
22 distributions were transformed or treated as non-normal. Biomarker data were handled appropriately in
23 analysis. The analyses performed were appropriate to a cross-sectional study of 84 male workers, 42 per
24 group, which was a small but reasonable number for this type of study.

25
26 **Utility (Adequacy) for CERHR Evaluation Process:** This survey was methodologically very sound and
27 mechanistically thoughtful. The multiplicity of exposures is an unavoidable limitation of the study;
28 however, study was well designed and executed. The authors observed a relation between decreasing
29 plasma FSH and increasing exposure to a metabolic precursor of bisphenol A and speculated that
30 bisphenol A was suppressing FSH. The relatively small number of men means that there is less power to
31 find smaller differences, and so there is less confidence in the negative results. The investigators
32 appropriately noted that the changes in FSH levels are “. . . within clinically normal conditions,” and
33 concluded that the “clinical significance is still unclear.”

34 35 **4.2 Experimental animal**

36 Studies in this section examine reproductive endpoints after administration of bisphenol A to sexually
37 mature animals. Reproductive endpoints after administration of bisphenol A during pregnancy, the
38 neonatal period, or puberty are discussed in Section 3.2.

39 40 *4.2.1 Female*

41 42 *4.2.1.1 Rat*

43 **Goloubkova et al. (216)**, supported by the Brazilian National Council of Scientific and Technological
44 Development and the National University of Rio Grande Do Sul, examined the effects of bisphenol A
45 exposure on the uterus and pituitary of ovariectomized rats. Wistar rats (60–67 days old) were fed a
46 standard certified rodent diet. [No information was provided on housing or bedding materials.] Rats
47 were subjected to bilateral ovariectomy or sham surgery. At 14 days post-surgery, rats were randomly
48 assigned to groups of at least 6 animals. Rats were sc injected with bisphenol A (>99% purity) at doses of
49 11, 78, 128, or 250 mg/kg bw/day for 7 days. An ovariectomized vehicle control group was exposed to
50 the 50% DMSO vehicle. A sham-operated control group was not exposed to the vehicle. Rats were killed
51 following the dosing period, and body and uterine weight were measured. Trunk blood was collected for

4.0 Reproductive Toxicity Data

1 measurement of serum prolactin level by RIA. The anterior pituitary was weighed and preserved in 10%
2 formalin. An immunohistochemical technique was used to identify pituitary cells expressing prolactin. A
3 total of 3 or 4 rats/group were evaluated for prolactin-positive cells in the pituitary and 6–8 rats were
4 evaluated for the other endpoints. Data were analyzed by ANOVA followed by post hoc Student-
5 Neuman-Keuls test or Kruskal-Wallis ANOVA followed by post hoc Dunn test.

6
7 In the 250 mg/kg bw/day group, final body weight was 7% lower than in the ovariectomized vehicle
8 control group, and body weight gain was lower compared to the ovariectomized vehicle and sham
9 controls. There was no effect of treatment on food intake. A dose-related increase in uterine weight
10 occurred in all groups of rats exposed to bisphenol A compared to the ovariectomized vehicle controls,
11 but uterine weight in the bisphenol A groups was lower than in the sham controls. Ovariectomy resulted
12 in decreased pituitary weight in ovariectomized vehicle controls and in the bisphenol A 11 and 78 mg/kg
13 bw/day dose groups compared to sham controls. Pituitary weight did not differ from sham controls after
14 128 mg/kg bw/day bisphenol A and was greater than in sham controls after 250 mg/kg bw/day bisphenol
15 A. Basal prolactin levels did not differ between the sham and ovariectomized vehicle controls. Serum
16 prolactin levels were increased in the 128 and 250 mg/kg bw/day bisphenol A groups compared to the
17 ovariectomized vehicle controls. Ovariectomy reduced the numbers of prolactin-positive cells in the
18 pituitary. The number of prolactin positive cells in the pituitary was increased by 64% in the 250 mg/kg
19 bw/day group compared to the ovariectomized controls. The study authors concluded that the
20 reproductive tract and neuroendocrine axis of Wistar rats can respond to bisphenol A.

21
22 **Strengths/Weaknesses:** This comprehensive neurocrine assessment demonstrated that bisphenol A
23 exhibits 17 β -estradiol-like activity on the hypothalamic-pituitary axis of the Wistar rat. Weaknesses are
24 the absence of a positive control to demonstrate maximal response in endpoints examined, dose levels
25 required to induce response that were excessively high, and the sc route of administration, which bypasses
26 potential first-pass metabolism. Maximal response was similar to that in sham-treated rats; the only
27 apparent adverse effect was hyperprolactemia in ovariectomized rats.

28
29 **Utility (Adequacy) for CERHR Evaluation Process:** This study clearly demonstrated that bisphenol
30 exhibits 17 β -estradiol-like activity in ovariectomized rats at high dose levels of exposure. However, the
31 relevancy of the model for human risk assessment is limited because the route of administration/dosing
32 paradigm was not relevant and the magnitude of response in the endpoints examined did not exceed the
33 levels in sham-treated rats.

34
35 **Funabashi et al. (416)**, supported by Yokoyama City University, examined the effects of bisphenol A
36 exposure on expression of progesterone receptor mRNA in the brain of ovariectomized rats. The effects
37 of butylbenzyl phthalate were also examined but will not be discussed. **[No information was provided**
38 **on feed, caging, or bedding materials.]** Wistar rats were ovariectomized at 7–8 weeks of age. Ten days
39 following ovariectomy, 6 rats/group were sc injected with sesame oil vehicle, 10 mg bisphenol A **[purity**
40 **not reported]**, or 10 μ g 17 β -estradiol. Rats were killed 24 hours later and the preoptic area, medial basal
41 hypothalamus, and anterior pituitary were removed. Expression of mRNAs for progesterone receptor,
42 preproenkephalin, and neurotensin were assessed by Northern blot. Data were analyzed by ANOVA
43 followed by Fisher protected least significant difference test. Exposure to bisphenol A resulted in
44 increased expression of progesterone receptor mRNA in the preoptic area and anterior pituitary.
45 Bisphenol A did not affect expression of mRNA for neurotensin in the preoptic area or preproenkephalin
46 in medial basal hypothalamus. 17 β -Estradiol increased expression of mRNA for progesterone receptor in
47 the preoptic area, medial basal hypothalamus, and anterior pituitary and increased preproenkephalin
48 mRNA expression in medial basal hypothalamus. The study authors concluded that bisphenol A increases
49 expression of progesterone receptor mRNA in the preoptic area of adult ovariectomized rats.

4.0 Reproductive Toxicity Data

1 **Strengths/Weaknesses:** This study demonstrated that bisphenol A induced progesterone receptor mRNA
2 in the preoptic area, basal hypothalamus, and anterior pituitary, similar to 17 β -estradiol. However,
3 potential concomitant changes in progesterone receptor protein were not examined. Only a single dose
4 level was used and the route was sc.

5
6 **Utility (Adequacy) for CERHR Evaluation Process:** This study shows estrogen-like activity for
7 bisphenol A in the central nervous system; Hhowever, the relevancy of the model for human risk
8 assessment is limited because the route of administration (sc)/dosing paradigm (single dose level) and no
9 functional/physiological correlate was examined.

10
11 **Yamasaki et al. (128)** conducted a 28-day exposure study that provided some information on the
12 reproductive organs of male and female rats. [**Complete details of this study are included in Section 2.**
13 **Results for females are discussed in this section, and results for males are discussed in Section**
14 **4.2.2.1.**] CD rats were fed a commercial diet (MF Oriental Yeast Co.) and housed in stainless steel wire
15 mesh cages. Ten 7-week-old rats/sex/group were gavaged with bisphenol A at 0 (olive oil vehicle), 40,
16 200, or 1000 mg/kg bw/day for 28 days. Due to the death of 1 animal exhibiting clinical signs in the 1000
17 mg/kg bw/day group, the high dose was reduced to 600 mg/kg bw/day on the 8th day of the study. In an
18 additional study, rats were exposed to ethinyl estradiol at 0, 10, 50, or 200 μ g/kg bw/day for 28 days.
19 There were no treatment-related alterations in blood levels of thyroid hormones, FSH, LH, 17 β -estradiol,
20 prolactin, or testosterone. The numbers of females with diestrus lasting 4 or more days was increased in
21 the high-dose group. Relative weights of ovary and uterus were unaffected. No gross or histopathological
22 alterations were reported for reproductive organs. The study authors concluded that change in estrous
23 cyclicity was the only useful endpoint for evaluating the endocrine-mediated effects of bisphenol A. In
24 comparison, females from the mid- and/or high-dose 17 β -estradiol group experienced alterations in
25 estrous cyclicity, decreased ovarian weight, increased uterine weight, and histopathological changes in the
26 ovary, uterus, and vagina.

27
28 **Strengths/Weaknesses:** This study was well-conducted and comprehensive. Dose levels were
29 appropriate for the induction of some measure of toxicity (decreased body weight). Bisphenol A
30 administration resulted in minimal effects on female reproductive parameters (estrous cyclicity) even in
31 the presence of general toxicity; histopathological findings were noted in (mostly) non-reproductive
32 tissues. An effect on the estrous cycle at the 1000/600 mg/kg bw/day dose level was clearly evident.
33 However, the authors did not explain their definition of an abnormal cycle. It is not uncommon for rats to
34 exhibit minor changes in their estrous cycles (this finding was seen at the other two dose levels). In
35 addition, the monitoring of the cycle was limited to ~2–3 weeks with no pretest data. Therefore, a
36 definitive NOAEL for reproductive effects in the female cannot be established.

37
38 **Utility (Adequacy) for CERHR Evaluation Process:** This GLP study was well conducted with an
39 appropriate route of administration. The 1000/600 mg/kg bw/day dose produced changes in estrous
40 cyclicity consistent with an estrogenic effect. A definitive NOAEL for reproductive effects in the female
41 cannot be established based on this study because of the limitations and variability of the data.

42
43 **Spencer et al. (417)**, supported by NIH, evaluated the uterine response to bisphenol A before and after
44 deciduoma formation in pseudopregnant Sprague Dawley rats. [**Cage and bedding materials and feed**
45 **were not indicated.**] Adult females underwent mechanical cervical stimulation to induce
46 pseudopregnancy [**pseudopregnancy day not indicated**]. On pseudopregnancy day 4, deciduoma
47 formation was induced under ether anesthesia by antimesometrial uterine epithelial trauma, applied
48 through a laparotomy under ether anesthesia. Rats were treated with sc bisphenol A [**97% purity**] 0 or
49 200 mg/kg bw in alcohol/saline on pseudopregnancy days 1–4 and killed on pseudopregnancy day 5, or
50 treated on pseudopregnancy days 5–8 and killed on pseudopregnancy day 9. Uteri and pseudopregnancy
51 day 9 endometria were harvested. Uteri were weighed and homogenized for measurement of protein and

4.0 Reproductive Toxicity Data

1 DNA content. Inducible nitric oxide synthase activity, decidual prolactin-related protein mRNA, *ER*
 2 mRNA, and cytosolic ER binding sites were measured in uteri and/or endometria. Blood was obtained for
 3 determination of serum 17 β -estradiol and progesterone. [**n = 5 was indicated for some of the data**
 4 **presentations.**] Results are summarized in Table 101. The authors called attention to the difference in
 5 bisphenol A effect depending on whether exposure was prior to or after decidual induction. They
 6 concluded that there was a decrease in proliferation when bisphenol A was given during decidual
 7 induction, with a decrease in decidual proteins, in spite of a lack of differential effect on *ER* mRNA or
 8 cytosolic ER binding sites. The authors also concluded that bisphenol A activity appeared to be
 9 antagonized by progesterone [**although they probably meant that bisphenol A antagonized the action**
 10 **of progesterone**].
 11

12 **Table 101. Bisphenol A Effects on Pseudopregnant Rats**

Endpoint	Treatment period, pseudopregnancy day	
	1–4	5–8
Uterus		
Wet weight	↑1.4-fold	↓63%
Protein content	↑1.4-fold	↓64%
DNA content	↔	↓53%
Decidual prolactin-related protein mRNA ^a	↔	↓44%
<i>ER</i> mRNA ^a	↓29%	↓50%
Cytosolic ER-binding sites	↓57%	↓37%
Nitric oxide synthase activity ^a	↔	↓50%
Pseudopregnancy day 9 endometrium		
Decidual prolactin-related protein mRNA ^a	Not applicable	↓48%
<i>ER</i> mRNA ^a	Not applicable	↓43%
Nitric oxide synthase activity ^a	Not applicable	↓40%
Serum		
17 β -Estradiol	↔	↔
Progesterone	↔	↓49%

↑, ↓, ↔ Statistically significant increase, decrease, or no change compared to vehicle control.

^aEstimated from a study graph by CERHR.

From Spencer et al. (417).

13
 14 **Strengths/Weaknesses:** The report characterizes the effects of 200 mg/kg bw/day bisphenol A sc on
 15 decidual formation in SD rats. These data suggest that sc administration of bisphenol A on GD 0–4
 16 would enhance implantation (a 17 β -estradiol-like action); whereas administration after GD 4 would be
 17 detrimental to early rat fetal development (perhaps by blocking progesterone action and acting as a weak
 18 17 β -estradiol agonist). These data are intriguing, but the functional consequences of bisphenol A
 19 administration on implantation were not assessed and the sc route of administration was not appropriate.
 20

21 **Utility (Adequacy) for CERHR Evaluation Process:** These data are interesting, identifying potential
 22 pathways of hormonal disruption by bisphenol A. However, given that only 1 dose level was examined
 23 and the sc route of administration, these data are of limited value for human risk assessment.
 24

25 **Funabashi et al. (418),** supported by Yokohama City University, examined the effects of bisphenol A
 26 exposure on sexual behavior and progesterone receptor expression in adult rats. Wistar rats were
 27 ovariectomized at 7–8 weeks of age. [**No information was provided on feed, caging, or bedding**
 28 **materials.**] In two sets of experiments, an immunohistochemistry technique was used to measure
 29 expression of progesterone receptor in the preoptic area and ventromedial hypothalamus following

4.0 Reproductive Toxicity Data

1 bisphenol A exposure. In the first experiment, 3–5 rats/group were sc injected with sesame oil vehicle, 10
2 mg bisphenol A (~40 mg/kg bw) [**purity not reported**], or 10 µg 17β-estradiol (~40 µg/kg bw) 2 weeks
3 following ovariectomy. In the second experiment, ovariectomized rats (3–4/group) were sc injected with
4 bisphenol A at 0.001, 0.010, 0.1, or 1 mg (~0.004, 0.040, 0.4, or 4 mg/kg bw). Rats were killed the day
5 following dosing, and brains were removed and fixed in 2% paraformaldehyde. Statistical analyses
6 included ANOVA followed by Scheffé post hoc test and Kruskal-Wallis test. Sexual behavior was
7 examined in a third experiment. Ovariectomized rats were sc injected with sesame oil vehicle, 10 mg
8 bisphenol A, or 10 µg 17β-estradiol. The next day, rats were injected with 1 mg progesterone or vehicle to
9 generate 4 treatment groups: sesame oil + progesterone (n = 5), bisphenol A + sesame oil (n = 5),
10 bisphenol A + progesterone (n = 8), or estradiol + progesterone (n = 6). Examination of behavior with a
11 sexually receptive male was conducted 5–7 hours following progesterone or vehicle injection. Statistical
12 analyses included ANOVA followed by Scheffé post hoc test.

13
14 In the first experiment, injection of rats with 10 mg bisphenol A increased progesterone-positive cells in
15 both the preoptic area and ventromedial hypothalamus. The dose-response experiment demonstrated that
16 dose-related increases in progesterone-positive cells in both brain regions occurred following exposure to
17 ≥0.1 mg bisphenol A. In sexual behavior testing, treatment with bisphenol A had no effect on lordosis
18 quotient. Rejection quotient was significantly higher in rats exposed to 10 mg bisphenol A and primed
19 with 1 mg progesterone than in the vehicle control rats primed with progesterone. Treatment with 17β-
20 estradiol resulted in increased numbers of progesterone positive cells in the preoptic area and ventral
21 medial hypothalamus and increased lordosis quotient. The study authors concluded that the findings
22 suggest that bisphenol A influences sexual behavior by altering the progesterone receptor system in the
23 hypothalamus.

24
25 **Strengths/Weaknesses:** This study appears to have been relatively well conducted and demonstrated that
26 rats injected with bisphenol A exhibit 17β-estradiol-like responses. This observation is consistent with the
27 estrogenic activity of bisphenol A. However, the route of administration was sc. The number of animals
28 per group is sufficient given the nature of this study design.

29
30 **Utility (Adequacy) for CERHR Evaluation Process:** This study provides support for bisphenol A
31 induction of estrogen-like responses; however, the route of administration was sc, limiting its utility for
32 human risk assessment.

33
34 **Funabashi et al. (419)**, supported by Yokohama City University and the Japanese Ministry of Education,
35 Culture, Sports, Science, and Technology, examined the effects of bisphenol A exposure on expression of
36 progesterone receptor mRNA in brain of adult ovariectomized rats. *p*-Nonylphenol and 4-tert-octyl
37 phenol were also examined, but will not be discussed. [**No information was provided on feed, housing,**
38 **or bedding materials.**] Wistar rats were ovariectomized at 7 weeks of age, and experiments were
39 conducted 10 days following ovariectomy. In the first experiment, 6 rats/group were sc injected with
40 sesame oil vehicle or 10 mg bisphenol A (~40 mg/kg bw) [**purity not reported**]. Rats were killed 24
41 hours following injection, and frontal, parietal, and temporal cortex were removed. In a second
42 experiment, frontal, temporal, and occipital cortex were collected from rats at 0, 6, 12, or 24 hours
43 following injection with 10 mg bisphenol A; 5–6 rats were killed and examined at each time point. In
44 both experiments, progesterone receptor mRNA expression was determined by Northern Blot in each area
45 of the cortex. Data were analyzed by ANOVA followed by Fisher protected least significant difference
46 post hoc test. At 24 hours following bisphenol A exposure, expression of progesterone receptor mRNA
47 was increased in the frontal cortex and decreased in the temporal cortex. In the time-course experiments,
48 expression of progesterone receptor mRNA was increased in the frontal cortex and decreased in the
49 temporal cortex from 6 to 24 hours following exposure. Bisphenol A had no effect on expression of
50 progesterone receptor mRNA in the parietal or occipital cortex. The study authors concluded that

4.0 Reproductive Toxicity Data

1 bisphenol A can alter the neocortical function through the progesterone receptor in adult rats, but the
2 physiological significance of the effect is not known.

3
4 **Strengths/Weaknesses:** This study links relatively high single-dose (10 mg) sc bisphenol A
5 administration to the induction of progesterone receptor mRNA, an estrogenic response. A weakness is
6 the absence of a positive control to demonstrate maximal response in estrogen-mediated increases in
7 progesterone mRNA. It was also not determined if increases in mRNA were associated with increases in
8 progesterone receptor protein. There was only one dose level administered and the rats were only dosed
9 once. The sc route of dose administration bypasses potential first-pass metabolism. The authors imply that
10 since progesterone receptor mRNA in the frontal cortex is affected; bisphenol A may have a potential
11 effect on neurobehavioral endpoints. No additional studies were conducted to ascertain this potential
12 relationship.

13
14 **Utility (Adequacy) for CERHR Evaluation Process:** This study demonstrated that a single dose of
15 bisphenol A administered once to ovariectomized rats is able to increase the mRNA for progesterone
16 receptor, an estrogen-like response. However, the potential physiological and functional consequences of
17 this increase were not explored, limiting its utility.

18
19 **Della Seta et al. (420)**, supported by a grant from MURST, Italy, examined the effects of bisphenol A
20 exposure on maternal behavior in rats. [No information was provided on the type of chow, bedding,
21 and caging used.] Female Sprague Dawley rats were trained to ingest peanut oil from a micropipette. At
22 14 weeks of age, female rats were mated for 48 hours. On the day following mating, females were
23 randomly assigned to groups administered peanut oil (n=23) or 0.040 mg/kg bw/day bisphenol A (n=17)
24 through a micropipette. Dosing was continued through the gestation and lactation periods. Two days
25 following delivery, litters were culled to 4 male and 4 female pups and were cross-fostered within
26 treatment groups. Pups were weighed on days 2, 7, and 21 following birth. Maternal behavior was tested
27 at 3 and 4 days and at 8 and 9 days following delivery. In 30-minute test sessions, frequency, duration,
28 and latency of behaviors such as retrieving pups, licking pups, postures, and nest building were evaluated
29 with pups of the same sex. Behavior with pups of the opposite sex was evaluated on the second day of the
30 test period, and the order of testing with male and female pups was reversed during each testing period
31 (days 3–4 and 8–9). Data were analyzed by general linear model, Duncan multiple range test, and/or
32 Mann-Whitney *U* test. The numbers of females giving birth were 9 of 17 in the bisphenol A group and 18
33 of 23 in the control group. Nine dams in the control group and 7 in the bisphenol A group were evaluated
34 for maternal behavior. The only significant effect reported for bisphenol A was reduced duration of
35 licking-grooming pups, which occurred with both sexes of pups during both observation periods [**~25–50**
36 **% decrease as estimated from a graph**]. Effects reported to be marginally significant were decreased
37 frequencies of licking-grooming of pups ($P < 0.09$), anogenital licking of pups ($P < 0.08$), and arched
38 back posture ($P < 0.07$). The study authors concluded that maternal behavior in rats is influenced by
39 prolonged exposure to low bisphenol A doses during pregnancy and lactation.

40
41 **Strengths/Weaknesses:** This behavioral study suggested that a low, oral dose of bisphenol A (0.040
42 mg/kg bw/day) affects pregnancy and maternal behavior. However, there was an unusually low
43 pregnancy rate observed in controls (18/23), raising concerns about the study design. Bisphenol A-treated
44 rats were only administered 1 dose level, so a dose-response relationship could not be established.
45 Moreover, although the behavioral data were collected electronically, it was not stated that the analysts
46 were blinded to treatment. Because bisphenol A/peanut oil was “fed” to the mice, residual bisphenol A
47 may have been retained in the oral cavity of the dam subsequently resulting in altered grooming via
48 altered tasted perception.

49
50 **Utility (adequacy) of the Evaluation Process:** Although this study suggested a relatively low dose
51 fertility effect, as a behavioral study it was not appropriately designed to test for this effect. Because the

4.0 Reproductive Toxicity Data

1 behavioral alterations observed may have been confounded by feeding (rather than gavaging) the rats with
2 bisphenol A, and only 1 dose level of bisphenol A was assessed, this study is of limited utility.

3 4 4.2.1.2 Mouse

5 **Park et al. (421)**, support not indicated, examined the effects of bisphenol A exposure on the
6 reproductive and hematological systems of male and female mice. **[Results for females are discussed**
7 **here, and results for males are discussed in Section 4.2.2.2.]** Adult ICR mice were fed mouse
8 formulation feed (Cheil Feed). **[No information was provided about caging or bedding materials.]**
9 Fifteen mice/sex/group were ip injected with bisphenol A in an ethanol/corn oil vehicle at 0.05, 0.5, or 5.0
10 mg/kg bw on 5 occasions (every 3 days over a 14-day period). One control group received no treatment
11 and a second control group was ip injected with corn oil. Females were examined 7 days following
12 administration. Reproductive organs were weighed and fixed in Bouin solution, and histopathological
13 examination was conducted. Hematological and clinical chemistry endpoints were also assessed. Data
14 were analyzed by least significant difference test.

15
16 Exposure to bisphenol A had no effect on body weight. Significant decreases were observed for right
17 ovary weight in the mid- and high-dose group and left ovary weight in the mid-dose group **[25–27%**
18 **lower]**. No treatment effects were observed for uterine or ovarian histology. There were no effects of
19 bisphenol A treatment on hematological endpoints in females. Blood urea nitrogen levels were
20 significantly decreased **[by 28–32%]** in females of all dose groups. The study authors did not report
21 conclusions regarding study findings.

22
23 **Strengths/Weaknesses:** The study design (frequency and route of administration) limits the relevance
24 bisphenol A for human risk assessment.

25
26 **Utility (Adequacy) for CERHR Evaluation Process:** This study is not useful in the evaluation.

27
28 **Al-Hiyasat et al. (422)**, supported by Jordan University of Science and Technology, examined the effect
29 of bisphenol A and dental composite leachate on fertility of female mice. In this study, Swiss mice were
30 fed a standard laboratory feed containing soy protein. **[No information was provided on caging and**
31 **bedding materials.]** At 60 days of age, 11 mice/group were gavaged with distilled water or composite
32 leachate for 28 days. Components of the composite leachate were identified by HPLC and included tri-
33 (ethylene glycol)-dimethacrylate (5945 mg/L), bisphenol A glycerolate dimethacrylate (2097 mg/L), and
34 bisphenol A (78 mg/L). **[Based on the reported volume of administration of 0.2 mL and a body**
35 **weight of 34.4 g, CERHR estimated bisphenol A intake from leachate at 0.45 mg/kg bw/day.]**
36 Additional 60-day-old mice (n = 15/group) were gavaged with bisphenol A (97% purity), at doses of 0
37 (ethanol/distilled water vehicle), 0.005, 0.025, or 0.1 mg/kg bw/day for 28 days. Five mice/group in the
38 bisphenol A study were killed at the end of the dosing period for measurement of body, uterus, and ovary
39 weights. All mice in the leachate study and 10 mice/group in the bisphenol A study were mated to
40 untreated males (2 females to 1 male) for 10 days. One week following the end of the mating period, the
41 mice were killed and examined for pregnancy, implantations, viable fetuses, and resorptions. Body,
42 ovary, and uterus weights were measured in mice from the leachate study. Data were analyzed by Student
43 *t*-test or Fisher exact test.

44
45 Statistically significant findings are summarized in Table 102. Effects in the leachate group included
46 increased relative (to body weight) ovarian weight and decreased percentages of pregnant mice. In mice
47 exposed to bisphenol A, body weights were decreased at all dose levels. Effects observed in mice exposed
48 to the mid and high dose of bisphenol A included increased uterine weight, increased percentages of
49 resorptions/implantations, and increased percentages of mice with resorptions. Ovarian weight was
50 increased in mice of the high-dose bisphenol A group. **[Although the effects were not statistically**
51 **significant, the percentages of pregnant females were 90, 77.7, 80, and 60% pregnant mice in the**

4.0 Reproductive Toxicity Data

1 **control and each respective dose group.]** In both the composite leachate and bisphenol A groups, there
 2 were no statistically significant effects on implantations or viable fetuses. The study authors concluded
 3 that bisphenol A and components leached from dental composite have adverse effect on fertility and the
 4 reproductive system of mice.

5
 6 **Table 102. Effects in Mice Gavaged with Dental Composite Leachate or Bisphenol A**

Endpoint	Leachate	Bisphenol A, mg/kg bw/day		
		0.005	0.025	0.100
Body weight	↔	↓25%	↓25%	↓29%
Relative ovarian weight	↑43%	↔	↔	↑2.1-fold
Relative uterine weight	↔	↔	↑56%	↑63%
Percent pregnant females	↓45%	↔	↔	↔
Resorptions	↔	↔	↑11%	↑10%
Mice with resorptions	↔	↔	↑5.6-fold	↑6-fold

↑,↓ Statistically significant increase, decrease compared to controls; ↔ no statistically significant effects compared to controls.

From Al-Hiyasat et al. (422).

7
 8 **Strengths/Weaknesses:** With only 10/group, this study was underpowered for determination of potential
 9 bisphenol A-related effects on fertility. Confirmation of mating was not performed (cohabitation was for
 10 10 days; if the mice mated on day 10, the necropsy would have been performed on GD 7. It is possible
 11 that pregnancy status in some mice may have been difficult to establish, because the uteri of apparently
 12 non-pregnant mice were not stained with ammonium sulfate). Mice are not as prolific breeders as rats.
 13 Control Swiss mice normally exhibit a fertility rate ranging from 80 to 100%, and it is likely that the
 14 apparent fertility rates observed in the low- and mid-dose groups are the result of variability. Mean body
 15 weight and reproductive organ weights of bisphenol A-treated animals were only collected from 5
 16 mice/dose level. Moreover, the normal body weight range for 10-week-old female Swiss mice is 28–35 g.
 17 Given that there are only 5 mice/group, it is hard to draw any meaningful conclusions from these data.

18
 19 **Utility (Adequacy) for CERHR Evaluation Process:** Given the design, this study is of limited utility.

20
 21 **Matsumoto et al. (371)**, support not indicated, examined the effect of maternal bisphenol A exposure on
 22 growth of offspring in mice; this paper was discussed in Section 3.2.7. Because the results of this study
 23 bear on lactation competence in treated dams, the study will also be considered here. Mice were fed
 24 standard rodent chow (CE-2, Japan Clea). **[No information was provided on caging and bedding**
 25 **materials.]** Mice of the ddY strain were exposed to bisphenol A (≥97% purity) through feed at 0 or 1%
 26 from GD 14 through PND 7. The study authors stated that the bisphenol A dose was equivalent to 1000
 27 mg/kg bw/day. **[The numbers of dams treated was not indicated. Day of vaginal plug and day of**
 28 **birth were not defined.]** Mice delivered pups on PND 21. Body weight of pups were monitored during
 29 the postnatal period in 31 pups from the control group and 61–89 pups from the bisphenol A group.
 30 Serum prolactin levels were measured by RIA in 3 dams/group 4 days following delivery. Pups were
 31 killed on PND 7, and stomach weight was measured. Data were analyzed by Student *t*-test.

32
 33 No differences were reported for live pups at birth. During the postnatal period, body weights of pups in
 34 the bisphenol A group were significantly lower **[by ~40%]** than control group pups. No deaths were
 35 reported for pups in the control group, but 30% of pups in the bisphenol A group died before PND 7. On
 36 PND 1, milk could be seen in stomachs of pups from the control group, but not the bisphenol A group.
 37 **[The number of pups evaluated for milk in stomach was not reported].** On PND 7, stomach weight
 38 was significantly lower **[by 40%]** in pups from the bisphenol A compared to control group. Serum
 39 prolactin level was significantly reduced **[by 46%]** in dams from the bisphenol A group. The authors
 40 concluded that administration of a high bisphenol A dose to mice resulted in suppressed postnatal growth

4.0 Reproductive Toxicity Data

1 of offspring which probably resulted from an insufficient supply of milk, which might have been due to
2 decreased prolactin secretion.

3
4 **Strengths/Weaknesses:** This study was conducted at a single high dose that likely induced maternal
5 toxicity (which was not assessed); therefore, it is difficult to delineate if the findings in the mouse pups
6 are the result of potential bisphenol A-related effects of maternal toxicity or an effect on the pup.

7
8 **Utility (Adequacy) for CERHR Evaluation Process:** Given the likely confounding effects of maternal
9 toxicity, this study is of no utility.

10 4.2.1.3 Other mammals

11 **Nieminen et al. (423)**, support not indicated, examined the effects of bisphenol A exposure on hormone
12 levels in the European polecat (*Mustela putorius*). Five animals/group/sex [**age not reported**] were
13 administered bisphenol A [**purity not reported**] in feed at concentrations providing doses of 0, 10, 50, or
14 250 mg/kg bw/day for 2 weeks. Body weight and length were measured during the study. Animals were
15 killed at the end of the exposure period, with sampling conducted in random double-blinded order. Liver
16 and kidney were weighed. Blood samples were obtained for measurement of hormone levels by RIA.
17 Microsomal enzyme activities were determined. Statistical analyses included ANOVA, post hoc Duncan
18 test, Student *t*-test, Spearman correlation coefficient, Kolmogorov-Smirnov test, and/or Levene test.

19
20
21 There were no clinical signs of toxicity and no effects on body weight or body mass index following
22 bisphenol A exposure. Absolute and relative liver weight were significantly increased in females of the
23 high-dose group. Plasma cortisol levels were significantly reduced in females of the mid-dose group.
24 Bisphenol A exposure had no significant effects on plasma levels of testosterone, estradiol, FSH, or
25 thyroid hormones. Glutathione-S-transferase (GST) activity was significantly increased in females of the
26 high-dose group. UDPGT activity was significantly higher in females of the mid- and high-dose group
27 and males of the high dose group. There was no effect on 7-ethoxyresorufin O-deethylase (EROD)
28 activity. The study authors concluded that the endocrine effects in this study were not as remarkable as
29 the effects on liver enzymes.

30
31 **Strengths/Weaknesses:** This study provides evidence that the bisphenol A administered to polecats
32 increases GST and UDPGT activity. Since these findings were dose-related it appears that in the polecat
33 bisphenol A increases phase 2 metabolism but has minimal effects on hormone levels. Due to the limited
34 number of animals and the absence of a dose-response relationship, the hormonal changes in this study
35 are difficult to interpret.

36
37 **Utility (Adequacy) for CERHR Evaluation Process:** Due to the small sample size and absence of
38 effects on reproductive endpoints, this study is of no utility in the evaluation.

39
40 **Nieminen et al. (424)**, support not indicated, examined the effects of bisphenol A exposure on endocrine
41 endpoints in field voles (*Microtus agrestis*). Animals were housed in plastic cages with wood shavings
42 and fed R36 diet (Lactamin, Sweden). Sexually mature field voles were randomly assigned to groups that
43 received bisphenol A [**purity not reported**] in propylene glycol by sc injection for 4 days. Doses of
44 bisphenol A (numbers of females in each group) were 0 (n = 5), 10 (n = 7), 50 (n = 5), and 250 (n = 8)
45 mg/kg bw/day. Animals were killed the day following the last dose. Body and liver weights were
46 measured. Blood was drawn for measurement of sex steroids, thyroxine, and weight-regulating hormone
47 levels in plasma using RIA or immunoradiometry methods. The activities of EROD, UDPGT, and GST
48 were measured in hepatic and renal microsomes using appropriate substrates. Statistical analyses included
49 ANOVA, post hoc Duncan test, Student *t*-test, Kolmogorov-Smirnov test, Levene test, Mann-Whitney *U*
50 test, chi-squared test, and Spearman correlation. [**Results for males are discussed in Section 4.2.2.3.**]

4.0 Reproductive Toxicity Data

1 Mortality was significantly increased by bisphenol A treatment, with incidences of 18, 36, and 20% in the
2 low- to high-dose groups. No mortality was observed in the control group. Bisphenol A treatment did not
3 significantly affect body or liver weight. Plasma testosterone levels increased with dose, and statistical
4 significance was attained in high-dose females compared to control females. 17β -Estradiol levels
5 decreased with dose in females. Pooled (male + female) LH levels were not significantly altered by
6 treatment. Liver EROD activity [**apparently combined for males and females**] was significantly
7 decreased at the mid and high dose, and liver GST activities [**not clear if for males or females or both**]
8 was significantly decreased at the highest dose level. There were no other significant effects on
9 microsomal enzymes examined. The study authors concluded that wild mammals such as field voles
10 could be more susceptible to bisphenol A-induced toxicity than laboratory rodents.

11
12 **Strengths/Weaknesses:** This study provides evidence that voles are more sensitive (based on mortality)
13 to sc bisphenol A administration than rats or mice. Bisphenol A also apparently increased plasma
14 testosterone levels.

15
16 **Utility (Adequacy) for CERHR Evaluation Process:** This study suggests that voles may be more
17 susceptible to the systemic toxic effects of bisphenol A compared to common laboratory animals.
18 However, the number of voles/dose level, route of administration, and lack of similar studies in the
19 literature with this species severely limits the utility of this study for human risk assessment.

20
21 **Razzoli et al. (425)**, supported by the Ministry of University Education and Research and the University
22 of Parma, examined the effects of bisphenol A on sociosexual and exploratory behavior in female
23 Mongolian gerbils, a monogamous species. Animals were fed Mil Morini Rodent Chow (Reggio Emilia,
24 Italy) and housed in Plexiglass cages with wood shaving/cotton nesting material. At 11–12 weeks of age,
25 female gerbils were trained to drink corn oil from a syringe, and 1 week later, they were paired with a
26 male. From the 1st through the 21st day of cohabitation, 12 females/group were fed 0 (corn oil vehicle),
27 0.002, or 0.020 mg/kg bw/day bisphenol A from a syringe. A group of 12 females received ethinyl
28 estradiol, the positive control, 0.04 μ g/kg bw/day during the same time period. During the cohabitation
29 period, social behavior (e.g., agonism, social investigation, huddling, and nest sharing) was observed and
30 body weights of females were measured. A free exploratory test, which measured the amount of time
31 females spent in an area of a cage with home nesting material compared to the time spent in an unfamiliar
32 area of a cage, was conducted following the 21-day cohabitation period. Exploratory behavior was
33 evaluated by an observer blinded to treatment groups. Statistical analyses included ANOVA and Duncan
34 test for multiple comparisons.

35
36 Bisphenol A treatment did not affect body weight. Social sniffing was significantly increased [**by 60%**]
37 in the low-dose bisphenol A group. Significant effects [**percent changes compared to control**] observed
38 in the exploratory test were decreased time in the unfamiliar area at the low [**60%**] and high [**44%**] dose,
39 fewer transitions to the unfamiliar area at the low [**60%**] and high [**50%**] dose, fewer transitions to the
40 home cage at the high dose [**29%**], and less time in the unfamiliar area at the low dose [**46%**]. Similar
41 results for both social sniffing and exploratory behavior were observed in the positive control group.
42 According to the study authors, this study demonstrated that chronic exposure of adult female gerbils to
43 environmentally relevant doses of bisphenol A during the hormonally sensitive period of cohabitation
44 resulted in subtly altered social and exploratory behavior.

45
46 **Strengths/Weaknesses:** This study appears to be a well conducted (e.g., respectable sample size, given
47 study complexity; blinded analyses; positive control) and suggests that 0.002 mg/kg bw/day bisphenol A
48 shows a greater effect on behavioral endpoints than does 0.020 mg/kg bw/day. Plasma analysis of parent
49 bisphenol A and potential metabolites was not performed. In addition, for 2 of the 5 endpoints, the
50 bisphenol A responses were similar between the bisphenol A dose groups (not a strong inverted
51 response). For 1 of the 5 endpoints the high dose bisphenol A was similar to the positive control (classic

4.0 Reproductive Toxicity Data

1 dose-response). Taken together, these data at best suggest that gerbils may be sensitive to the estrogenic
2 effects of bisphenol A, .but there is no strong evidence of dose response.

3
4 **Utility (adequacy) for CERHR Evaluation Process:** This study examined behavioral endpoints in
5 gerbils, and included a positive control (17 β -estradiol) and 2 doses of bisphenol A. Significant bisphenol
6 A-induced estrogen-like behavioral alterations were observed, without a clear dose response. Given the
7 absence of concurring/supporting data from other laboratories in the same species and the difficulty in
8 extrapolating from the observed changes in behavioral endpoints in gerbils to potential effects in humans,
9 these data are of limited utility.

10 11 4.2.1.4 Invertebrates

12 **Oehlmann et al. (426)**, supported by the Berlin Federal Environmental Agency, reported the effects of
13 bisphenol A on reproductive organs in the freshwater ramshorn snail (*Marisa cornuarietis*) and the
14 marine dog whelk (*Nucella lapillus*). In the first experiment, adult ramshorn snails were exposed for 5
15 months to bisphenol A in ethanol at 0, 1, 5, 25, or 100 μ g/L. Thirty snails/group were removed every
16 month for evaluation of reproductive organs. **[The purity of bisphenol A and its stability during the
17 exposure period were not reported. The snails removed for evaluation were adults; this species
18 requires 8 months to attain sexual maturity. Octylphenol was also evaluated, but is not discussed
19 here.]** In the second experiment, ramshorn snails were exposed to bisphenol A in ethanol at 0, 1, or 100
20 μ g/L for 1 year. Thirty F₁ snails per time point were removed for evaluation at 6, 8, and 12 months. In the
21 third experiment, dog whelk were exposed to bisphenol A in glacial acetic acid at 0, 1, 25, or 100 μ g/L
22 for 3 months. Thirty specimens were removed for evaluation each month. Evaluations included
23 measurements of sex organs and the identification of sperm or oocytes in the genital tract. Statistical
24 analyses included ANCOVA followed by Tukey or Student-Newman-Keuls test, Kruskal-Wallis test, chi-
25 squared test, and Weir test.

26
27 Adult ramshorn snails were reported to show increases in volume of the capsule and albumen glands
28 (portions of the oviduct). **[Apparently, the increase in volume was based on appearance rather than
29 measurements. The measured lengths of the sex organs were not affected by treatment.]** Occasional
30 specimens that had been exposed to bisphenol A showed rupture of the oviduct with protrusion of the egg
31 mass. Enumeration of spawning masses and eggs showed statistically significant time-dependent
32 increases in all bisphenol A groups. Histologic examination of the gonads did not suggest abnormalities
33 of spermatogenesis or oogenesis. The F₁ snails also demonstrated a statistically significant increase in
34 spawning mass and oocyte production at the 100 μ g/L bisphenol A concentration, and some specimens
35 showed rupture of the oviduct at 12 months of age in both bisphenol A groups. An increase in imposex
36 **[the presence of vas deferens tissue]** was noted significantly more often in snails exposed to bisphenol
37 A 100 μ g/L than controls. Adult dog whelk demonstrated a significant increase in the length and weight
38 of the sex glands and an increase in number of females with oocytes in the oviduct. The authors
39 concluded that invertebrates are sensitive to bisphenol A toxicity at environmentally relevant
40 concentrations.

41
42 **Strengths/Weaknesses:** The study appears to be well conducted and suggests that bisphenol A has
43 stimulatory (17 β -estradiol-like) effects on the spawning masses and eggs of snails. These changes
44 occurred in the absence of a histological correlate. The potential stability/biotransformation was discussed
45 in the introduction but not determined during the exposure period.

46
47 **Utility (Adequacy) for CERHR Evaluation Process:** Because this study was conducted in a non-
48 mammalian species, it has no utility for human risk assessment.

49
50 **Forbes et al. (427)**, supported by the Bisphenol A Global Industry Group, evaluated the effects of
51 bisphenol A on reproduction in the freshwater ramshorn snail (*Marisa cornuarietis*). Bisphenol A

4.0 Reproductive Toxicity Data

1 concentrations in test water were 0, 0.10, 1.0, 16, 160, and 640 µg/L. Concentrations were periodically
2 checked. Thirty breeding pairs per treatment level were observed for a 12-week period. The number of
3 egg masses and number of eggs/egg mass were recorded. Hatchability was evaluated using 5 consecutive
4 egg masses collected from 5 females/replicate (75 egg masses/treatment). Juvenile growth rates were
5 calculated for a subset of the offspring. Nested ANOVAs were used for data analysis. All snails survived.
6 There were no significant treatment-related differences in adult egg production, hatchability, or juvenile
7 growth rate. Interindividual variability in these parameters was prominent, and the authors concluded that
8 a large number of replicates would be necessary using this animal model to detect reproductive effects.
9 The authors believed the study of Oehlmann et al. (426) to have used invalid statistical tests because there
10 was only one replicate for each treatment.

11
12 **Strengths/Weaknesses:** This study examined doses response over a 12-week exposure of freshwater
13 snails to bisphenol A with egg masses and number of eggs/egg mass as endpoints. Although no treatment-
14 related effects were observed, interindividual variability was high.

15
16 **Utility (Adequacy) for CERHR Evaluation Process:** This study of snails is of no utility in the
17 evaluation process.

18 19 4.2.1.5 *In vitro*

20 **Xu et al. (428)**, supported by the Japan Society for the Promotion of Science, examined the effects of
21 bisphenol A exposure on mouse ovarian granulosa cells in a series of experiments. Ovarian granulosa
22 cells were obtained from 4-week-old B6C3F₁ mice. Following incubation of cells with 0 or 100 fM [23
23 pg/L] to 100 µM [23 mg/L] bisphenol A in ethanol vehicle for 72 hours, the CellTiter 96 assay was used
24 to evaluate cell viability, and the TUNEL assay and 4',6-diamidino-2-phenylindole staining were used to
25 evaluate apoptosis. In cells that were incubated in 100 µM [23 mg/L] bisphenol A for 24, 48, or 72 hours,
26 the TUNEL method was used to evaluate apoptosis and a flow cytometry technique was used to assess
27 apoptosis and the cell cycle. Bcl2 and Bax protein expression was examined by Western blot, and mRNA
28 expression was assessed by RT-PCR in cells that were exposed to 100 µM [23 mg/L] bisphenol A for 72
29 hours. Experiments were repeated a minimum of 3 times. Statistical analyses included ANOVA followed
30 by Fisher protected least significant difference test. [Statistical significance was not clearly indicated
31 for some endpoints.]

32
33 A dose-related reduction in cell viability was observed at bisphenol A concentrations ≥100 pM [23 ng/L].
34 Examination of cells by the TUNEL method indicated a concentration-related increase in apoptosis at
35 bisphenol A concentrations ≥100 pM [23 ng/L]. Features noted in apoptotic cells included cellular
36 shrinkage, membrane blebbing, and nuclear condensation. Apoptotic cells, as determined by TUNEL and
37 the presence of sub-G₁ cells were increased in a time-related manner following incubation with 100 µM
38 [23 mg/L] bisphenol A from 24 to 72 hours. An increase in G₂-M arrest was also observed and reached a
39 maximum value following a 48-hour incubation of cells with 100 µM [23 mg/L] bisphenol A (18 vs. 12%
40 in controls). Expression of Bax protein was increased and Bcl2 protein was decreased following
41 incubation with 100 µM [23 mg/L] bisphenol A for 72 hours. Similar expression patterns were observed
42 for *Bax* and *Bcl2* mRNA expression [data were not shown by study authors]. The study authors
43 concluded that bisphenol A at doses of 100 pM [23 ng/L] and higher, presumably relevant to
44 environmental concentrations, decreases viability and increases apoptosis in granulosa cells. The study
45 authors postulated that apoptosis may have been induced by decreases in the anti-apoptotic protein Bcl2
46 and increases in the pro-apoptotic protein Bax.

47
48 **Strengths/Weaknesses:** Because this study used *in vitro* study PMSG-stimulated murine cells,
49 metabolism is likely to have been minimal (if present at all) and the *in vitro* dosimetry of bisphenol A is
50 difficult to extrapolate to *in vivo* dosimetry. Bisphenol A is known to induce reactive oxygen species,
51 which may influence the tetrazolium salt-based assay. Moreover, based on the data presented the

4.0 Reproductive Toxicity Data

1 mechanism by which bisphenol A may be inducing cell cytotoxicity/apoptosis is likely not “endocrine
2 disruptor” mediated.

3
4 **Utility (Adequacy) for CERHR Evaluation Process:** These data suggest that granulosa cells may be
5 sensitive to bisphenol A (induce cytotoxicity/apoptosis), but because these data were conducted in an in
6 vitro system without the appropriate information to link this information to in vivo, it is very difficult to
7 utilize this information for human risk assessment. This study is not useful in the evaluation.

8
9 **Mlynarčíková et al. (429)**, supported by the European Union, examined the effects of bisphenol A
10 exposure on hormone production by porcine ovarian granulosa cells. Granulosa cell cultures were
11 prepared from porcine ovaries collected from a slaughter house. The cells were incubated for 72 hours in
12 media containing bisphenol A at 10^{-8} to 10^{-4} M [**2.3 µg/L to 23 mg/L**] or the DMSO vehicle, with or
13 without addition of 1 µg/mL FSH or LH. Following the incubation period, media were collected for
14 measurement of progesterone and 17β -estradiol concentrations by RIA. Experiments were replicated 5–8
15 times. Data were analyzed by ANOVA and Bonferroni post test. Significant changes in progesterone
16 production, included an increase at 10^{-5} M [**2.3 mg/L**] and decrease at 10^{-4} M [**23 mg/L**] bisphenol A.
17 Bisphenol A significantly increased FSH-stimulated progesterone synthesis at 10^{-6} M [**0.23 mg/L**] and
18 inhibited FSH-stimulated progesterone production at 10^{-4} M [**23 mg/L**]. LH-induced progesterone
19 production was inhibited by 10^{-4} [**23 mg/L**] bisphenol A. FSH-induced 17β -estradiol production was also
20 inhibited by bisphenol A at all concentrations tested, but statistical significance was only attained at doses
21 $\geq 10^{-6}$ M [**0.23 mg/L**]. Bisphenol A dimethylacrylate was also tested, and most results were similar to
22 those observed with bisphenol A. The study authors concluded that ovarian steroidogenesis might be a
23 target of bisphenol A toxicity.

24
25 **Strengths/Weaknesses:** Potential estrogenic effects were observed at 10^{-5} M bisphenol A. Decreases in
26 responses observed at the 10^{-4} M concentration are likely due to nonspecific cytotoxicity. Bisphenol A-
27 mediated responses in progesterone endpoints appeared to reach a near maximum at the lowest dose level
28 examined. There was no mention of whether phenol red-free media were used or whether fetal bovine
29 serum was charcoal-stripped. The serum likely contained steroids, which would have been potential
30 confounding factors. Also, it appears that cell viability was not examined after the incubation period.
31 With exception of the highest dose level, there was no dose response (inconsistent trends); the statistical
32 flags are potentially due to random chance. Since this was an in vitro study, the potential effects of
33 metabolism could not be assessed.

34
35 **Utility (Adequacy) for CERHR Evaluation Process:** Due the weaknesses and limitation in the
36 experimental design, this study is not useful in the evaluation.

37
38 **Mohri and Yoshida (430)**, supported by the Japanese Ministry of Education, Science, Sports, and
39 Culture, examined the effects of bisphenol A and 17β -estradiol exposure on calcium oscillations in
40 immature mouse oocytes. Immature oocytes with intact germinal vesicles were obtained from 8–12-week-
41 old CD-1/ICR mice and incubated in bisphenol A in a DMSO vehicle at concentrations of 0 or 10 nM
42 [**2.3 µg/L**] to 100 µM [**23 mg/L**] for 60 minutes. Calcium oscillations were measured using a Fura-2 dye
43 and image analyzer. Data were analyzed by Student *t*-test. At 100 µM [**23 mg/L**] bisphenol A, the
44 duration of calcium oscillations was significantly shortened and the oscillations became irregular. The
45 same findings were observed following exposure to 17β -estradiol at concentrations that were 10,000-fold
46 lower than that of bisphenol A, producing the same effect. The study authors stated that estrogens may
47 affect the oocyte by regulating calcium oscillations and that bisphenol A could affect oocyte maturation.

48
49 **Strengths/Weaknesses:** This study appears to have been well conducted; however, because this study
50 used an in vitro system, metabolism could not be assessed. It is unclear if calcium oscillations play a role
51 in oocyte maturation in other species, including humans.

4.0 Reproductive Toxicity Data

1
2 **Utility (Adequacy) for CERHR Evaluation Process:** Calcium oscillations in immature mouse oocytes
3 do not appear to be a particularly sensitive target of bisphenol A; therefore, this study is not useful in the
4 evaluation.

5 6 *4.2.2 Male*

7 Studies on the androgenicity of bisphenol A, including Hershberger assays, are discussed in Section 2.2.3.

8 9 *4.2.2.1 Rat*

10 **Yamasaki et al. (128)**, support not indicated, conducted a 28-day exposure study that provided some
11 information on the reproductive organs of male and female rats. **[Complete details of this study are**
12 **included in Section 2. Results for males are discussed in this section, and results for females are**
13 **discussed in Section 4.2.1.1.]** CD rats were fed a commercial diet (MF Oriental Yeast Co.) and housed in
14 stainless steel wire mesh cages. Ten 7-week-old rats/sex/group were gavaged with bisphenol A at 0 (olive
15 oil vehicle), 40, 200, or 1000 mg/kg bw/day for 28 days. Due to the death of 1 animal exhibiting clinical
16 signs in the 1000 mg/kg bw/day group, the high dose was reduced to 600 mg/kg bw/day on the eighth day
17 of the study. In an additional study, rats were exposed to ethinyl estradiol at 0, 10, 50, or 200 µg/kg
18 bw/day for 28 days. There were no treatment-related abnormalities in sperm or alterations in blood levels
19 of thyroid hormones, FSH, LH, 17β-estradiol, prolactin, or testosterone. Changes in relative reproductive
20 organ weights **[assumed to be relative to body weight]** included a 28% decrease in relative ventral
21 prostate weight and 21% increase in relative testis weight in the high-dose group. No gross or
22 histopathological alterations were reported for reproductive organs. The study authors concluded that
23 change in estrous cyclicity was the only useful endpoint for evaluating the endocrine-mediated effects of
24 bisphenol A. In comparison, male rats exposed to the mid and/or high doses of ethinyl estradiol
25 experienced decreased prostate, seminal vesicle, and pituitary weights; increased testis weight; and
26 histopathological alterations in prostate, seminal vesicle, mammary gland, and testis.

27
28 **Strengths/Weaknesses:** This was a well conducted, comprehensive study. Dose levels were appropriate
29 for the induction of some measure of toxicity (decrease in body weight). Bisphenol A administration
30 resulted in minimal effects on male reproductive normalized organ weights. There was an apparent
31 statistically significant increase in testicular weight; however, testicular (as well as other reproductive
32 organ) weight of the same age animals has been shown to be independent of body weight.

33
34 **Utility (Adequacy) for CERHR Evaluation Process:** This was a well conducted GLP-study with an
35 appropriate route of administration. The apparent increase in testis weight after bisphenol A
36 administration is likely an artifact of decreased terminal body weight. There appears to be a bisphenol A-
37 related decrease in the weight of the ventral prostate at the high dose level. The effects on the adrenal are
38 likely secondary to stress/general toxicity, which is common in rats. The NOAEL for male reproductive
39 effects (decrease ventral prostate weight) is 200 mg/kg bw/day.

40
41 **Takahashi and Oishi (431)**, support not indicated, examined the effects of bisphenol A exposure on
42 testis of rats. F344 rats were fed standard, soy-containing diet (CE-2, Clea Japan, Inc. Tokyo) and housed
43 in stainless steel suspended cages. Four-week-old male rats (n = 8/group) were administered bisphenol A
44 (99.0% purity) through diet at concentrations of 0, 0.25, 0.5, or 1.0% for 44 days. The study authors
45 estimated bisphenol A intake at 235, 466, and 950 mg/kg bw/day. The stability of bisphenol A in the diet
46 was verified. Food intake was measured, and animals were weighed and observed daily for clinical signs.
47 Rats were killed when mean body weight of controls reached ~200 g. Testosterone levels were measured
48 in serum using an ELISA method. Preputial gland, testes, epididymides, prostate, seminal vesicles,
49 kidneys, and liver were weighed. The testis was fixed in buffered 6% formaldehyde and examined
50 histologically. Statistical analyses included Bartlett test, ANOVA, Dunnet or Scheffé parametric test,

4.0 Reproductive Toxicity Data

1 Kruskal-Wallis test, Dunnet non-parametric test, Wilcoxon rank sum test, chi-squared test, Mantel-Haenzel test, and Fisher exact test.

2
3
4 Statistically significant findings are summarized in Table 103. Body weight gain and terminal body
5 weights were reduced in males of the mid- and high-dose groups. Food intake was said to be slightly
6 decreased according to dose. Absolute organ weight effects included decreased weight of preputial glands
7 at all doses; liver in the mid and high dose group; and seminal vesicles with coagulation glands, dorsal
8 and lateral prostate, and hypophysis at the high dose. **[The Expert Panel assumes that by coagulation
9 gland, the authors mean the anterior prostate or coagulating gland.]** Significant organ weight weight
10 effects relative to body weights are summarized in Table 103. Changes in relative organ weights included
11 decreased preputial gland weight and increased kidney weights at all doses, decreased liver weight at the
12 mid and high dose, and decreased dorsal and lateral prostate weight at the high dose. Testicular lesions
13 observed with bisphenol A treatment included seminiferous tubule degeneration at the mid and high dose,
14 disorganized spermatids at all dose levels, and differences in percentages of seminiferous tubules in
15 spermatogenic stages at all dose levels. Although it does not appear that statistical significance was
16 attained, dose-related increases in arrested spermatogenesis and disappearance of elongated spermatids
17 were also reported. There were no significant effects on serum testosterone concentrations. The study
18 authors concluded that bisphenol A was toxic to the testis and accessory sex organs of F344 rats at a
19 minimum toxic dose of 235 mg/kg bw/day.

20
21 **Table 103. Effects Observed in Male Rats Exposed to Bisphenol A Through Diet**

Endpoint	Dose, % in diet [mg/kg bw/day]			BMD ₁₀	BMDL ₁₀	BMD _{1SD}	BMDL _{1SD}
	0.25	0.5	1.0				
Terminal body weight	↔	↓13%	↓18%	0.55 [522]	0.42 [399]	0.41 [389]	0.30 [285]
Relative weight							
Dorsal and lateral prostate	↔	↔	↓32%	0.29 [276]	0.22 [209]	0.52 [494]	0.36 [342]
Preputial gland ^a	↓22%	↓26%	↓25%	0.13 [124]	0.09 [86]	0.18 [171]	0.12 [114]
Liver	↔	↓10%	↓14%	0.69 [656]	0.56 [532]	0.30 [285]	0.23 [218]
Kidney	↑8%	↑8%	↑12%	0.99 [940]	0.69 [656]	0.50 [475]	0.34 [323]
No. rats with							
Seminiferous tubule degeneration ^b	↔	↑ to 6/8	↑ to 5/8				
Disorganization of stage I-VI spermatids (+ severity) ^b	↑ to 4 of 8	↔	↔				
Disorganization of stage I-VI spermatids (2+ severity) ^b	↔	↔	↑ to 6 of 8	0.36 [342]	0.22 [209]		
% Seminiferous tubules in stages							
I-VI	↓59%	↓70%	↓53%				
IX-XI	↑3.4-fold	↑5.2-fold	↑4-fold				
XII-XIV	↑3.2-fold	↑3.6-fold	↑3-fold				

↑,↓ Statistically significant increase, decrease compared to controls; ↔no statistically significant effect compared to controls.

^aBenchmark doses were estimated using a polynomial model.

^bControl value = 0 of 8.

From Takahashi and Oishi (431).

22
23 **Strengths/Weaknesses:** This paper reports a relatively well conducted study with a relevant route of
24 administration. General toxicity was demonstrated. Formalin is not recommended for fixation of testes
25 with paraffin embedding, especially when staging is conducted.

26
27 **Utility (Adequacy) for CERHR Evaluation Process:** Findings suggest a hormonal effect on hormone-
28 dependent reproductive tissues at all doses examined. The lowest dose level, 0.25% in diet, exhibited

4.0 Reproductive Toxicity Data

1 histopathological changes in the testes, most strikingly described as a large alteration in the relative
2 frequency of the different stages of the seminiferous epithelium. Due to techniques used for fixation and
3 embedding of the testes, the histopathological analyses may be of limited value. Overall, this study used
4 an appropriate route of exposure and indicates the induction of male reproductive tract changes at
5 bisphenol A exposures of 235 mg/kg bw/day and above.

6
7 **Sakaue et al. (432)**, supported by the Japanese Science and Technology Agency, examined the effect of
8 bisphenol A exposure on spermatogenesis in the adult rat. Animals were fed CE-2 chow (CLEA Japan)
9 and housed in stainless steel wire caging. Thirteen-week old male Sprague Dawley rats (5/group) were
10 gavaged for 6 days with the ethanol/corn oil vehicle or bisphenol A (99.6% purity) at doses 0.020, 0.200,
11 2, 20, or 200 mg/kg bw/day. The high dose was based upon a preliminary experiment that demonstrated
12 reduced daily sperm production in a Holtzman rat gavaged with 200 mg/kg bw/day bisphenol A for 6
13 days. In this study, rats were killed 2 days following dosing (at 14 weeks of age) or at 18 weeks of age.
14 Testes were weighed. Sperm endpoints were measured from one testis. Histopathological examinations
15 were conducted on the other testis after fixation in Bouin fluid, paraffin embedding, and staining with
16 hematoxylin and eosin. Statistical analyses included Student *t*-test, ANOVA, and Fisher protected least
17 significant difference test.

18
19 There were no changes in daily sperm production/g testis at 14 compared to 18 weeks of age. **[No data
20 were shown for 14-week-old rats, and results of bisphenol A treatment were not discussed.]**

21 Bisphenol A did not significantly affect body or testis weight at 18 weeks of age. In the 18-week-old rats,
22 daily sperm production and daily sperm production/g tissue were significantly reduced **[by ~25%]** in all
23 bisphenol A treatment groups. The study authors noted the lack of a dose-response relationship and that
24 daily sperm production in treated groups at 18 weeks of age was comparable to that of 14-week-old
25 controls. Histopathological evaluations of testis revealed no evidence of atrophy or disrupted
26 spermatogenesis in the seminiferous tubules. **[Data were not shown by study authors.]**

27
28 To obtain more dose-response information, Sakaue et al. (432) repeated the study in 8 rats/group dosed
29 **[assumed by gavage as in the first study]** with 0.000002, 0.00002, 0.0002, 0.002, 0.020, 0.200, or 2
30 mg/kg bw/day bisphenol A. **[It is assumed that ages of rats, treatment period, and observation
31 periods were the same as in the first study.]** Body and testis weights were not affected by bisphenol A
32 treatment at week 18. At week 18, significant decreases in daily sperm production and daily sperm
33 production/g tissue were observed at 0.020, 0.200, and 2 mg/kg bw/day. **[The decrease compared to
34 control was estimated from a graph. For daily sperm production, the decreases were ~30% at 0.020
35 mg/kg bw/day, ~34% at 0.200 mg/kg bw/day, and ~32% at 2 mg/kg bw/day. For daily sperm
36 production/g tissue, the decreases were ~24% at 0.020 mg/kg bw/day, ~32% at 0.200 mg/kg bw/day,
37 and ~28% at 2 mg/kg bw/day.]**

38
39 In a third experiment, rats were given a single oral dose of 0.020 mg/kg bw bisphenol A. Six hours later,
40 the rats were killed, the right testis was homogenized, and the cytosol was examined for protein
41 expression using two-dimensional polyacrylamide gel electrophoresis. Changes in intensity and mobility
42 were noted for 3 unidentified proteins. The study authors concluded that the dose-response curve for
43 bisphenol A affects on spermatogenesis in the adult rat was monotonic rather than having an inverted U-
44 shape.

45
46 **Strengths/Weaknesses:** This study used a relevant route of administration and may have shown a
47 potential estrogenic effect. Variability in control daily sperm production between the first and second
48 study is disturbing; given the small sample (5/group), this variability severely decreases confidence in the
49 data. No histopathologic correlate was presented. The apparent decrease in daily sperm production is
50 unlikely to affect fertility (the reproductive consequence were not determined); therefore, the NOAEL
51 cannot be established. Confidence in the control values is limited.

4.0 Reproductive Toxicity Data

1
2 **Utility (adequacy) for CERHR Evaluation Process:** This study is suggestive of a reproductive tract
3 effect of bisphenol A, but design considerations (small sample, variable control values between
4 experiments) limit the utility of the study.
5

6 **Ashby et al. (433)**, support not indicated, examined the effects of bisphenol A exposure on sperm
7 production in rats. The study attempted to replicate earlier findings from Sakaue et al. (432). Five
8 independent experiments were conducted, and the conditions for each experiment are summarized in
9 Table 104. Some of the experiments used the same conditions as the Sakaue et al. (432) study, including
10 stainless steel cages with no bedding, CE2 diet (CLEA, Tokyo, Japan), and glass water bottles. In the first
11 4 studies, 10–20 adult (~13-week-old) Sprague Dawley rats/group were gavaged with bisphenol A (99%
12 purity) at 0 (ethanol/corn oil vehicle), 0.020, 2, or 200 mg/kg bw/day for 6 days. Concentrations of dosing
13 solutions were verified. In the fifth study, rats fed different diets were gavaged with vehicle for 6 days.
14 Rats were fed 1 of 3 diets as indicated in Table 104. Phytoestrogen aglycone content of the feed was
15 measured. Respective concentrations of daidzein, genistein, and coumestrol in each feed were reported at
16 94, 62, and 0.6 µg/g diet for Rat and Mouse No. 3 (RM3; Special Diet Services Ltd.); 40, 23, and 0.1 µg/g
17 diet for 5002 (Purina Mills); and 157, 106, and 2.2 µg/g CE2 diet. Ten rats were used in each group,
18 except in third and fourth studies, where 20 control rats were split into 2 groups prior to dosing. Rats were
19 administered drinking water through an automatic system in the first study and via glass bottles in the
20 other studies. In the first study, rats were housed 3/cage at the start of the study and 2/cage later in the
21 study. In the other 4 studies, rats were housed 2/cage. Rats were weighed during the study. Animals were
22 killed at 18 weeks of age, 5 weeks after the start of dosing. Liver, kidney, and reproductive organs were
23 weighed, and sperm counts were obtained. In the first 4 studies, data were analyzed by ANOVA,
24 ANCOVA for organ and body weights, and Dunnett test. Results from all 4 studies were also analyzed by
25 ANOVA in an attempt to increase study power. Data from the fifth study were analyzed by Fisher least
26 significant difference test.
27

28 In the four studies that compared the effects of bisphenol A exposure to a vehicle control group, there
29 were no significant effects of bisphenol A exposure on sperm count, daily sperm production, or weights
30 of body, liver, kidney, testis, prostate, epididymis, or seminal vesicle. One animal exposed to 200 mg/kg
31 bw/day bisphenol A in the third study was reported to have unexpectedly small testes and epididymides,
32 but the study authors indicated that inclusion of this animal in later statistical analyses had no effect on
33 outcome. One animal in the 200 mg/kg bw/day group in the fourth study had a small testis. No significant
34 effects were observed when data from the first 4 experiments were pooled and analyzed. The study
35 authors noted that some endpoints were variable from one experiment to the other. It was noted that
36 prostate weights were 10% lower in animals from Experiment 1 than from Experiments 2–4. Sperm
37 counts and daily sperm production were reportedly different in control animals from Experiment 1
38 compared to Experiment 2. It was noted that rats were fed different diets in Experiment 1 (RM3) and
39 Experiment 2 (5002), and a study to examine the effects of feed was conducted. In the study examining
40 effects in rats fed different diets but exposed to vehicle, no effects of diet on daily sperm production were
41 observed. The only significant effect reported was a 9% lower weight of right epididymis in rats fed CE2
42 compared to RM3 or 5002 feed. The study authors stated that the effect was likely spurious due to lack of
43 effect on other endpoints, no effect on left or total epididymis weight, and lack of the effect in the first 4
44 experiments. The study authors concluded that there was no evidence in their study that bisphenol A
45 affected reproductive organ weights or daily sperm production. Lack of bisphenol A-induced effect on
46 daily sperm production was in contrast to observations of the Sakaue et al. (432) study, which reported a
47 decrease in this endpoint. Subtle genetic differences in the rats were suggested as a possible reason for
48 differences in results between the 2 studies.

4.0 Reproductive Toxicity Data

Table 104. Conditions Used in Experiments to Study Bisphenol A Effects on Sperm Production in Rats

Experiment	Bisphenol A doses, mg/kg bw/day	No. rats/group	Diet/water	Caging
1	0, 0.020, 2, or 200	10	RM3/Automatic system	Stainless steel, unspecified bedding
2	0, 0.020, 2, or 200	10	5002/Glass bottles	Stainless steel, no bedding
3	0, 0.020, 2, or 200	10/bisphenol A group; 20 in control group	5002/ Glass bottles	Stainless steel, no bedding
4	0, 0.020, 2, or 200	10/bisphenol A group; 20 in control group	CE2/ Glass bottles	Stainless steel, no bedding
5	0	10	RM3, 5002 or CE2/not specified	Not specified

From Ashby et al. (433).

Strengths/Weaknesses: This paper reports a very well conducted, comprehensive assessment of the potential effects of bisphenol A on daily sperm production. Absence of confirmation of the work of Sakaue et al. (432) led to an extensive study of the potential variables (e.g. diet, housing, etc.) that might account for the discrepancies. These data suggests that subtle changes in study endpoints, especially daily sperm production and organ weights, may occur by random chance or genetic differences in the respective lab's supplier of rats may play a role. These data also strongly suggest bisphenol A administered orally has no effect on sperm production.

Utility (Adequacy) for CERHR Evaluation Process: Given the robustness and comprehensiveness of this study, it is highly useful. It strongly suggests that the NOAEL for potential bisphenol A-mediated effects on the adult rat reproductive system exceeds 200 mg/kg/day.

Tohei et al. (434), supported in part by the Japan Society for the Promotion of Science, examined the effects of bisphenol A exposure on testicular function of Wistar-Imamichi rats. **[No information was provided about composition of chow, bedding, or caging.]** In a series of studies, rats were dosed with bisphenol A in sesame oil by sc injection for 2 weeks. Bisphenol A doses were 0.1 or 1 mg/day [**~0.3 or 3 mg/kg bw/day based on the reported body weights of 300–350 g**]. The dose of 1 mg/day bisphenol A was stated to be similar to the highest exposures reported in humans, which were based on saliva levels measured in patients receiving composite dental sealants. Doses and exposure duration were based on results of preliminary studies. Five or 6 animals/dose group were used in each experiment. Statistical analyses included ANOVA, Fisher protected least significant difference test, and Mann-Whitney *U* test.

In the first study conducted to examine testicular and pituitary function, LH, FSH, prolactin, testosterone, and inhibin were measured in plasma, pituitary, and/or testis by RIA in rats sc dosed with 1 mg/day bisphenol A for 2 weeks. Statistically significant effects **[percent differences compared to controls, as estimated from a graph]** included increases in plasma levels of LH **[150%]** and prolactin **[1067%]** and decreases in levels of plasma testosterone **[29%]** and testicular inhibin **[36%]**.

In a second experiment to examine testicular response, rats were sc dosed with 0.1 or 1 mg/day bisphenol A for 2 weeks. The rats then received 10 IU hCG through an atrial cannula. Blood samples were drawn

4.0 Reproductive Toxicity Data

1 for measurement of progesterone and testosterone levels before and at various time intervals between 30
2 and 180 minutes following the hCG challenge. Plasma progesterone and testosterone levels were increased
3 following the hCG challenge in control rats. In the bisphenol A-treated rats, only a slight increase in
4 progesterone levels occurred 30 minutes following challenge, and plasma progesterone levels were
5 significantly lower compared to the control group at 60–150 minutes following challenge. There was an
6 increase in plasma testosterone level following challenge of the bisphenol A group, but values were
7 significantly lower than control values at 90–120 minutes following the challenge.

8
9 In a third experiment examining pituitary response, adult male rats were castrated 5 days before bisphenol
10 A treatment. Castrated rats were sc injected with 1 mg/day bisphenol A and 75 µg/day testosterone
11 propionate for 2 weeks. The rats then received 250 ng gonadotropin-releasing hormone by sc injection.
12 Plasma LH was measured before and at various time intervals between 0.25 and 4 hours following the
13 gonadotropin-releasing hormone challenge. No statistically significant effects were observed.

14
15 In a fourth study, males were dosed with 1 mg/day bisphenol A for 2 weeks and then paired with females
16 in proestrus. Sexual function was evaluated by scoring mounts, intromissions, and ejaculations. No
17 significant effects were observed for sexual function. Based on the findings reported in all studies, the
18 study authors concluded that “The testis is probably a more sensitive site for [bisphenol A] action than the
19 hypothalamus-pituitary axis.”

20
21 **Strengths/Weaknesses:** RIAs appear to have been competently conducted. SC is not a relevant route of
22 exposure, and the sample size was limited. Blood collection via decapitation is not appropriate, because
23 decapitation stress affects plasma prolactin and LH secretion. No mention is made of the order of killing.
24 If controls were killed first and the guillotine was not cleaned between uses (and animals were not in
25 separate rooms), there may be serious confounding of the data. Because rat plasma testosterone levels are
26 normally highly variable, the low degree of variability in this study, given the small sample size, is
27 remarkable (~ ±0.12 ng/mL). No functional consequence of the alterations in hormone levels were
28 described.

29
30 **Utility (adequacy) for CERHR Evaluation Process:** Due to the study design these data are of minimal
31 utility.

32
33 **Kim et al. (I25)**, supported by the Korean Ministry of Health and Social Welfare, examined the effects of
34 bisphenol A exposure on the male reproductive system. A translation of the study was provided by the
35 American Plastics Council. Four-week-old male F344 rats were given bisphenol A in drinking water at 0
36 (ethanol vehicle), 0.1, 1, 10, or 100 ppm for 13 weeks. According to the study authors, these values were
37 equivalent to 0.011, 0.116, 1.094, and 11.846 mg/kg bw/day. **[No information was provided about the**
38 **numbers of animals treated/group, bisphenol A purity, or feed, caging, or bedding materials.]** Body
39 weight and food and water consumption were measured during the study. Urine was collected for 24
40 hours following completion of dosing, and then animals were killed. Blood was collected. Organs,
41 including those of the male reproductive system, were weighed. Parts of organs were preserved in
42 formalin and examined histologically. Testes and epididymides were preserved in liquid nitrogen to
43 obtain sperm counts and for measurement of levels of testicular enzymes. Data were analyzed by
44 ANOVA.

45
46 Bisphenol A treatment had no significant effect on body weight or food or water intake. There were no
47 effects on absolute or relative weights of the testis, epididymis, prostate, seminal vesicle, liver, kidney,
48 heart, lung, spleen, or brain. Daily sperm production and number of sperm heads were unaffected by
49 bisphenol A treatment. No significant effects were observed for activities of testicular γ -glutamyl
50 transpeptidase, sorbitol dehydrogenase, acid phosphatase, or β -glucuronidase. No histopathological
51 alterations were reported for the testis, epididymis, seminal vesicle, prostate, spleen, or brain. Bisphenol

4.0 Reproductive Toxicity Data

1 A levels in urine are reported in Section 2. The study authors concluded that sperm density and the male
2 reproductive system do not appear to be affected in F344 rats exposed to bisphenol A.

3
4 **Strengths/Weaknesses:** The absence of information about the animal group size and other study design
5 features is a weakness

6
7 **Utility (Adequacy) for CERHR Evaluation Process:** This study is not useful in the evaluation.

8
9 **Chitra et al. (435)**, supported by the Lady Tata Memorial Trust, Indian Council of Medical Research, and
10 the Population Council, examined the effects of bisphenol A on the reproductive system of male rats.
11 Animals were given “standard commercial laboratory chow.” **[Bedding and caging materials were not**
12 **reported.]** Six 45-day-old male Wistar rats/group were orally dosed **[gavage assumed]** with bisphenol A
13 (97% purity) in olive oil at 0, 0.0002, 0.002, and 0.020 mg/kg bw/day for 45 days. Rats were killed 24
14 hours following the last treatment. Testes, epididymides, seminal vesicles, and ventral prostate were
15 weighed. Epididymal sperm counts and motility were assessed. Antioxidant enzyme activities were
16 measured in sperm. Statistical analyses included ANOVA followed by Student *t*-test. Significant effects
17 on organ weights and sperm endpoints are summarized in Table 105. Bisphenol A treatment did not affect
18 body weight. Absolute and relative (to body weight) weights of testis and epididymis and were reduced,
19 and absolute and relative ventral prostate weights were increased at all dose levels. Effects on relative
20 organ weights are summarized in Table 105. Sperm motility was decreased at all dose levels, and sperm
21 counts were reduced at the mid and high dose. There were dose-related decreases in activity of superoxide
22 dismutase, catalase, glutathione reductase, and glutathione peroxidase in sperm at all dose levels.
23 Hydrogen peroxide generation and lipid peroxidation in sperm increased dose-dependently at all dose
24 levels. The study authors concluded that adverse effects of bisphenol A on the male reproductive system
25 may be due to oxidative stress.

26
27 **Table 105. Reproductive Effects in Male Rats Orally Dosed with Bisphenol A**

Endpoint	Dose, mg/kg bw/day						
	0.0002	0.002	0.020	BMD ₁₀	BMDL ₁₀	BMD _{1SD}	BMDL _{1SD}
Relative organ weight							
Testis	↓5%	↓6%	↓7%	0.056	0.021	0.014	0.0087
Epididymis	↓13%	↓17%	↓26%	0.011	0.0082	0.0069	0.0050
Ventral prostate	↑13%	↑34%	↑29%	0.014	0.0083	0.015	0.0089
Epididymal sperm motility ^a	↓23%	↓37%	↓41%				
Epididymal sperm count	↔	↓18%	↓27%				

↑,↓ Statistically significant increase, decrease; ↔ no statistically significant effect.

^aValues estimated from a graph by CERHR; data estimated from graphs were not modeled.

From Chitra et al. (435).

28
29 **Strengths/Weaknesses:** Although these studies have a limited number of animals per group, they appear
30 to be relatively well conducted, and there are apparently consistent dose-dependent changes in testis and
31 epididymis weights and sperm parameters. The epididymal (portion not mentioned) sperm numbers
32 measured in this study are consistent with the daily sperm production measured by Sakaue et al. (432). A
33 potential significant concern in this study is the use of olive oil as the vehicle. The stability/reactivity of
34 bisphenol A was not determined and it is possible that bisphenol A interacted with olive oil, resulting in
35 the observed findings.

36
37 **Utility (adequacy) of CERHR Evaluation Process:** This study provides suggestive data that bisphenol
38 A induces oxidative stress in epididymal sperm and alters testis and epididymis weights at low doses.
39 However, concern with the small group size and the vehicle used for dosing limits utility of the study.

4.0 Reproductive Toxicity Data

1 **Chitra et al. (436)**, supported by the Population Council, New York, examined the effects of bisphenol A
2 and vitamin C exposure on epididymis and sperm counts in rats. Wistar rats (45-days old) were fed
3 standard commercial laboratory chow and housed in plastic cages. **[No information was provided about**
4 **bedding.]** Four rats/group were orally dosed with bisphenol A (97% purity) at 0 (olive oil vehicle),
5 0.0002, 0.002, or 0.020 mg/kg bw/day for 60 days. Additional rats received the same bisphenol A doses
6 in conjunction with 40 mg vitamin C. **[The specific method of oral dosing was not stated. A vehicle**
7 **control group administered vitamin C was not included.]** Rats were killed 24 hours following the last
8 dose. Epididymides were fixed in Bouin solution and examined histologically. Sperm were counted and
9 examined for viability and motility. Levels of antioxidant enzymes were measured in sperm and
10 epididymis. Data were analyzed by ANOVA followed by Student *t*-test.

11
12 There was no effect on sperm viability, but significant dose-related reductions were observed in sperm
13 motility and count in all dose groups. **[In the low- to high-dose group, sperm motility was reduced to**
14 **~70, 60, and 55% of control levels. Sperm counts in the low to high dose group were ~12, 30, and**
15 **40% lower than control values.]** Complete degeneration of epithelia of caput, corpus, and cauda
16 epididymis was reported at all dose levels. **[It was not clear if the effect occurred in every rat of each**
17 **dose group.]** Significant dose-related decreases in glutathione peroxidase and superoxide dismutase
18 activity and increased lipid peroxidation were observed in sperm and epididymis of rats from each
19 bisphenol A treatment group. No changes in sperm motility, sperm count, antioxidant enzyme activity, or
20 lipid peroxidation were observed when bisphenol A was administered with vitamin C. The study authors
21 concluded that bisphenol A induced oxidative stress and degeneration of epididymal epithelium, and
22 vitamin C protected against those effects.

23
24 **Strengths/Weaknesses:** There were a limited number of animals/group, and there is low confidence in
25 the control values (e.g. ~95% motile sperm when literature values range from 60–85%; minimal
26 variability in this study is surprising). The bisphenol A-treated animals exhibited an apparent decrease in
27 motile sperm; however, these levels are consistent with those that are published in the literature. These
28 values may be the result of strain/substrain differences. The stability of the bisphenol A/olive oil mixture
29 was not assessed, and there is potential for interactions/potential of reactive oxygen species. It appears
30 the bisphenol A data in this study are the same data noted in the previous study (this report was quoted in
31 the previous manuscript).

32
33 **Utility (Adequacy) for CERHR Evaluation Process:** This data set may be the same as that above and
34 suggests that bisphenol A induces oxidative stress. However, concerns with group size and the vehicle
35 limits the utility of the study.

36
37 **Saito et al. (437)**, support not indicated, examined the effects of bisphenol A exposure on sex hormone
38 levels in male rats. Wistar rats were fed MF feed (Oriental Yeast Co.). **[No information was provided**
39 **about bedding and caging materials.]** Eight or 9 rats/group were sc injected with bisphenol A **[purity**
40 **not reported]** at 0 (corn oil vehicle), 0.005, or 5 mg every 2 days from 3 to 11 weeks of age. **[Based on a**
41 **graph showing body weights of ~50 g at the beginning of treatment and ~300 g at the end of**
42 **treatment, the bisphenol A doses would have been 0.1 and 100 mg/kg bw at the beginning of the**
43 **treatment period and 0.017 and 17 mg/kg bw at the end of the treatment period.]** Additional groups
44 of 8–9 rats were injected with 5 µg/day 17β-estradiol or diethylstilbestrol. Rats were killed at 13 weeks of
45 age, 2 weeks following the last treatment. Body, testes, and other reproductive organs were weighed.
46 Levels of 17β-estradiol and testosterone were measured in plasma by RIA. Data were analyzed by
47 Student *t*-test and Dunnett test. No clinical signs of toxicity or changes in behavior were observed.
48 Exposure to bisphenol A did not affect body weight gain or absolute or relative testis weight. No effects
49 were observed for weights of prostate, preputial gland, or epididymis. **[Data were not shown by study**
50 **authors.]** Plasma testosterone levels were significantly reduced in the low bisphenol A group **[by ~1.5**
51 **fold]** and plasma estradiol levels were significantly increased in the high bisphenol A dose group **[by ~8-**

4.0 Reproductive Toxicity Data

1 **fold].** Effects observed with 17 β -estradiol and diethylstilbestrol exposure included decreased body weight
2 gain, reduced absolute and relative testis weight, decreased plasma testosterone levels, and increased
3 plasma 17 β -estradiol levels. The study authors concluded that bisphenol A disturbed sex steroid
4 production in male rats.

5
6 **Strengths/Weaknesses:** The sc route of exposure was used and is not relevant to human exposure. The
7 method of anesthesia was inappropriate for measuring hormone levels. Single point testosterone
8 measurements are normally highly variable; the apparent significant decrease in testosterone observed in
9 this study may be spurious and due to the small group size, an unusual low variability in testosterone, and
10 the use of the Student *t*-test, an inappropriate statistical test for this analysis. There is some concern with
11 the dynamic range of the 17 β -estradiol RIA as 17 β -estradiol is normally measured in pg/mL.

12
13 **Utility (Adequacy) for CERHR Evaluation Process:** Based on experimental design concerns, this study
14 is not useful in the evaluation.

15
16 **Takahashi and Oishi (438),** support not indicated, examined species, strain, and route differences in
17 reproductive systems of male rodents exposed to bisphenol A. The studies in rats are discussed in this
18 section, and the studies in mice are discussed in Section 4.2.2.2. Animals were housed in stainless steel
19 suspended cages or “chip-bedded” plastic cages. **[No information was provided about the type of chow**
20 **used.]** Animals used in this study were 4 weeks old at the start of dosing. In the dietary portion of the
21 study, male Wistar rats or Holtzman SD rats were given feed containing 0 or 0.25% bisphenol A (>99.0%
22 purity) for 2 months. There were 8 animals in each dose group. The 0.25% dose group was reported to
23 produce minimal testicular effects in a previous study. Mean bisphenol A intakes were estimated by study
24 authors at ~200 mg/kg bw/day in rats. In parenteral exposure studies, 4-week-old male Wistar rats were sc
25 dosed with bisphenol A in propylene glycol at 0 or 200 mg/kg bw on 4 days/week for 1 month.
26 Additional male Wistar rats were given ip injections of bisphenol A in propylene glycol at 0, 2, or 20
27 mg/kg bw 4 days/week for 1 month. An ip dose of 200 mg/kg bw was originally administered but resulted
28 in death. There were 5–6 animals/group in the parenteral exposure studies. In both the dietary and
29 parenteral exposure studies, animals were observed daily for clinical signs, and body weight and food
30 intake were measured. Animals were killed at the end of the dosing period. Liver, kidney, and
31 reproductive organs were weighed. Testes were fixed in formaldehyde solution and examined
32 histologically. The study authors noted that the appropriate fixative for the testis is Bouin solution but that
33 obvious and severe injuries could be detected with the method used in the present study. Testosterone was
34 measured in serum by ELISA. Daily sperm production and efficiency and epididymal sperm reserves
35 were evaluated. Statistical analyses included *F* test, Student *t*-test, Aspin-Welch test, Bartlett test,
36 ANOVA, Dunnett test, Kruskal-Wallis test, Dunnett non-parametric test, Wilcoxon rank-sum test, chi-
37 squared test, Mantel-Haenzel test, and Fisher exact test.

38
39 In rats exposed through diet, there was no effect on body weight or absolute organ weight. Relative liver
40 weight was significantly increased in Wistar rats exposed to bisphenol A. **[Data were not shown by**
41 **study authors.]** The study authors indicated that they forgot to weigh seminal vesicles and prostate
42 glands. No effects were reported for reproductive organ histopathology, daily sperm production or
43 efficiency of production, epididymal sperm reserves, or serum testosterone levels in rats exposed to
44 bisphenol A through diet. **[Data were not shown by study authors.]**

45
46 In the portion of the study where rats were administered 200 mg/kg bw bisphenol A, stiffness was
47 observed at the injection site. Terminal body weight was lower **[by 20%]** in treated rats. Treatment
48 resulted in **[~20%]** decreases in absolute liver, kidney, preputial gland, and testis weight and **[~40–80%]**
49 decreases in epididymis, seminal vesicle, and prostate weight. The study authors also reported decreases
50 in relative weights of epididymis, seminal vesicle and coagulation gland, and prostate. **[Data were not**
51 **shown. The Expert Panel assumes that by coagulation gland, the authors mean the anterior**

4.0 Reproductive Toxicity Data

1 **prostate or coagulating gland.** No histopathological alterations were observed in the seminiferous
 2 tubules of control animals. In the bisphenol A group, histopathological observations (incidence) in
 3 seminiferous tubules included focal atrophy (60%), exfoliation (60%), detachment (20%), missing stage
 4 VII/VIII spermatids (40%), retention of stage IX/XI spermatids (60%), and loss of basement membrane
 5 (20%). Bisphenol A treatment reduced daily sperm production [**by ~25%, as estimated from a graph**
 6 **for total production but not per g testis.**] Reserves in head and body of the epididymis and the cauda
 7 epididymis were also reduced/g of tissue in bisphenol A-treated rats [**by ~43 % in the head and body of**
 8 **epididymis and 63% in the cauda epididymis, as estimated from a graph**]. There was no significant
 9 effect on serum testosterone level.

10
 11 Effects in rats administered bisphenol A by ip injection are summarized in Table 106. At 20 mg/kg bw,
 12 terminal body weight and prostate, liver, and kidney weight were reduced. Serum testosterone levels were
 13 also reduced in rats from the 20 mg/kg bw/day group. There were no effects on testicular histopathology
 14 or sperm endpoints. [**Data were not shown by study authors.**] Enlarged ileum was observed at necropsy
 15 in the 20 mg/kg bw group and histopathological examination revealed mucosal degeneration and
 16 hyperplastic duodenum, jejunum, ileum, and cecum. The study authors concluded that bisphenol A is
 17 more toxic through sc and ip exposure routes than by oral exposure in the diet.

18
 19 **Table 106. Effects in Rats Given Bisphenol A by IP Injection**

Endpoint	Dose, mg/kg bw					
	2	20	BMD ₁₀	BMDL ₁₀	BMD _{1SD}	BMDL _{SD}
Weight						
Terminal body	↔	↓12%	19	12	17	5
Ventral prostate	↔	↓29%	7	5	9	6
Liver	↔	↓18%	14	8	12	6
Kidney	↔	↓12%	20	11	19	6
Serum testosterone	↔	↓69%	3	2	16	9

↑,↓ Statistically significant increase, decrease; ↔ no statistically significant effect.
 From Takahashi and Oishi (438).

20
 21 **Strengths/Weaknesses:** This paper reports a comprehensive and well conducted study with an adequate
 22 number of animals per group. The paper contrasted the effects of the route of bisphenol A administration
 23 as well as the potential sensitivity of different strains of rat to bisphenol A-related toxicity. The data
 24 suggest that systemic exposure is necessary for bisphenol A estrogenic activity to be exhibited and
 25 strongly indicate that route of administration (oral vs. ip) is an important consideration. A minimal
 26 exposure range; the study did not explore low doses.

27
 28 **Utility (adequacy) of CERHR Evaluation Process:** Due to differences in strain sensitivities, a NOAEL
 29 was not established. Nevertheless, it is likely to be near 0.25% in the diet. This study is useful.

30
 31 **Herath et al. (439),** supported by Japan Society for Promotion of Science and the Japanese Ministry of
 32 Education, Culture, Sports, Science, and Technology, examined the effects of bisphenol A exposure on
 33 reproductive hormones and sperm endpoints in male rats. Octylphenol was also examined in this study,
 34 but results will not be discussed. Wistar-Imamichi rats were fed a soy-containing commercial feed (Nosan
 35 Corporation, Japan) and housed in metal cages. Rats were randomly assigned to groups and beginning at
 36 50 days of age, 10–11 rats/group were sc injected with bisphenol A (≥95% purity) at 0 (DMSO vehicle)
 37 or 3 mg/kg bw/day for 5 weeks. Rats were weighed during the study. LH, testosterone, and progesterone
 38 concentration were measured in blood after 2 weeks of treatment and on the following day, 1 hour after a
 39 challenge with gonadotropin-releasing hormone. Rats were killed after 5 weeks of treatment. Blood was
 40 obtained for measurement by RIA of LH, progesterone, testosterone, immunoreactive inhibin, and insulin
 41 growth factor 1 levels. The testis, seminal vesicle, epididymis, and prostate were weighed, and sperm

4.0 Reproductive Toxicity Data

1 counts and motility were determined. A total of 5–11 rats/group were examined for each endpoint.
2 Statistical analyses included ANOVA and Duncan Multiple Range test.

3
4 No statistically significant effects on baseline LH, testosterone, or progesterone levels were observed
5 following 2 weeks of bisphenol A treatment. Following injection with gonadotropin-releasing hormone,
6 LH levels were significantly increased in the bisphenol A group and progesterone levels were
7 significantly increased in the vehicle control group. In the bisphenol A group compared to the control
8 group, incremental increases following injection with gonadotropin-releasing hormone were smaller for
9 testosterone [**~410 vs. 875%**] and progesterone [**~75 vs. 510%**]; statistical significance was reported for
10 the progesterone effect. Following 5 weeks of bisphenol A treatment, significant effects on plasma
11 hormone levels compared to controls included decreased testosterone [**by ~55%**] and increased insulin-
12 like growth factor 1 [**by ~20%**]. Ventral prostate weight was significantly higher [**by ~29%**] in the
13 bisphenol A versus control group, but there were no effects on testis, seminal vesicle, or right epididymis
14 weight. [**Relative reproductive organ weights were not reported.**] Epididymal sperm counts were
15 significantly reduced [**by ~10%**] in the bisphenol A group, but there was no significant effect on sperm
16 motility. The study authors concluded that bisphenol A exposure can affect basal and gonadotropin-
17 releasing hormone-stimulated LH production and reduced daily sperm production in rats.

18
19 **Strengths/Weaknesses:** This study appears to have been relatively well conducted. A major weakness of
20 this paper is the inconsistency in the hormone data (control data after 2 weeks were dramatically different
21 than after 5 weeks even though both are from sexually mature rats), which severely detracts from the
22 utility of these data. The route of administration was not relevant for human risk assessment.

23
24 **Utility (Adequacy) for CERHR Evaluation Process:** Due to variable responses in the control rat
25 hormone levels, these data are of no utility. Moreover, the route of administration is not relevant for
26 human risk assessment.

27
28 **Toyama et al. (440)**, supported in part by the Japanese Ministry of Environment and Ministry of
29 Education, Science, Sports, and Culture, examined the effects of bisphenol A exposure on the
30 reproductive system of male rats and mice. [**No information was provided about feed, caging, or**
31 **bedding materials. The mouse portion of the study is discussed in Section 4.2.2.2.**] Adult male Wistar
32 rats (n = 12/group) were sc injected with bisphenol A at 0.020 or 0.200 mg/kg bw/day for 6 consecutive
33 days. Three control animals were sc injected with the DMSO/olive oil vehicle for 6 days. Ten
34 animals/bisphenol A group and 2 controls were killed the day following treatment and perfused with
35 glutaraldehyde. Testes were weighed and examined by light and electron microscopy. Epididymis,
36 preputial gland, ventral prostate, and seminal vesicle with coagulating glands were also weighed. The
37 remaining animals, 2 in each bisphenol A group and 1 in the control group, were held an additional 2
38 months and then subjected to fertility tests. In fertility testing, each male was mated to 2 untreated
39 females. One of the 2 mated females was kept until parturition. [**The males were apparently killed for**
40 **an examination of reproductive organs following fertility testing.**] Results were qualitatively reported,
41 and statistical analyses were not conducted.

42
43 The description of the results was limited primarily to rats in the 0.020 mg/kg bw/day group. Body and
44 male accessory reproductive organ weights were not affected by bisphenol A treatment. [**Data were not**
45 **shown by study authors.**] In the bisphenol A group, examination by light microscopy revealed
46 exfoliation of round spermatids, deformed heads of mature spermatids, and multinucleated giant cells in
47 seminiferous epithelium. Testicular effects observed by electron microscopy included abnormal
48 acrosomal caps and invagination and/or vacuole formation in nuclei of spermatids beyond step 1.
49 Ectoplasmic specialization around Sertoli cells was also affected by bisphenol A treatment. No
50 histological or ultrastructural abnormalities were observed in the testis 2 months following exposure.
51 Sexual behavior was observed to be normal in treated males. Females delivered normal pups and litter

4.0 Reproductive Toxicity Data

1 sizes were similar between groups. The study authors concluded that bisphenol A exposure did not affect
2 fertility in rats and that adverse effects were transient.

3
4 **Strengths/Weaknesses:** Definite conclusions cannot be drawn from such a limited data set; the fertility
5 assessment was not meaningful due to the sample size (2/group). The background incidence of the
6 electron microscope findings was not discussed.

7
8 **Utility (Adequacy) for CERHR Evaluation Process:** This study is not useful in the evaluation.

9 10 4.2.2.2 Mouse

11 **Takao et al. (441)**, support not indicated, examined the effects of bisphenol A exposure on the
12 reproductive system of mice. Five-week-old male C57BL/6 mice were exposed to bisphenol A in
13 drinking water at 0 (0.005% ethanol in water vehicle), 0.0005, or 0.050 g/L for 4 or 8 weeks. **[Based on**
14 **daily water intakes and body weights reported in the study, bisphenol A intake was estimated by**
15 **CERHR at 0.14 and 13 mg/kg bw/day.]** To maintain bisphenol A at a stable concentration, drinking
16 water was changed twice a week, but the stability of bisphenol A was not verified. Mice were killed, and
17 both testes and spleen were removed and weighed. One testis was processed for histopathological
18 evaluation. Plasma testosterone, corticosterone, and LH levels were measured in 7 mice/group using RIA
19 or enzyme immunoassay. **[No information was provided on the number of mice exposed/group,**
20 **purity of bisphenol A, time between last dose and sacrifice, or the type of chow, caging, or bedding**
21 **materials used. Very few details were provided on the methods, including histopathological**
22 **evaluation.]** Statistical analyses included ANOVA followed by Fisher protected least significant
23 difference test.

24
25 Water intake was significantly reduced **[by 8%]** in the high-dose group exposed for 4 weeks. There were
26 no effects on body weight or absolute or relative (to body weight) testis or spleen weight. Plasma
27 testosterone levels were reduced **[by 87–89%]** in the high-dose group, but statistical significance was
28 attained only in the group exposed for 8 weeks. No statistically significant changes were reported for
29 plasma corticosterone or LH levels. The number of multinucleated cells in the seminiferous tubules was
30 increased in high-dose mice treated for 8 weeks. The study authors concluded that exposure to bisphenol
31 A around the peripubertal period may disrupt the reproductive tracts of male mice.

32
33 **Strengths/Weaknesses:** This study appears to have been well conducted, and it appears that bisphenol A
34 in the drinking water results in a dose-related decrease in plasma testosterone. However, this endpoint is
35 highly variable because testosterone is secreted in a pulsatile manner, and controls for the week 4 and 8
36 varied by ~30%. The study lacks important details on the methods used for fixation and embedding of the
37 testes and the incidence of multinucleated germ cells in individual animals. In the absence of incidence
38 data, it cannot be determined if 1 mouse exhibited a greater response by chance, or if it is a generalized
39 effect.

40
41 **Utility (Adequacy) for CERHR Evaluation Process:** Due to the paucity of important experimental
42 details and the variability of the testosterone data, this study has limited utility.

43
44 **Al-Hiyasat (442)**, supported by the Deanship of Scientific Research at Jordan University of Science and
45 Technology, examined the effect of bisphenol A exposure on fertility of male mice. **[No information was**
46 **provided about composition of chow, bedding, or caging.]** Ten 60-day-old male Swiss mice/group
47 were gavaged with the ethanol/distilled water vehicle or bisphenol A (97% purity) for 30 days. **[The**
48 **study listed the bisphenol A doses as 5, 25, and 100 ng/kg bw. An erratum was later released that**
49 **indicated the correct units were µg/kg bw (0.005, 0.025, and 0.1 mg/kg bw/day).]** Following the
50 dosing period, each male was mated for 10 days with 2 untreated female mice, who were placed inside the
51 cage of the male during the same time period. The males were then killed for an evaluation of testes,

4.0 Reproductive Toxicity Data

1 seminal vesicles, and preputial gland weights. Sperm counts and daily sperm production were determined.
 2 Mated females were killed 10 days later to determine numbers of pregnancies, implantation sites, viable
 3 fetuses, total resorptions, and females with resorptions. [There was no indication that mating was
 4 confirmed by checking for sperm in the vagina.] Data were analyzed by Student *t*-test or Fisher exact
 5 test.

6
 7 Results that obtained statistical significance are summarized in Table 107. Body weights were lower in all
 8 dose groups compared to controls. There were no evident dose-response relationships for organ weights.
 9 Absolute testis weight was decreased at the low dose, and absolute seminal vesicle weight was reduced at
 10 the mid and high dose. Effects on relative organ weights are summarized in Table 107. Decreases in
 11 testicular sperm counts and daily sperm production were observed at the mid and high dose. Total sperm
 12 counts in the epididymis were decreased at all dose levels, and sperm counts/mg epididymis were
 13 decreased at the mid and high dose. The total number of resorptions and females with resorptions were
 14 increased at all dose levels. The percentage of pregnant females was reduced at the mid and high dose.
 15 The study authors concluded that bisphenol A could adversely affect fertility and reproduction of adult
 16 male mice.

17
 18 **Table 107. Effects Observed Following Gavage of Male Mice with Bisphenol A and Mating with**
 19 **Untreated Females**

Endpoint	Dose, mg/kg bw/day				BMD ₁₀	BMDL ₁₀	BMD _{1SD}	BMDL _{1SD}
	0.005	0.025	0.1					
Body weight	↓18%	↓21%	↓13%					
Relative weight								
Testis	↔	↑26%	↔					
Seminal vesicle	↔	↓27%	↔					
No. sperm/testis	↔	↓17%	↓29%	0.035	0.029	0.036	0.028	
No. sperm/mg testis	↔	↓16%	↓37%	0.027	0.023	0.029	0.023	
Daily sperm production	↔	↓17%	↓29%	0.035	0.029	0.036	0.028	
Efficiency of sperm production	↔	↓16%	↓37%	0.027	0.023	0.029	0.023	
No. sperm/epididymis	↓14%	↓25%	↓35%	0.033	0.026	0.040	0.030	
Sperm/mg epididymis	↔	↓17%	↓31%	0.033	0.025	0.053	0.038	
Percent pregnant females	↔	↓40%	↓33%					
Resorptions/implantation site (3% control rate)	13%	15%	13%					
Percent females with resorption sites	↑2.5-fold	↑3.8-fold	↑3.4-fold					

↑,↓ Statistically significant increase, decrease, ↔ no statistically significant effect.
 From Al-Hiyasat (442).

20
 21 **Strengths/Weaknesses:** The number of animals per group was too small for a definitive assessment. The
 22 method of randomization (or initial body weights) was not presented. There is also an absence of a dose
 23 response in several of the endpoints assessed. Given that mice usually have poor (relative to rats) fertility
 24 rates, collectively the controls in this study are suspect. The male mice were killed shortly after the
 25 mating period, which may have influenced/confounded the number of sperm in the epididymis. Student *t*-
 26 test is an inappropriate analysis for organ weights (ANOVA with appropriate post hoc test would be
 27 appropriate). Statistical significance is suspect, and the changes in organ weights are minimal in
 28 magnitude.

29
 30 **Utility (adequacy) of CERHR Evaluation Process:** Given the limitations of the study design, this study
 31 is of minimal utility.

4.0 Reproductive Toxicity Data

1
2 **Nagao et al. (369)**, support not indicated, examined the effects of bisphenol A in mice following
3 exposure during different life stages. An initial experiment, described in more detail in Section 3.2.7,
4 found that C57BL/6N mice were more sensitive to 17 β -estradiol than ICR mice, and the study authors
5 therefore used C57BL/6N mice to examine the effects of bisphenol A. Life stages examined included
6 prenatal development, adolescence, and adulthood. The study conducted in adult mice is described here,
7 while the studies conducted during prenatal development and adolescence are described in Section 3.2.7.
8 C57BL/6N mice were fed PLD (phytoestrogen-low diet, Oriental Japan). They were housed in
9 polycarbonate cages with wood bedding. Daidzein and genistein levels were analyzed in diet, tap water,
10 and bedding and found to be below 0.5 mg/100 g. At 10 weeks of age, 20 male mice/group were gavaged
11 with bisphenol A (99.0% purity) at 0.002, 0.020, or 0.200 mg/kg bw/day for 6 days. Twenty control
12 males/group were given 0.5% carboxymethyl cellulose [**assumed to be the vehicle**]. Six weeks after the
13 final dose was administered, the mice were weighed and 15 males/group were killed and necropsied. The
14 testis, epididymis, and seminal vesicles with coagulating glands were weighed. The ventral prostate was
15 not weighed due to difficulties in obtaining only prostate and determining the precise weight. Epididymal
16 sperm counts were obtained. Histopathological examinations were conducted for organs fixed in Bouin
17 solution. Data were analyzed by Bartlett test to determine homogeneity of variance, followed by ANOVA
18 when homogeneity of variance was confirmed or Kruskal-Wallis analysis of ranks when variance was not
19 homogenous. Dunnett test was used for multiple comparisons.

20
21 In the bisphenol A group, there were no significant differences in body weight gain or terminal body
22 weights. [**Data were not shown.**] There were no significant differences in absolute or relative (to body
23 weight) weights of the testis, epididymis, or seminal vesicles. There were no significant effects on sperm
24 count. No histopathological alterations in reproductive organs were reported. The study authors concluded
25 that low-dose bisphenol A exposure of mice did not reduce sperm density.

26
27 **Strengths/Weaknesses:** This study was extremely well conducted and significantly adds to the
28 understanding of the potential effects of low doses of bisphenol A administered by a relevant route of
29 exposure. Strengths are an appropriate number of mice per group, examination of 2 strains (1 which
30 demonstrated a greater sensitivity to 17 β -estradiol), inclusion of 17 β -estradiol as a positive control, and
31 the presentation of sperm data in light of historical control data.

32
33 **Utility (Adequacy) for CERHR Evaluation Process:** This study identified a NOAEL at >200 μ g/kg
34 bw/day in the adult mouse (the highest dose level tested). This study is highly useful.

35
36 **Peknicová et al. (443)**, supported by the Czech Republic and EU, examined the effects of bisphenol A
37 exposure on mouse sperm. CD-1 mice were given ST1 feed (Velaz a.s., Prague). Three generations of
38 mice were exposed to bisphenol A through drinking water at doses of 0.000002 and 0.000020 mg
39 “/animal’s weight/day.” It was stated that there were 6 pairs of mice in the control group. Litter size was
40 evaluated in 3 generations; 1 litter was examined in the first and second generation and 2 litters were
41 examined in the third generation. In each generation, samples of sperm were collected from all males and
42 a histopathological investigation of testes was conducted in ≥ 3 males/group. Sperm acrosomal status was
43 assessed using an immunohistochemical and Western blot method. Statistical analyses included ANOVA
44 and Newman–Keuls test. [**Very few experimental details were provided. No information was**
45 **provided on bedding and caging materials, bisphenol A purity, the numbers of mice in each**
46 **treatment group, treatment of the control group, ages of mice during treatment, durations of**
47 **treatment, sample sizes and litter representation for sperm effects, and mating procedures. It was**
48 **not clear if female rats were also treated.**] Litter sizes were significantly reduced in the first and second
49 generation of mice treated with the low dose (5–6.7 pups/litter vs. 11.5–12 pups/litter in controls). There
50 were no effects of bisphenol A treatment on testes weight. [**Data were not shown by authors.**]
51 Pathological changes observed in testes from the low-dose group included damaged seminiferous tubule

4.0 Reproductive Toxicity Data

1 and reduced spermatogenesis. Acrosome integrity, evaluated as percent cells binding monoclonal
2 antibodies to acrosin and intra-acrosomal proteins, was significantly reduced in all 3 generations of the
3 low-dose group (48.5–57.7 compared to 93.3–95% integrity in controls) and the third generation of the
4 high-dose group (62.5 compared to 93.3% integrity in controls). **[While the text of the study stated that
5 acrosomal integrity was significantly affected only in the third generation of the high-dose group,
6 the caption for Figure 7 of the study stated that both the second and third generations were
7 significantly affected. Based on findings reported in the figure, it appears that the description in the
8 text is correct.]** The study authors concluded that bisphenol A exposure negatively impacts fertility,
9 spermatogenesis, and sperm quality in mice.

10
11 **Strengths/Weaknesses:** Although potentially interesting findings are presented, the study lacks important
12 details.

13
14 **Utility (Adequacy) for CERHR Evaluation Process:** Due to the absence of critical study design
15 information, this study has no utility for the evaluation.

16
17 **Takahashi and Oishi (438)**, support not indicated, examined species, strain, and route differences in
18 reproductive systems of male rodents exposed to bisphenol A. Studies on mice are discussed here, and
19 studies on rats are discussed in Section 4.2.2.1. Animals were housed in stainless steel suspended cages or
20 “chip-bedded” plastic cages. **[No information was provided about the type of chow used.]** Animals
21 used in this study were 4 weeks old at the start of dosing. In the dietary portion of the study, CD-1 (ICR)
22 mice and C57BL/6CrSlc mice were given feed containing 0 or 0.25% bisphenol A (>99.0% purity) for 2
23 months. There were 8 animals in each dose group. The 0.25% dose was reported to produce minimal
24 testicular effects in a previous study. Mean bisphenol A intakes were estimated by study authors at ~400
25 mg/kg bw/day in mice. The parenteral exposure studies were performed only in rats. Animals were
26 observed daily for clinical signs, and body weight and food intake were measured. Animals were killed at
27 the end of the dosing period. Liver, kidney, and reproductive organs were weighed. Testes were fixed in
28 formaldehyde solution and examined histologically. The study authors noted that the appropriate fixative
29 for the testis is Bouin solution, but that obvious and severe injuries could be detected with the method
30 used in the present study. Testosterone was measured in serum by ELISA. Daily sperm production and
31 efficiency and epididymal sperm reserves were evaluated. Statistical analyses included *F* test, Student *t*-
32 test, Aspin-Welch test, Bartlett test, ANOVA, Dunnett test, Kruskal-Wallis test, Dunnett non-parametric
33 test, Wilcoxon rank-sum test, chi-squared test, Mantel-Haenzel test, and Fisher exact test.

34
35 There were no significant effects on organ or body weights in C57BL/6CrSlc mice exposed through diet.
36 In CD-1 (ICR) mice exposed through diet, there were increases in absolute testis **[16%]**, liver **[12%]**, and
37 kidney **[20%]** weights and a decrease in absolute epididymis **[12%]** weight. The study authors reported
38 that relative testis weight was not significantly affected, but when the value from 1 mouse with a high
39 relative testis weight was deleted, the effect attained statistical significance. **[Data were not shown by
40 study authors.]** No effects were reported for testis histopathology, daily sperm production or efficiency
41 of production, epididymal sperm reserves, or serum testosterone levels in mice exposed to bisphenol A
42 through diet. **[Data were not shown by study authors.]** The study authors concluded that the testicular
43 toxicity of bisphenol A is “relatively weak,” based on the co-occurrence of liver and kidney toxicity at
44 exposure levels causing testicular effects.

45
46 **Strengths/Weaknesses:** This study appears to have been well designed and conducted. Although not
47 statistically significant, there is an apparent ~2-fold increase in testosterone levels. It is unfortunate that
48 an additional dose level was not used to determine if the potential trends were dose-related. The route of
49 administration was appropriate.

50

4.0 Reproductive Toxicity Data

1 **Utility (adequacy) of CERHR Evaluation Process:** Because this study has used 1 dose level of
2 bisphenol A (the maximum tolerated dose) it has marginal utility. The NOAEL is <400 mg/kg bw/day.

3
4 **Park et al. (421)**, support not indicated, examined the effects of bisphenol A exposure on the
5 reproductive and hematological systems of male and female mice. **[Results for males are discussed**
6 **here, and results for females are discussed in Section 4.2.1.2.]** Adult ICR mice were fed mouse
7 formulation feed (Cheil Feed). **[No information was provided about caging or bedding materials.]**
8 Fifteen mice/sex/group were ip injected with bisphenol A in an ethanol/corn oil vehicle at 0.05, 0.5, or 5.0
9 mg/kg bw on 5 occasions (every 3 days over a 14-day period). One control group received no treatment,
10 and a second control group was ip injected with corn oil. Males were examined 2 days following
11 administration. Semen was collected and assessed for sperm number, viability, and motility. Reproductive
12 organs were weighed and fixed in Bouin solution, and histopathological examination was conducted.
13 Hematological and clinical chemistry endpoints were also assessed. Data were analyzed by least
14 significant difference test.

15
16 Exposure to bisphenol A had no effect on body weight or on weights of male reproductive organs
17 including testis, epididymis, vesicular gland, or coagulating gland. Reductions in sperm concentrations
18 **[by 18%]** and increases in sperm abnormalities **[by 28%]** were significant in the high-dose group. There
19 were no treatment effects on testicular histology. There were no significant effects on hematological or
20 clinical chemistry endpoints in males treated with bisphenol A. The study authors did not report
21 conclusions regarding study findings.

22
23 **Strengths/Weaknesses:** Frequency of administration was every 3 days and, given the half-life of the
24 chemical, it is unlikely that sufficient blood chemical levels were sustained to induce “maximal”
25 bisphenol A-mediated estrogenic responses. The route of administration was not relevant for human risk
26 assessment. Sperm viability appears low, perhaps as a result of differences in methodology and/or
27 substrain of mouse.

28
29 **Utility (Adequacy) for CERHR Evaluation Process:** Given the dosing paradigm (ip injection every 3
30 days) this study is of minimal value.

31
32 **Toyama et al. (440)**, supported in part by the Japanese Ministry of Environment and Ministry of
33 Education, Science, Sports, and Culture, examined the effects of bisphenol A exposure on the
34 reproductive system of male rats and mice. **[No information was provided about feed, caging, or**
35 **bedding materials. The mouse study is discussed here, and the rat study is discussed in Section**
36 **4.2.2.1.]** Adult male ICR mice (n = 12/group) were sc injected with bisphenol A at 0.020 or 0.200 mg/kg
37 bw/day for 6 consecutive days. Three control animals were sc injected with the DMSO/olive oil vehicle
38 for 6 days. Ten animals/bisphenol A group and 2 controls were killed the day following treatment and
39 perfused with glutaraldehyde. Testes were weighed and examined by light and electron microscopy.
40 Epididymis, preputial gland, ventral prostate, and seminal vesicle with coagulating glands were also
41 weighed. The remaining animals, 2 males in each bisphenol A treatment group and 1 control male, were
42 held an additional 2 months and then subjected to fertility tests. In fertility testing, each male was mated
43 to 2 untreated females. One of the 2 mated females was kept until parturition. **[The males were**
44 **apparently killed for an examination of reproductive organs following fertility testing.]** Results were
45 qualitatively reported, and statistical analyses were not conducted.

46
47 The study authors noted that all effects were similar between rats and mice and between dose groups, and
48 their description of results was primarily limited to rats in the 0.020 mg/kg bw/day group. Body and male
49 accessory reproductive organ weights were not affected by bisphenol A treatment. **[Data were not shown**
50 **by study authors.]** In the bisphenol A group, examination by light microscopy revealed exfoliation of
51 round spermatids, deformed heads of mature spermatids, and multinucleated giant cells in seminiferous

4.0 Reproductive Toxicity Data

1 epithelium. Testicular effects observed by electron microscopy included abnormal acrosomal caps and
2 invagination and/or vacuole formation in nuclei of spermatids beyond step 1. Ectoplasmic specialization
3 around Sertoli cells was also affected by bisphenol A treatment. No histological or ultrastructural
4 abnormalities were observed in testes 2 months following exposure. Sexual behavior was observed to be
5 normal in treated males. Females delivered normal pups and litter sizes were similar between groups. The
6 study authors concluded that bisphenol A exposure did not affect fertility in mice and that adverse effects
7 were transient.

8
9 **Strengths/Weaknesses:** It is not possible to draw definite conclusions from such a limited data set; the
10 fertility assessment was not meaningful due to the small sample size (2/group). The background incidence
11 of the electron microscopy findings was not discussed.

12
13 **Utility (Adequacy) for CERHR Evaluation Process:** Due to the limited number of animals per group,
14 definite conclusions cannot be made. Therefore, this study has minimal value.

15
16 **Anahara et al. (444)**, supported by the Japanese Ministry of Environment and Ministry of Education,
17 Culture, Sports, Science, and Technology, examined the effects of bisphenol A exposure on expression of
18 cortactin protein in the mouse testis. Cortactin is an actin binding protein that makes up the apical
19 ectoplasmic specialization between Sertoli cells and spermatids and the basal ectoplasmic specialization
20 between Sertoli cells. Cortactin is one of several proteins that control spermatid development. Adult (12-
21 week-old) male ICR mice (n = 5–7/group) were sc injected with corn oil vehicle, 0.0024 mg/kg bw/day
22 bisphenol A, 2.5 µg/kg bw/day diethylstilbestrol, or 1.2 µg/kg bw/day 17β-estradiol for 5 days. **[No
23 information was provided on purity of bisphenol A or the types of feed, caging, or bedding used.]**
24 Animals were killed on the day following the last injection. Testes were homogenized and expression of
25 cortactin protein was determined in testes from 5–7 rats/group by Western blot, immunohistochemistry,
26 and immunoelectron microscopy techniques. Data were analyzed by *t*-test. Exposure to bisphenol A
27 resulted in a significant decrease in testicular cortactin protein expression **[to ~60% of control levels]**.
28 Immunohistochemical analysis revealed that cortactin staining was reduced in the apical ectoplasmic
29 specialization but not in the basal ectoplasmic specialization. Examination by immunoelectron
30 microscopy revealed no expression of cortactin around heads of spermatid and deformation of nuclei and
31 acrosomes. Effects observed with 17β-estradiol and diethylstilbestrol were similar to those observed with
32 bisphenol A, with the exception that diethylstilbestrol also reduced cortactin protein expression in the
33 basal ectoplasmic specialization and did not result in deformation of spermatids. The authors concluded
34 that exogenous chemicals can damage junctional proteins like cortactin and have adverse effects on
35 Sertoli cell protein regulation.

36
37 **Strengths/Weaknesses:** The route of administration was not relevant. The lack of additional dose levels
38 of bisphenol A makes interpretation of the significance of these data challenging. Western blot analysis of
39 cortactin was inappropriately presented as a function of the control value with no variability in the control
40 sample. There were no apparent differences in levels of protein expression between various estrogenic
41 agents/treatments. No adverse outcomes of the changes in cortactin were explored.

42
43 **Utility (Adequacy) for CERHR Evaluation Process:** While these data are interesting, the route of
44 exposure and the absence of a correlating adverse outcome makes these data difficult to interpret for risk
45 assessment purposes, limiting the utility of the study.

4.2.2.3 Other mammals

46
47
48 **Moon et al. (445)**, supported by Korea University Medical Science Research Center and the Korean
49 Ministry of Education, examined the effects of bisphenol A exposure on penile function in rabbits. **[No
50 information was provided on feed or caging and bedding materials.]** Male, 8–12 week-old New
51 Zealand white rabbits were ip injected with corn oil vehicle or 150 mg/kg bw bisphenol A **[purity not**

4.0 Reproductive Toxicity Data

1 **reported**], every other day for 12 days to a cumulative dose of 900 mg/kg bw [75 mg/kg bw/day].
2 Rabbits were killed at 4 weeks (n = 15/group) and 8 weeks (n=15/group) following bisphenol A
3 treatment. In 5 rabbits/group, the penis was removed and fixed in 10% neutral buffered formalin for
4 histological examination. In 10 rabbits/group, the corpora cavernosa were removed from the penis, and in
5 vitro responses to norepinephrine, acetylcholine, sodium nitroprusside, and L-arginine were studied. Data
6 were analyzed by Student *t*-test. Treatment with bisphenol A significantly suppressed contraction of
7 corpora cavernosa in response to norepinephrine and relaxation in response to acetylcholine, sodium
8 nitroprusside, and L-arginine at both stages of evaluation. Histopathological observations in the bisphenol
9 A-treated rabbits but not control rabbits at both ages included intracavernosal fibrosis in conjunction with
10 decreased sinusoidal spaces. Compared to rabbits in the control group, both age groups of rabbits exposed
11 to bisphenol A had significantly increased trabecular smooth muscle content (73.3–83.2 versus 33.2% in
12 controls) and a non-significant difference in thickness of tunica albuginea (0.93–1.12 mm versus 0.32–
13 0.43 mm in controls). The study authors concluded that bisphenol A may affect erectile responses by
14 inducing histological alterations in the penis.

15
16 **Strengths/Weaknesses:** There is no evidence that bisphenol A has any effect on the ability to attain an
17 erection resulting in copulation in mice or rats. This study does not have a concurrent control (e.g., 17 β -
18 estradiol) to ascertain if the observed effects are the result of estrogenic responses in the penis. The route
19 of administration is not relevant for human risk assessment.

20
21 **Utility (Adequacy) for CERHR Evaluation Process:** Although interesting this study provides no utility
22 for bisphenol A risk assessment.

23
24 **Nieminen et al. (423)**, support not indicated, examined the effects of bisphenol A exposure on hormone
25 levels in the European polecat (*Mustela putorius*). There were no significant effects on plasma levels of
26 testosterone, estradiol, FSH, or thyroid hormones. Details of this study are discussed in Section 4.2.1.3.

27
28 **Strengths/Weaknesses:** This study provides evidence that the bisphenol A administered to polecats
29 increases GST and UDPGT activity. Since these findings were dose-related it appears that in the polecat
30 bisphenol A increases phase 2 metabolism but has minimal effects on hormone levels. Due to the limited
31 number of animals and the absence of a dose-response relationship, the hormonal changes in this study
32 are difficult to interpret.

33
34 **Utility (Adequacy) for CERHR Evaluation Process:** Due to the small sample size and absence of
35 effects on reproductive endpoints, this study is of no utility in the evaluation.

36
37 **Nieminen et al. (424)**, support not indicated, examined the effects of bisphenol A exposure on endocrine
38 endpoints in field voles (*Microtus agrestis*). Animals were housed in plastic cages with wood shavings
39 and fed R36 diet (Lactamin, Sweden). Sexually mature field voles were randomly assigned to groups that
40 received bisphenol A [**purity not reported**] in propylene glycol by sc injection for 4 days. Doses of
41 bisphenol A (numbers of males in each group) were 0 (n = 6), 10 (n = 4), 50 (n = 6), and 250 (n = 7)
42 mg/kg bw/day. Animals were killed the day following the last dose. Body and liver weights were
43 measured. Blood was drawn for measurement of sex steroids, thyroxine, and weight regulating hormone
44 levels in plasma using RIA or immunoradiometry methods. The activities of EROD, UDPGT, and GST
45 were measured in hepatic and renal microsomes using appropriate substrates. Statistical analyses included
46 ANOVA, post hoc Duncan test, Student *t*-test, Kolmogorov-Smirnov test, Levene test, Mann-Whitney *U*
47 test, chi-squared test, and Spearman correlation. [**Results for males are discussed in Section 4.2.1.3.**]

48
49 Mortality was significantly increased by bisphenol A treatment, with incidences of 18, 36, and 20% in the
50 low-to high-dose groups. No mortality was observed in the control group. Bisphenol A treatment did not
51 significantly affect body, liver, or testis weight. Plasma testosterone levels increased with dose, and

4.0 Reproductive Toxicity Data

1 statistical significance was attained in high-dose males and females combined. Pooled (male + female)
2 LH levels were not significantly altered by treatment. Liver EROD activity [**apparently combined for**
3 **males and females**] was significantly decreased at the mid and high dose and liver GST activities [**not**
4 **clear if for males or females or both**] was significantly decreased at the highest dose level. There were
5 no other significant effects on microsomal enzymes examined. The study authors concluded that wild
6 mammals such as field voles could be more susceptible to bisphenol A-induced toxicity than laboratory
7 rodents.

8
9 **Strengths/Weaknesses:** Weaknesses are the number of voles/dose levels, the sc route of administration,
10 and lack of similar studies in the literature for comparison purposes.

11
12 **Utility (Adequacy) for CERHR Evaluation Process:** This study is not useful in the evaluation.

13 14 4.2.2.4 Fish and invertebrates

15 **Shioda and Wakabayashi (446)**, supported by the Japanese Ministry of Education, examined the effects
16 of bisphenol A exposure on reproductive capability of male medaka (*Oryzias latipes*). Adult male medaka
17 were housed for 2 weeks in glass beakers containing distilled water and bisphenol A at 0, 0.3, 1, 3, or 10
18 μM [**0, 0.07, 0.23, 0.69, or 2.3 mg/L**]. [**The number of male fish treated was not reported. Though**
19 **not specifically stated, it was suggested that fish in the negative control group were exposed to the**
20 **acetone vehicle.**] Following exposure, each male was housed with two females in beakers containing
21 distilled water. The numbers of eggs spawned, fertilized, and hatched were determined. Statistical
22 analyses included *F* test followed by *t*-test or Welch test. Exposure to bisphenol A 10 μM [**2.3 mg/L**]
23 significantly reduced the number of eggs produced and hatched compared to the negative control group.
24 Additional compounds were also examined, and it was reported that eggs and hatchings were significantly
25 reduced following exposure to 17 β -estradiol (≥ 3 nM), but not nonylphenol or diethylhexyl phthalate. The
26 study authors concluded that the reproductive effects induced by bisphenol A in this study occurred at a
27 higher concentration than results observed in a yeast estrogen screen.

28
29 **Strengths/Weaknesses:** This study appears to have been well conducted study and suggests that
30 bisphenol A 2.3 mg/L in water decreases the number of medaka eggs produced and hatched

31
32 **Utility (Adequacy) of CERHR Evaluation Process:** Because this study uses a non-mammalian model,
33 it is not useful in the evaluation.

34
35 **Kinnberg and Toft (447)**, supported by the Danish Environmental Research Programme, examined the
36 effects of bisphenol A exposure on the reproductive system of male guppies (*Poecilia reticulata*). Thirty
37 sexually mature male guppies/group were exposed for up to 30 days to bisphenol A at nominal
38 concentrations of 0 (acetone vehicle) 5, 50, 500, or 5000 $\mu\text{g/L}$. Levels of bisphenol A in water were
39 verified. Exposure to the 5000 $\mu\text{g/L}$ concentration was stopped after 21 days because of a high mortality
40 rate. All fish in the high-dose group and 6 fish/group in the lower dose groups were killed and fixed in
41 neutral buffered formalin. Histopathological examination was conducted in whole fish. The mortality rate
42 in the 5000 $\mu\text{g/L}$ group was 77%, but no increase in mortality was observed in the lower concentration
43 groups. Testes of fish from the high-dose group contained spermatozeugmata (bundles of spermatozoa
44 with heads pointing outward and tails in the center) in ducts, and the authors stated the effect indicated
45 blocked spermatogonial mitosis. [**No information was provided on incidence or severity of testicular**
46 **lesions, and it does not appear that statistical analyses were conducted.**] Additional compounds were
47 also tested, and it was indicated that effects induced by flutamide, 1,1-dichloro-2,2-bis(p-
48 chlorophenyl)ethylene, and 4-tert-octylphenol were similar to those observed with bisphenol A exposure.
49 In contrast, exposure to 17 β -estradiol resulted in hypertrophy of Sertoli cells and efferent duct cells. The
50 study authors concluded that a high bisphenol A induced adverse effects on testicular structure.

4.0 Reproductive Toxicity Data

1 **Strengths/Weaknesses:** This study appears to have been well conducted. The metabolism of bisphenol A
2 in fish is unknown. It appears the bisphenol A does not exhibit the typical 17β -estradiol-like effect on the
3 testis. Findings occurred at high relative exposures. There was no apparent low-dose effect.
4

5 **Utility (Adequacy) for CERHR Evaluation Process:** Because this study uses a non-mammalian model,
6 it is not useful in the evaluation.
7

8 **Oehlmann et al. (426)**, supported by the Berlin Federal Environmental Agency, reported the effects of
9 bisphenol A on reproductive organs in the freshwater ramshorn snail (*Marisa cornuarietis*) and the
10 marine dog whelk (*Nucella lapillus*). Details of this study are discussed in Section 4.2.1.4, and most of
11 the findings pertained to female snails. Adult ramshorn snails did not show abnormalities of male sexual
12 organs or gonads after exposure to bisphenol A concentrations up to 100 $\mu\text{g/L}$ for 5 months or after
13 exposure for the first year of life. In the dog whelk, a 1 month exposure to 1, 25, or 100 $\mu\text{g/L}$ bisphenol A
14 significantly decreased the proportion of males with sperm in the seminal vesicles compared to the
15 vehicle-exposed control. The length of the penis and prostate gland were also reduced by all
16 concentrations of bisphenol A in this animal. The authors concluded that bisphenol A toxicity occurs in
17 invertebrates at environmentally relevant concentrations.
18

19 **Strengths/Weaknesses:** The study appears to have been well conducted and suggests that bisphenol A
20 has an effect on the dog whelk. The potential stability/biotransformation was discussed in the introduction
21 but not determined during the exposure period.
22

23 **Utility (Adequacy) for CERHR Evaluation Process:** Because this study uses a non-mammalian model,
24 it is not useful in the evaluation.
25

26 4.2.2.5 *In vitro*

27 **Nikula et al. (448)**, support not indicated, examined the *in vitro* effects of bisphenol A on steroidogenesis
28 in mouse Leydig tumor cell cultures. Octyl phenols were also examined in this study, but results will not
29 be discussed. In the first experiment, cells were incubated for 48 hours in media containing bisphenol A at
30 0 (ethanol vehicle) or 10^{-7} – 10^{-4} M [**0.023–23 $\mu\text{g/L}$**] or estradiol at 10^{-8} M. Production of cyclic adenosine
31 monophosphate (cAMP) and progesterone was measured following the incubation period and at 1 and 3
32 hours following a challenge with 10 ng/mL hCG. In additional experiments, the cells were exposed to
33 bisphenol A at 0 or 10^{-6} M [**0.23 $\mu\text{g/L}$**] or 17β -estradiol or diethylstilbestrol at 10^{-8} M. Production of
34 cAMP and progesterone was measured following the incubation period and at 1 and/or 3 hours following
35 challenge with hCG, forskolin, cholera toxin, or 8-bromo-cAMP. An additional study measured binding
36 of ^{125}I -hCG to the LH receptor following a 48-hour exposure to bisphenol A at 0 or 10^{-6} M [**0.23 $\mu\text{g/L}$**].
37 Each experiment contained 5–8 replicates, and results from 3 independent experiments were pooled. Data
38 were analyzed by ANOVA followed by Fisher test.
39

40 Bisphenol A had no effect on basal cAMP or progesterone production. At 3 hours following the hCG
41 challenge, the increase in cAMP production was attenuated following previous exposure to bisphenol A at
42 concentrations $\geq 10^{-7}$ M [**0.023 $\mu\text{g/L}$**] and increase in progesterone production was reduced at bisphenol A
43 concentrations $\geq 10^{-6}$ M [**0.23 $\mu\text{g/L}$**]. At 3 hours following challenge, 10^{-6} M [**0.23 $\mu\text{g/L}$**] bisphenol A
44 decreased hCG-induced cAMP production but had no effect on forskolin- or cholera toxin-induced cAMP
45 production. Following 3-hour challenges, hCG-induced progesterone production was reduced following
46 exposure to 10^{-6} M [**0.23 $\mu\text{g/L}$**] bisphenol A, but there were no effects on forskolin-, cholera toxin-, or 8-
47 bromo-cAMP-induced progesterone production. Generally, 17β -estradiol and diethylstilbestrol attenuated
48 hCG-, forskolin, and 8-bromo-cAMP-induced progesterone production. Bisphenol A exposure had no
49 effect on binding of ^{125}I -hCG to the LH receptor. The study authors concluded that bisphenol A appears to
50 inhibit cAMP formation and steroidogenesis in rat Leydig tumor cells by preventing coupling between the
51 LH receptor and adenylate cyclase. Because no inhibition of cAMP production was observed following

4.0 Reproductive Toxicity Data

1 incubation of cells with 17 β -estradiol, the study authors concluded that the effects of bisphenol A may not
2 be estrogen related.

3
4 **Strengths/Weaknesses:** This appears to be a well conducted in vitro study. Stimulation occurred in the
5 absence of steroid-rich fetal bovine serum. There was no mention of whether phenolred-free media were
6 used. Cell viability does not appear to have been determined. Because this study used an in vitro system,
7 the effects of metabolism were limited. Nonetheless, this study provides compelling evidence that the
8 actions of bisphenol A maybe non-estrogen mediated.

9
10 **Utility (Adequacy) for CERHR Evaluation Process:** Because this study was performed in vitro, it has
11 minimal value other than suggesting that not all bisphenol A effects on reproductive tissues are
12 hormonally mediated.

13
14 **Murono et al. (449)**, from the Centers for Disease Control and Prevention, examined the effects of
15 bisphenol A exposure on steroidogenesis in cultured rat Leydig cells. Leydig cell cultures were prepared
16 from testes of 55–65-day-old Sprague Dawley rats (n = 8–10). Cells were incubated in 0 or 1–1000 nM
17 [0.23–230 $\mu\text{g/L}$] bisphenol A in DMSO vehicle, with and without 10 IU/mL hCG for 24 hours.
18 Following the incubation period, testosterone level was measured by RIA and ¹²⁵I-hCG binding to LH
19 receptors was assessed. Media containing hydroxycholesterol was then added to the cultures, and
20 testosterone production following a 4-hour incubation period was measured. The effects of 17 β -estradiol
21 and 4-*tert*-octylphenol were also examined, but will not be discussed. Cell viability was evaluated by
22 trypan blue exclusion and found to be unaffected at the bisphenol A concentrations used in this study.
23 Three experiments with 4 samples/experiment were conducted. Data were analyzed by ANOVA and
24 Student-Newman-Keuls test. Bisphenol A had no effect on basal or hCG-induced testosterone production
25 or hCG binding to LH receptors. [Data wer not shown by study authors.] Conversion of
26 hydroxycholesterol to testosterone was also unaffected by exposure of Leydig cells to bisphenol A. No
27 effect on testosterone production was observed following exposure of cells to 17 β -estradiol. The study
28 authors noted the similarity of effect between bisphenol A and 17 β -estradiol, which differed from the
29 modest effects observed with 4-*tert*-octylphenol exposure.

30
31 **Strengths/Weaknesses:** This study appears to have been well conducted. Phenol red-free media were
32 used and cell viability after treatment was assessed. There was likely limited metabolism of bisphenol A,
33 and the activity of metabolites cannot be assessed.

34
35 **Utility (Adequacy) for CERHR Evaluation Process:** These data indicate that 1–1000 nM bisphenol A
36 in vitro has no effect on basal hCG-induced testosterone production. However, because this study was
37 performed in vitro, it has limited utility.

38
39 **Akingbemi et al. (306)**, supported by NIEHS, US EPA, NICHHD, and NIH, conducted in vitro studies to
40 examine the effects of bisphenol A exposure on Leydig cell cultures. In vivo studies were also conducted
41 and are described in Section 3 because exposures were commenced in immature animals. In a series of
42 studies, testosterone production by Leydig cells was assessed following incubation of cells with various
43 doses of bisphenol A or bisphenol A in combination with other compounds. Leydig cells were obtained
44 from 90-day-old rats. In a dose-response study, testosterone and 17 β -estradiol levels were measured in
45 Leydig cells that were incubated with bisphenol A at 0 (ethanol vehicle), 0.01, 0.1, 1, 10, 100, or 1000
46 nM [0, 0.0023, 0.023, 0.23, 2.3, 23, and 230 $\mu\text{g/L}$] bisphenol A for 18 hours. To determine if bisphenol A
47 induces estrogenic effects on Leydig cells, testosterone production was also measured in cells incubated
48 with diethylstilbestrol or 2,2-bis(*p*-hydroxyphenyl)-1,1,1-trichloroethane, a metabolite of methoxychlor,
49 at the same concentrations as bisphenol A. In mechanistic studies, Leydig cells were incubated with 0.01
50 nM [0.0023 $\mu\text{g/L}$] bisphenol A, with and without the addition of LH or the antiestrogenic compound ICI
51 182,780. Endpoints assessed included testosterone and 17 β -estradiol production and expression of mRNA

4.0 Reproductive Toxicity Data

1 for steroidogenic metabolizing enzymes, ER, and steroidogenic acute regulatory protein, a substance that
2 transports the cholesterol used in testosterone synthesis. Levels of hormones in media were measured
3 using RIA methods, and mRNA expression was evaluated using RT-PCR techniques. Statistical analyses
4 included ANOVA and the Duncan multiple range test.

5
6 In the concentration-response study, production of testosterone by Leydig cells was decreased following
7 exposure to bisphenol A at 0.01 nM [**0.0023 µg/L**] but not at higher doses. Diethylstilbestrol reduced
8 testosterone production at all dose levels, and 2,2-bis(*p*-hydroxyphenyl)-1,1,1-trichloroethane reduced
9 testosterone production at concentrations ≥ 100 nM. Some statistically significant effects were observed in
10 the mechanistic studies in which cells were exposed to 0.01 nM bisphenol A. In one study, LH-stimulated
11 but not basal testosterone production was reduced by bisphenol A exposure. A second study demonstrated
12 a decrease in basal testosterone production following bisphenol A exposure, but no decrease in
13 testosterone level was observed following incubation of cells with bisphenol A in combination with ICI
14 182,270. 17β -Estradiol production was decreased in cells exposed to bisphenol A. Changes in mRNA
15 expression following bisphenol A exposure included reduced expression of mRNA for the steroidogenic
16 enzymes P45017 α -hydroxylase and aromatase. ER β was not detected in Leydig cells, and expression of
17 ER α mRNA was not affected. The study authors concluded that environmentally relevant concentrations
18 of bisphenol A act directly on Leydig cells to inhibit steroidogenesis, presumably via the ER.

19
20 **Strengths/Weaknesses:** This study appears to have been very well-conducted by a respected lab. The
21 study used a wide dose range and showed decreased testosterone production in in vitro Leydig cell
22 cultures at low (0.1 nM) but not at higher concentrations. The response of multiple endpoints provides
23 compelling evidence of a biological effect at 0.01 nM. An explanation for the selective effect of bisphenol
24 A at this single low concentration (0.1 nM) was not provided, nor was the dose range of this effect
25 explored.

26
27 **Utility (Adequacy) for CERHR Evaluation Process:** Although this study was performed in vitro (with
28 all the caveats), given the compelling effects in a multitude of endpoints examined, these data are highly
29 relevant and suggest the occurrence of selective low dose effects in the absence of high dose effects.

30
31 **Song et al. (450)**, supported by the Hormone Research Center and the Korean Andrological Society,
32 examined the role of bisphenol A in inducing expression of orphan nuclear receptor *Nur77*, a receptor that
33 plays an important role in the regulation of LH-induced steroidogenesis in Leydig cells. Methods used in
34 this study are described in conjunction with the results. **[It does not appear that statistical analyses
35 were conducted in this study.]** Following treatment of the mouse Leydig cell line K28 with bisphenol A
36 at ≥ 0.01 μ M, expression of *Nur77* mRNA was increased in a dose-related manner, with saturation of
37 expression observed at 1 μ M [**0.23 mg/L**]. In a time-response study with 1 μ M [**0.23 mg/L**] bisphenol A,
38 maximal expression of *Nur77* mRNA was observed at 30 minutes following treatment, basal levels of
39 expression were observed from 2 to 12 hours following treatment, and expression was again increased at
40 24 hours following treatment. When K28 cells were pretreated with the protein kinase inhibitor H89 or
41 the mitogen-activated protein kinase (MAPK) inhibitor PD98059, induction of *Nur77* mRNA by
42 bisphenol A was reduced by 40–45%. Induction of *c-fos* and *c-jun* mRNA occurred concurrently with
43 induction of *Nur77* mRNA. Bisphenol A-induced increases in *Nur77* promoter activity were greater
44 following transfection of cells with *Nur77* promoter reporter and *c-jun* but not with *c-fos*. Possible
45 activation of MAPK by bisphenol A was examined using an immunoblot method with an antibody
46 specific for phosphorylated MAPK. Phosphorylation of MAPK reached a maximum level at 10 minutes
47 following bisphenol A treatment. No changes in bisphenol A-induced induction of *Nur77* were observed
48 following pretreatment with a protein kinase C inhibitor or P13K inhibitor. The study authors stated that
49 together these results suggest possible involvement of the protein kinase A and MAPK pathways in
50 bisphenol A-induced induction of *Nur77*.

4.0 Reproductive Toxicity Data

1 In K28 cells transfected with Nur77 promoter or monomer binding site-luciferase reporters, gene
2 promoter activities and transactivation were increased following treatment with $\geq 0.1 \mu\text{M}$ [**0.023 mg/L**]
3 bisphenol A, thus suggesting similar responses between promoter activity and mRNA induction. In a
4 yeast assay, bisphenol A had no effect on interactions between *Nur77* and its corepressor, silencing
5 mediator of retinoid and thyroid receptor.

6
7 Exposure of K28 cells to $1 \mu\text{M}$ [**0.23 mg/L**] bisphenol A resulted in increased progesterone production,
8 which was inhibited 25% by the overexpression of dominant negative *Nur77*, which reduces the
9 transactivation activity of *Nur77*. Expression of mRNA for steroidogenic enzymes was investigated and it
10 was found that bisphenol A treatment increased expression of steroidogenic acute regulatory mRNA,
11 cholesterol side-chain cleavage enzyme, and 3β -hydroxysteroid dehydrogenase. Effects of bisphenol A on
12 expression of mRNA for *Nur77* and steroidogenesis enzymes was tested in prepubertal mice (18 days
13 old). Injection of 5 mice/group with 125 mg/kg bw/day bisphenol A resulted in increased expression of
14 *Nur77* mRNA and testosterone levels in mouse testis from 1–6 hours following exposure. [**Very few**
15 **details were provided for the in vivo experiment.**] The study authors concluded that the results of these
16 studies indicate that bisphenol A induces *Nur77* gene expression and alters steroidogenesis in Leydig
17 cells, indicating a possible novel mechanism of toxicity.

18
19 **Strengths/Weaknesses:** This study appears to have been well conducted and links in vitro bisphenol A
20 administration to dose-related (classic, not inverted) activation of *Nur77* and subsequent downstream
21 signal transducing proteins. Various confirmatory experiments supported this relationship. These data
22 strongly suggest that bisphenol A ($>0.1 \mu\text{M}$) activates *Nur77*. The toxicological implications of these
23 findings were not addressed.

24
25 **Utility (Adequacy) for CERHR Evaluation Process:** This study provides an additional potential
26 mechanism of action for bisphenol A; however, the link between these gene changes and adverse
27 outcomes is unclear. Moreover, further studies with other agents that activate *Nur77* are necessary to
28 understand the significance of these findings. This study is not useful in the evaluation.

29
30 **Hughes et al. (451)**, supported by the Medical Research Council, the British Heart Fund, and the
31 European Chemical Industry Council, examined the effects of bisphenol A on rat testicular calcium
32 pumps. Other phenolic compounds were examined, some in greater detail than bisphenol A, but this
33 discussion is limited to bisphenol A. Studies were conducted to determine the effects of bisphenol A
34 exposure on calcium ATPase pump activity, calcium uptake in testicular microsomes, calcium levels in
35 the TM4 Sertoli cell line, and TM4 cell viability. In the cell-viability study, cells were exposed to
36 bisphenol A at 0, 100, 300, or 600 μM [**0, 23, 68, or 137 mg/L**] for 16 hours. In each study, 2–12
37 samples/group were analyzed. [**For most studies, very few details were provided about procedures**
38 **such as exposure concentrations used and time that cells were incubated. There was no discussion**
39 **of statistical procedures, and it was not clear if statistical analyses were conducted for some**
40 **endpoints.**]

41
42 Bisphenol A inhibited calcium ATPase activity in rat testis microsomes. Mean \pm SEM median inhibitory
43 concentration (IC_{50}) values were reported at $0.40 \pm 0.15 \mu\text{M}$ [**91 \pm 34 $\mu\text{g/L}$**] for inhibition of calcium
44 ATPase activity and $2.5 \pm 1.0 \mu\text{M}$ [**571 \pm 228 $\mu\text{g/L}$**] for calcium uptake. Exposure to 200 μM [**47 mg/L**]
45 bisphenol A increased intracellular calcium levels in TM4 cells. A viability study was conducted to
46 determine if increased intracellular calcium levels resulted in cell death. Bisphenol A exposure resulted in
47 reduced TM4 cell viability (percent viability compared to control cells was 93, 64, and 17% at
48 concentrations of 100, 300, and 600 μM). The study authors concluded that these results provide evidence
49 that environmental estrogens may induce toxicity in male reproductive development by disrupting
50 calcium homeostasis.

4.0 Reproductive Toxicity Data

1 **Strengths/Weaknesses:** This interesting mechanistic study examined the role of bisphenol A in
2 modulating intracellular calcium levels. It is difficult to interpret the relationship between microsomal and
3 intact cell effects of bisphenol A given the large difference in concentrations needed to produce an effect.
4 Moreover, it is not clear if bisphenol A caused cytotoxicity by a calcium-dependent or non-calcium-
5 mediated process.

6
7 **Utility (Adequacy) for CERHR Evaluation Process:** This study is not useful in the evaluation.
8

9 **Tabuchi et al. (452)**, supported by the Japanese Ministry of Education, Culture, Sports, Science, and
10 Technology and Takeda Science Foundation, examined the effects of bisphenol A exposure on viability
11 and gene expression in TTE3 cells, a mouse Sertoli cell line. The cells were incubated for 24 hours in
12 media containing 0 or 24–400 μM [5.5–91 mg/L] bisphenol A (99.7% purity) in a DMSO vehicle. Cell
13 viability was determined, and gene expression changes were examined using microarray and PCR
14 techniques. Data were analyzed by Dunnett multiple comparison test or Student *t*-test. Compared to values
15 in control cells, bisphenol A exposure reduced cell viability by 25% at 100 μM [23 mg/L], 33% at 200
16 μM [46 mg/L], and 96% at 400 μM [91 mg/L]. Based on the results of the cell-viability studies, a
17 bisphenol A concentration of 200 μM [46 mg/L] was selected for the gene expression studies. Of 1081
18 genes examined by microarray, mRNA was downregulated in 3 cases and upregulated in 10 cases. Six
19 genes were selected for evaluation of mRNA expression by PCR, and of those genes, 1 was
20 downregulated (*ER α*) and 5 were upregulated (*iNOS*, *chop-10*, *odc*, *BipGRP78*, and *osip*). The study
21 authors concluded that microarray analysis is a useful tool for investigating molecular mechanisms of
22 bisphenol A-induced toxicity in testicular cells.

23
24 **Strengths/Weaknesses:** This interesting mechanistic study appears to have been well conducted, but it is
25 unclear from the data if bisphenol A-related changes in *chop-10* are a primary (or secondary) effect or are
26 the result of cytotoxicity.

27
28 **Utility (Adequacy) for CERHR Evaluation Process:** This study is not useful in the evaluation.
29

30 **Tabuchi and Kondo (453)**, supported by Japanese Ministry of Education, Culture, Sports, Science, and
31 Technology, Takeda Science Foundation, and Toyama Daiichi Bank Foundation, conducted a series of
32 experiments to examine the effects of in vitro bisphenol A exposure on gene expression in mouse Sertoli
33 cells. The experiments used TTE3 cells, an immortalized Sertoli cell line established from transgenic
34 mice expressing temperature-sensitive simian virus large T-antigen. Cells were exposed to bisphenol A
35 (99.7% purity) in a DMSO vehicle. The majority of experiments were repeated 2–4 times, and data were
36 analyzed by Student *t*-test. [Statistical significance was not reported in the results section of the
37 study.] Prior to conducting gene expression studies, cells were exposed to 25–400 μM [5.7–91 mg/L]
38 bisphenol A for 3–24 hours, and viability was determined using a tetrazolium compound. Cell viability
39 was reduced at bisphenol A concentrations \geq 200 μM [46 mg/L], and reductions in viability were
40 increased with longer durations of exposure. Intracellular calcium levels were measured using a
41 fluorescence imaging technique over a 15-minute period in cells exposed to 0–400 μM [0–91 mg/L]
42 bisphenol A, and a dose-related increase in calcium influx was observed at \geq 100 μM [23 mg/L]. Based on
43 results for cell viability and calcium influx studies, a concentration of 200 μM [46 mg/L] was selected for
44 the gene-expression experiments.

45
46 Using a PCR technique, it was determined that expression of mRNA for transferrin was decreased and
47 glucose-regulated protein mRNA was increased by bisphenol A exposure of up to 24 hours. Observations
48 of increased intracellular calcium concentration and upregulated glucose-regulated protein mRNA
49 expression led the study authors to conclude that bisphenol A stresses the endoplasmic reticulum. Gene
50 expression was analyzed by a cDNA microarray technique after exposure for 3, 6, 12, and 24 hours, and it
51 was determined that 31 of the 865 genes examined were upregulated by exposure to bisphenol A; no

4.0 Reproductive Toxicity Data

1 downregulation of genes was observed. The greatest change in gene expression was observed for *chop-*
2 *10*, a stress-response gene. Upregulation of 4 genes, *c-myc*, *fra-2*, *odc*, and *chop-10*, were confirmed by
3 quantitative PCR. *Chop-10* was determined to be the most responsive gene. To determine if *chop-10* was
4 required for development of endoplasmic reticulum stress and cell injury, a stably transfected cell line
5 expressing *chop-10* antisense RNA (*chopR14*) was developed. Mock cells were used as negative controls
6 in studies where cells were exposed to 200 μ M [46 mg/L] bisphenol A for up to 24 hours. Production of
7 *chop-10* protein, as determined by Western blot analysis, was reduced in the *chopR14* cells compared to
8 the mock cells following exposure to bisphenol A. In contrast to the mock cells, no reductions in cell
9 viability or transferrin mRNA expression were observed in the *chopR14* cells following bisphenol A
10 exposure. There were no changes in glucose-regulated protein mRNA expression in *chopR14* versus
11 mock cells. The study authors postulated that bisphenol A may disrupt the male reproductive system by
12 altering calcium homeostasis in Sertoli cell endoplasmic reticulum without interacting with the ER and
13 that genes such as *chop-10* may be involved in the process.

14
15 **Strengths/Weaknesses:** This mechanistic study appears to have been well conducted, but it is unclear
16 from the data if bisphenol A-related changes in *chop-10* are a primary (or secondary) effect or are the
17 result of cytotoxicity. Calcium levels were also affected and collectively these changes may be the result
18 of apoptosis initiated by some other mechanism.

19
20 **Utility (Adequacy) for the CERHR Evaluation Process:** This paper is not useful in the evaluation.

21
22 **Tabuchi et al. (454)**, supported in part by the Japanese Ministry of Education, Culture, Sports, Science,
23 and Technology, examined the effects of bisphenol A on gene expression in mouse Sertoli cell cultures.
24 TTE3 cells were incubated in media containing bisphenol A [**purity not reported**] at 0 (DMSO vehicle)
25 or 200 μ M [46 mg/L] for up to 12 hours. Cells were examined for viability using dye exclusion assays
26 and for apoptosis by formation of DNA ladders. RNA was extracted from cells, and gene expression was
27 determined by PCR and microarray analyses. Data were analyzed by Student *t*-test. Cell viability was
28 decreased in a time-related manner between 3 and 12 hours of bisphenol A exposure, but there was no
29 evidence of apoptosis. PCR analysis indicated that bisphenol A exposure significantly and time-
30 dependently increased mRNA transcripts for 2 endoplasmic reticulum stress markers, *hspa5* and *ddit3*.
31 Microarray analysis demonstrated that 661 sets of genes were downregulated and 604 sets of genes were
32 upregulated more than 2-fold following bisphenol A exposure. Pathway analysis of decreased gene
33 clusters revealed 2 significant genetic networks associated with the cell cycle or cell growth and
34 proliferation. In increased gene clusters, two genetic networks were associated with cell death, DNA
35 replication, recombination and repair, or injuries and abnormalities. The study authors concluded that the
36 genes, genetic clusters, and genetic networks identified in this study are likely involved in Sertoli cell
37 injury following bisphenol A exposure.

38
39 **Strengths/Weaknesses:** State-of-the-art technology was used in this study to examine gene expression
40 changes after in vitro bisphenol A exposure of a Sertoli cell line. Only one dose level was examined. The
41 use of hormone rich fetal bovine serum in the media may be a confounder. The absence of DNA
42 laddering is not conclusive evidence of the absence of apoptosis (e.g., adherent cells undergoing apoptosis
43 often are released into the culture media). Moreover, it is not surprising that given this “high” bisphenol A
44 concentration, “novel” and likely non-specific gene changes were noted.

45
46 **Utility (Adequacy) for CERHR Evaluation Process:** This study is not useful in the evaluation.

4.0 Reproductive Toxicity Data

4.2.3 Male and female

4.2.3.1 Rat

Emm et al. (292), supported by the Japanese Ministry of Health and Welfare, conducted a multigeneration reproductive toxicity study of bisphenol A in CD rats. Animals were housed in suspended stainless steel cages at the beginning of the study. From GD 17, wood chips were used as bedding. Rats were fed CRF-1 chow (Oriental Yeast Co). In the study that was conducted according to GLP, F₀ male rats and female rats with 4–5-day estrous cycles were randomly assigned to groups of 25/sex. Five-week-old males and 10-week-old females were gavaged with 0 (distilled water vehicle), 0.0002, 0.002, 0.020, or 0.200 mg/kg bw/day bisphenol A (99.9% purity). Males were dosed for 10 weeks prior to mating and during the mating period, which lasted up to 2 weeks. Females were dosed from 2 weeks prior to mating, and during the mating, gestation, and lactation periods. Doses were based on results of studies by Nagel et al. (205) and vom Saal et al. (341). Stability and concentration of dosing solutions were verified. Dams delivered and nursed their pups. At weaning on PND 22 (day of birth defined as PND 0), 1 or 2 F₁ weanlings/litter/sex (25/sex/group) were selected to continue in the study. Dosing of F₁ animals began on PND 23 and continued for 10 weeks prior to mating and through the mating period, which lasted up to 3 weeks. Dosing was continued through the gestation and lactation periods. Twenty-five F₂ weanlings/sex/group were selected on PND 22. Beginning on PND 22, male F₂ rats were dosed for 4 weeks and females were dosed for 11 weeks prior to being killed.

Endpoints examined in adult rats included clinical signs, body weight, and food intake. Fertility, copulation, and gestational indices were examined in mating rats. Vaginal smears were evaluated for two weeks prior to mating in F₀ and F₁ females and at 9–11 weeks of age in F₂ females. Dams were killed and necropsied following weaning of their pups, and uterine implantation sites were examined. Males were killed following mating. Organs were weighed and histopathology examinations were conducted in control and high-dose animals. Sperm endpoints were measured in F₀ and F₁ adult males. Serum hormone levels were measured in 6 adult F₀ and F₁ males and proestrous females. At birth, pups were counted, sexed, and examined for viability and external malformations. On PND 4, litters were culled to 4 male and 4 female pups. At weaning, 1 male and female F₁ and F₂ weanling was killed for organ weight measurement; histopathology exams were conducted in seminal vesicles and coagulating glands of F₂ weanlings. Survival and growth were monitored during the postnatal period. Pups were examined for developmental landmarks and attainment of vaginal opening or preputial separation. Anogenital distance in pups was examined at numerous time points during the lactation period and through adulthood. Behavioral testing was conducted at 5–7 weeks of age. The litter was considered the experimental unit in data obtained prior to weaning. Statistical analyses included Bartlett test for homogeneity of variance, ANOVA, and/or Dunnett multiple comparison, Kruskal-Wallis, Mann-Whitney *U*, chi-squared, or Fisher exact tests.

In F₀ and F₁ adult animals, there were no treatment-related effects on clinical signs, body weight gain, or death. The only significant reproductive effects reported in adult animals were non-dose-related decreases in percentages of females with normal estrous cycles (76 versus 96% in controls) and reduced gestation duration (by 0.5 days) in the F₁ group treated with 0.020 mg/kg bw/day. Bisphenol A did not significantly affect the precoital interval, copulation index, fertility index, gestation index, number of implantations, or delivery index. There were no adverse effects on sperm endpoints such as count, motility, or morphology in F₀ or F₁ males. A significant decrease in abnormal and tailless sperm was observed in F₁ males of the 0.020 mg/kg bw/day group. There was no evidence of histopathological effects in reproductive organs of F₀ animals that did not copulate or had totally resorbed litters or in F₁ animals of the high-dose group. **[Data were not shown by study authors.]** In F₀ females, there were significant decreases in serum LH concentrations at 0.0002, 0.002, and 0.020 mg/kg bw/day and in serum triiodothyronine levels at 0.200 mg/kg bw/day. **[Data were not shown by study authors.]** Organ weight changes in F₁ adult males included decreased absolute weights of lung at 0.0002 and 0.200 mg/kg bw/day, kidney at 0.2 mg/kg

4.0 Reproductive Toxicity Data

1 bw/day, and testis at 0.020 mg/kg bw/day. Absolute ovarian weight was decreased in females of the
2 0.0002 mg/kg bw/day group. Seminal vesicle weight was decreased in F₂ males of the 0.200 mg/kg
3 bw/day group. **[Data were not shown by study authors].**
4

5 There were no significant effects on number of F₁ or F₂ pups delivered, sex ratio, or pup survival during
6 the lactation period. Body weights of F₁ pups in the 0.020 mg/kg bw/day group were significantly lower
7 **[by 6–7%]** on PND 14 and 21. Testicular descent was delayed by 0.7 days in F₂ offspring from the 0.020
8 and 0.200 mg/kg bw/day groups. There were no significant effects on age of pinna detachment, incisor
9 eruption, or eye opening. Some significant but non-dose-related effects on reflex development were
10 observed. Day of mid-air righting reflex was accelerated by 1.2 days in F₁ males and 1.5 days in F₁
11 females of the 0.020 mg/kg bw/day group. In F₂ males, negative geotaxis was delayed by 0.8 days at
12 0.0002 mg/kg bw/day, 0.5 days at 0.002 mg/kg bw/day, and 0.8 days at 0.020 mg/kg bw/day. Bisphenol
13 A treatment did not significantly affect age of vaginal opening or preputial separation in F₁ or F₂
14 offspring. Some sporadic and small (within 5% of control values) changes in anogenital distance were
15 observed in F₁ and F₂ offspring. In F₁ males, decreased anogenital distance was observed in the 0.0002
16 mg/kg bw/day group on PND 57 and in the 0.020 mg/kg bw/day group on PND 106, 113, and on the day
17 of sacrifice. In F₁ females, anogenital distance was decreased in the 0.200 mg/kg bw/day group on PND 4
18 and increased in the 0.002 and 0.020 mg/kg bw/day group on PND 7. Decreases in anogenital distance of
19 F₂ females were observed in the 0.020 mg/kg bw/day group on PND 64, 71, 85, 92, and on the day of
20 sacrifice and in the 0.200 mg/kg bw/day group on PND 57, 64, and on the day of sacrifice. In F₁
21 offspring, there were no significant effects on behavior, as determined by open-field testing and
22 performance in a T-maze. **[Data were not shown by study authors.]** There was no evidence of
23 histopathological effects in seminal vesicle or coagulating gland of F₂ pups from the high-dose group.
24 **[Data were not shown by study authors.]** Organ weight changes in F₁ male weanlings included
25 decreased absolute lung weight at 0.020 and 0.200 mg/kg bw/day group and decreased kidney weight at
26 0.020 mg/kg bw/day. In male F₂ weanlings, significant decreases were observed in absolute and relative
27 seminal vesicle weight and absolute thyroid weight at 0.002 mg/kg bw/day, absolute lung weight at 0.020
28 mg/kg bw/day, and relative heart weight at 0.200 mg/kg bw/day; relative liver weight was significantly
29 increased in F₂ males of the 0.002 mg/kg bw/day group. The study authors concluded that oral
30 administration of bisphenol A at 0.0002 to 0.200 mg/kg bw/day to 2 generations of rats did not cause
31 changes in reproduction or development.
32

33 [The NTP Statistics Subpanel (295) reviewed an unpublished study that appeared to be the same study
34 later published as Ema et al. (292). The subpanel noted that in general they agreed with the statistical
35 methodology used in the study but stated that the Dunnett test does not require significance of ANOVA.
36 It was noted that the anogenital distance findings were the most difficult to interpret. The Subpanel noted
37 that many of the anogenital distance effects remained statistically significant when analyzed by
38 ANCOVA, a method they considered superior to adjustment by body weight. The NTP Subpanel agreed
39 with the author's conclusion that effects on anogenital distance were not biologically significant. They
40 noted an error in the unpublished study abstract that described increases in anogenital distance in F₁ and
41 F₂ females in the 0.020 and 0.2 mg/kg bw/day groups when actually the effect should have been
42 decreased anogenital distance. **[It was not clear to CERHR if this error was carried forward to the
43 published report.]**
44

45 **Strengths/Weaknesses:** This well-designed comprehensive low-dose assessment of potential bisphenol
46 A-related effects on multiple generations of rats examined a wide variety of hormonally sensitive
47 endpoints. The study had appropriate power with an appropriate number of rats per group. Route of
48 administration (oral) was appropriate. The concentrations of the dosing solutions were verified (both prior
49 and after). It would have been helpful if a dose level that caused maternal toxicity was also used;
50 however, given the objective of this study it is a minor point.
51

4.0 Reproductive Toxicity Data

1 **Utility (Adequacy) for CERHR Evaluation Process:** This thorough multiple generation rat study is
2 highly valuable for human risk assessment of low dose oral exposure to bisphenol A. This study indicates
3 that the NOAEL for bisphenol A exceeds 0.2 mg/kg bw/day under the conditions of this study.
4

5 **Tyl et al. (293, 411)**, sponsored by The Society of the Plastics Industry, Inc., conducted a multigeneration
6 study of bisphenol A in rats. In the study that was conducted according to GLP, Sprague Dawley rats
7 were fed Purina Certified Rodent Chow® 5002. F₀ rats (30/sex/group) were exposed to bisphenol A
8 (99.5% purity) in feed for 10 weeks prior to mating. **[Age at start of exposure was not reported, but**
9 **based on information provided in the discussion, it appears that the animals were adults at the start**
10 **of exposure.]** Vaginal smears were evaluated during the last 3 weeks of the prebreeding period. Exposure
11 continued through a 2-week mating period. Males were exposed an additional 3 weeks following mating,
12 and females were exposed through gestation and lactation. Concentrations of bisphenol A added to feed
13 were 0, 0.015, 0.3, 4.5, 75, 750, or 7500 ppm. Target intakes were ~0, 0.0009, 0.018, 0.27, 4.5, 45, and
14 450 mg/kg bw/day in males and 0.001, 0.02, 0.30, 5, 50, and 500 mg/kg bw/day in females. Actual
15 intakes were 0.0007–0.003, 0.015–0.062, 0.22–0.73, 4.1–15.4, 37.6–167.2, and 434–1823 mg/kg bw/day.
16 The study was designed to include low-dose exposures reported to increase prostate weights (205, 455)
17 and maximally tolerated doses expected to result in toxicity. Concentration, stability, and homogeneity of
18 bisphenol A in feed were verified. During the study, body weight and food intake were measured and
19 animals were examined for clinical signs. F₀ males were killed and necropsied following delivery of the
20 F₁ litter. Histopathological evaluation of organs was conducted in all control animals and 10
21 animals/bisphenol A dose group. Reproductive organs were weighed and sperm endpoints were
22 evaluated. F₀ females were killed and necropsied following weaning of their litters. Selected organs were
23 weighed and ovarian primordial follicles were counted.
24

25 On PND 4, F₁ litters were culled to 10 pups, with equal numbers of each sex when possible. Endpoints
26 examined in pups included growth and survival in the prenatal period and retained areolae or nipples on
27 PND 11–13. At weaning on PND 21, 30 F₁ offspring/sex/group were randomly selected and exposed to
28 bisphenol A in the diet according to the same protocol as F₀ rats. Those selected offspring were monitored
29 for vaginal opening and preputial separation and later mated. Up to 3 F₁ weanlings/sex/litter were killed
30 for organ weight measurement. Mating and evaluation of F₁ offspring were conducted according to the
31 same procedures described for F₀ rats. The same procedures were repeated in F₂ rats and F₃ litters during
32 the lactation period. Anogenital distance was measured in F₂ and F₃ rats at birth. Following weaning of F₃
33 offspring, up to 3/sex/litter were randomly selected for necropsy. Thirty/sex/dose were selected for
34 evaluation of vaginal patency, preputial separation, and estrous cyclicity. Bisphenol A exposure was
35 continued in those offspring until they were killed ~10 weeks following weaning. F₃ offspring were not
36 mated, but necropsy evaluations were conducted as described above for previous generations.
37

38 Statistical analyses for quantitative continuous data included Bartlett test for homogeneity of variances,
39 ANOVA, Dunnett, linear trend, Kruskal-Wallis, or Mann-Whitney *U* tests. Frequency data were analyzed
40 by chi-squared, Fisher exact, and Cochran-Armitage tests. Covariance and correlations analyses were also
41 conducted.
42

43 Treatment-related systemic findings with available quantitative information in adult rats are summarized
44 in Table 108. Body weights and body weight gain were consistently lower in F₀, F₁, F₂, and F₃ adult rats
45 of the 750 and 7500 ppm dose groups, including during gestation and lactation periods. Terminal body
46 weight effects are summarized in Table 108. Terminal body weight was reduced in all generations at 7500
47 ppm and in F₁ females and F₁ and F₂ males at 750 ppm. There were no consistent or clearly treatment-
48 related effects on feed intake. No treatment-related clinical signs were reported. In the 7500 ppm group,
49 absolute weights of the liver in males and the kidney in both sexes were decreased across generations.
50 Relative weights were either increased or did not attain statistical significance. **[According to Table 2 of**
51 **the study, absolute liver weights were also decreased in males of the 750 ppm group. The study**

4.0 Reproductive Toxicity Data

1 **authors also mentioned reductions in weights of adrenal glands, spleen, pituitary, and brain at the**
2 **high dose, but there were no data shown in the report for those endpoints.]** Other changes in
3 nonreproductive organ weight occurred sporadically at lower dose and were not dose-related or consistent
4 across generations. Relative organ weight changes that consistently attained statistical significance at the
5 highest dose are summarized in Table 108. Histopathological analyses revealed a higher incidence of mild
6 renal tubular degeneration and chronic hepatic inflammation in F₀, F₁, and F₂ but not F₃ females of the
7 7500 ppm group.

8
9 Treatment-related effects on reproductive endpoints in adult animals are summarized in Table 108. In
10 evaluating organ weights, the study authors only considered organ weight effects to be biologically
11 significant if statistically significant results were obtained in the same direction for absolute and relative
12 weights. Therefore, the study authors concluded that the only treatment-related organ weight effects were
13 reduced absolute and relative ovary weights. **[Numerous statistically significant effects on**
14 **reproductive organ weights were reported in Table 2 of the study. Reductions in testes,**
15 **epididymides, prostate, and seminal vesicle weights were observed in most generations of the 7500**
16 **ppm group. When adjusted for body weight, organ weights were either increased or did not differ**
17 **significantly from controls.]** Relative reproductive organ weight changes that consistently attained
18 statistical significance at the highest dose are summarized in Table 108. The authors reported no effect on
19 mating, fertility, pregnancy, or gestational indices. **[With the exception of gestational length, data**
20 **were not shown by study authors.]** Precoital interval, postimplantation loss, estrous cyclicality, and
21 reproductive organ histopathology were also unaffected by bisphenol A treatment. In the high-dose group,
22 there was no adverse effect on paired ovarian primordial follicle counts but counts were significantly
23 increased by 43% in the F₀ generation. Implantation sites were decreased in F₀, F₁, and F₂ dams of the
24 7500 ppm group. The only significant effects on sperm endpoints were decreased epididymal sperm
25 concentration in F₁ males and decreased daily sperm production in F₃ males of the 7500 ppm dose group.
26 There were no effects on sperm morphology or motility. The study authors considered sperm to be
27 unaffected by treatment.

28
29 Treatment-related effects observed in developing rats are summarized in Table 109. The number of live
30 pups/litter was reduced in F₁, F₂, and F₃ litters of the 7500 ppm group. Body weights of F₁, F₂, and F₃
31 pups of the 7500 mg/kg bw/day groups were lower during the lactation period. Some small (~5%)
32 decreases in pup body weight during the lactation period at lower doses were apparently not considered
33 treatment-related by study authors. Postnatal survival was unaffected by bisphenol A treatment. In male
34 rats, there were no effects on anogenital distance or the presence of areolas or nipples. Anogenital
35 distance was significantly increased in F₂ females at all doses except 75 and 7500 ppm; there was no
36 affect on anogenital distance in F₃ females. The study authors did not consider anogenital distance effects
37 to be biologically or toxicologically significant. Vaginal patency was delayed in F₁, F₂, and F₃ females,
38 and the effect remained significant following adjustment for body weight. Preputial separation was
39 delayed in F₁ males of the 750 and 7500 ppm groups, F₂ males in the 0.3, 75, 750, and 7500 ppm groups,
40 and F₃ males of the 7500 ppm group. When adjusted for body weight, the effect remained significant in F₁
41 males of the 750 and 7500 ppm groups and F₂ and F₃ males of the 7500 ppm group. The study authors
42 stated that reduced body weights were the most likely cause of puberty delay in males and females. **[In**
43 **rats killed at weanling, absolute organ weights were said to be decreased at the high dose but**
44 **increased when adjusted for body weight. The specific organs affected were not reported and no**
45 **data were presented. The exception was ovarian weights, which were reported to parallel effects**
46 **observed in adult females with decreases in both absolute and relative weight at 7500 ppm.]**

47
48 The study authors concluded that there was no evidence of low-dose bisphenol effects (1 µg to 5 mg/kg
49 bw/day) at any stage of the life cycle. They identified NOAELs of 75 ppm (~5 mg/kg bw/day) for adult
50 systemic toxicity and 750 ppm (~50 mg/kg bw/day) for offspring and reproductive effects. The study
51 authors concluded that bisphenol A should not be considered a selective reproductive toxicant.

4.0 Reproductive Toxicity Data

1 **Table 108. Treatment-related Effects in Adult Rats Fed Bisphenol A Through Diet in a Multigeneration Reproductive Toxicity**
 2 **Study**

Endpoint	Dose, ppm diet [mg/kg bw/day ^a]						BMD ₁₀	BMDL ₁₀	BMD _{1SD}	BMDL _{1SD}	
	0.015 [0.0095]	0.3 [0.019]	4.5 [0.285]	75 [4.75]	750[47.5]	7500 [475]					
Terminal body weight											
F ₀ males	↔	↔	↔	↔	↔	↔	↓22%	3554 [225]	3137 [199]	3133 [198]	2701 [171]
F ₁ males	↔	↔	↔	↔	↔	↔	↓6%	2811 [178]	2548 [161]	2443 [155]	2153 [136]
F ₂ males ^b	↔	↔	↔	↔	↔	↔	↓12%	733 [46]	554 [35]	648 [41]	484 [31]
F ₃ males ^b	↔	↔	↔	↔	↔	↔	↓26%	1456 [92]	913 [58]	1260 [80]	786 [50]
F ₀ females	↔	↔	↔	↔	↔	↔	↓13%	5722 [362]	4753 [301]	4741 [300]	3876 [245]
F ₁ females	↔	↔	↔	↔	↔	↔	↓6%	4600 [291]	3950 [250]	3730 [236]	3142 [199]
F ₂ females ^b	↔	↔	↔	↔	↔	↔	↓14%	3863 [245]	1576 [100]	3115 [197]	1291 [82]
F ₃ females	↔	↔	↔	↔	↔	↔	↓20%	3664 [232]	3194 [202]	3456 [219]	2949 [187]
Relative paired kidney weight											
F ₀ males	↔	↔	↔	↔	↔	↔	↑14%	5903 [374]	4555 [288]	6536 [414]	5035 [319]
F ₁ males	↔	↔	↓5%	↔	↔	↔	↑10%	5729 [363]	4662 [295]	5053 [320]	4088 [259]
F ₂ males	↔	↔	↔	↔	↔	↔	↑5%	4524 [287]	3893 [247]	3471 [220]	2950 [187]
F ₃ males	↔	↔	↔	↔	↔	↔	↑16%	6986 [442]	4319 [274]	6720 [426]	3403 [216]
F ₀ females	↔	↔	↔	↔	↔	↔	↑7%	8008 [507]	7521 [476]	7712 [488]	6578 [417]
F ₂ females	↔	↔	↔	↔	↔	↔	↑6%	7930 [502]	7515 [476]	7621 [483]	6247 [396]
Relative paired testis weight											
F ₀ males	↔	↔	↔	↔	↔	↔	↑27%	2924 [185]	2567 [163]	2998 [190]	2596 [164]
F ₁ males	↔	↔	↔	↔	↔	↔	↑18%	3287 [208]	2763 [175]	4106 [260]	3428 [217]
F ₂ males	↔	↔	↔	↔	↔	↔	↑24%	3086 [195]	2874 [182]	3245 [206]	2779 [176]
F ₃ males	↔	↔	↔	↔	↔	↔	↑19%	4329 [274]	2593 [164]	5010 [317]	3298 [209]
Relative paired epididymis weight											
F ₀ males	↔	↔	↔	↔	↔	↔	↑19%	3804 [241]	3072 [195]	5044 [319]	4068 [258]
F ₁ males	↔	↔	↔	↔	↔	↔	↑19%	2963 [188]	2566 [163]	3255 [206]	2786 [17]
F ₂ males ^b	↔	↔	↔	↔	↔	↔	↑8%	884 [56]	596 [38]	951 [60]	641 [41]
F ₃ males	↔	↔	↔	↔	↔	↔	↑22%	3449 [218]	2516 [159]	4117 [261]	3095 [196]
Relative liver weight											
F ₀ females	↔	↔	↔	↔	↔	↔	↑11%	7663 [485]	5848 [370]	7965 [504]	7439 [471]
F ₂ females	↑	↔	↔	↔	↔	↔	↑19%	6912 [438]	3650 [231]	7454 [472]	5533 [350]
Relative paired ovary weight]											
F ₀ females	↔	↔	↔	↔	↔	↔	↓19%	4103 [260]	3149 [199]	7126 [451]	5387 [341]
F ₁ females	↔	↔	↔	↔	↔	↔	↓15%	5754 [364]	3964 [251]	10,237 [648]	6966 [441]

4.0 Reproductive Toxicity Data

Endpoint	Dose, ppm diet [mg/kg bw/day] ^a						BMD ₁₀	BMDL ₁₀	BMD _{1SD}	BMDL _{1SD}
	0.015 [0.0095]	0.3 [0.019]	4.5 [0.285]	75 [4.75]	750[47.5]	7500 [475]				
F ₂ females	↓15%	↔	↓15%	↓11%	↔	↓24%	7053 [447]	3520 [223]	7646 [484]	6360 [403]
Number with renal tubule degeneration										
F ₀ females	0/12	0/12	0/12	0/14	0/12	4/13	6491 [411]	3848 [244]		
F ₁ females	0/10	0/10	0/10	0/10	0/10	8/11	5498 [348]	2470 [156]		
F ₂ females	0/11	0/10	0/12	0/11	0/12	7/13	5884 [373]	3018 [191]		
Number females with chronic liver inflammation										
F ₀ females	0/12	1/12	0/12	0/14	1/12	3/13	4867 [308]	3214 [204]		
F ₁ females	0/10	0/10	3/10	1/10	1/10	3/11				
F ₂ females	1/11	0/10	2/12	2/11	2/12	5/13	3029 [192]	1856 [118]		
Number of implantation sites										
F ₀ dams	↔	↔	↔	↔	↔	↓16%	4088 [259]	3021 [191]	8020 [508]	5832 [369]
F ₁ dams ^b	↔	↔	↔	↔	↔	↓26%	6120 [388]	2383 [151]	7000 [443]	4713 [298]
F ₂ dams	↔	↓8%	↔	↔	↔	↓18%	4917 [311]	3597 [228]	7679 [486]	5631 [357]
Epididymal sperm concentration, F ₁	↔	↔	↔	↔	↔	↓18%	5012 [317]	3407 [216]	11,050 [700]	7407 [469]
Daily sperm production, F ₃	↔	↔	↔	↔	↔	↓19%	7399 [469]	4025 [255]	8279 [524]	7596 [481]

↑,↓ Statistically significant increase, decrease, ↔ no statistically significant effect.

^aBased on target doses provided by the study authors and expressed as an average of the dose for males and females.

^bBenchmark dose values were estimated using a polynomial model.

From Tyl et al. (293).

1

4.0 Reproductive Toxicity Data

1 Table 109. Treatment-related Effects in Developing Rats in a Multigeneration Reproductive Toxicity Study of Bisphenol A

Endpoint	Dose, ppm diet [mg/kg bw/day ^a]						BMD ₁₀	BMDL ₁₀	BMD _{1SD}	BMDL _{1SD}	
	0.015 [0.0095]	0.3 [0.019]	4.5 [0.285]	75 [4.75]	750[47.5]	7500 [475]					
Live pups/litter											
F ₁	↔	↔	↔	↔	↔	↔	↓20%	4232 [268]	3033 [192]	8823 [559]	6225 [394]
F ₂	↔	↔	↔	↔	↔	↔	↓26%	6661 [422]	2405 [152]	7241 [459]	4645 [294]
F ₃	↔	↓11%	↔	↔	↔	↔	↓26%	3733 [236]	2742 [174]	5943 [376]	4518 [286]
Pup body weight											
F ₁ , PND 4	↔	↔	↔	↔	↔	↔	↓11%	6412 [406]	4473 [283]	8860 [561]	6317 [400]
F ₁ , PND 7	↔	↔	↔	↔	↔	↔	↓23%	3432 [217]	2891 [183]	4179 [265]	3448 [218]
F ₂ , PND 7	↔	↔	↔	↔	↔	↔	↓15%	5179 [328]	4059 [257]	6023 [381]	4653 [295]
F ₃ , PND 7	↔	↔	↔	↔	↔	↔	↓13%	4976 [315]	3854 [244]	6474 [410]	4940 [313]
F ₁ , PND 14	↔	↔	↔	↔	↔	↔	↓27%	2890 [183]	2570 [163]	2789 [177]	2415 [153]
F ₂ , PND 14	↔	↔	↔	↔	↔	↔	↓20%	3840 [243]	3302 [209]	3579 [227]	3013 [191]
F ₃ , PND 14	↔	↔	↔	↔	↔	↔	↓20%	3704 [235]	3224 [204]	3323 [210]	2827 [179]
F ₁ , PND 21	↔ ^b	↔	↔	↔ ^b	↔ ^b	↔ ^b	↓27%	3284 [208]	2621 [166]	3523 [223]	2763 [175]
F ₂ , PND 21	↔ ^b	↔	↔	↔ ^b	↔ ^b	↔ ^b	↓20%	4253 [269]	3566 [226]	4219 [267]	3473 [220]
F ₃ , PND 21	↔ ^b	↔	↔	↔ ^b	↔ ^b	↔ ^b	↓19%	3972 [252]	3423 [217]	3575 [226]	3016 [191]
Anogenital distance, F ₂ females	↑3%	↑3%	↑3%	↔	↔	↔	↑4%	↔	↔	↔	↔
Age of vaginal opening adjusted for body weight											
F ₁	↔	↔	↔	↔	↔	↔	↑3.6 days	6225 [394]	5422 [343]	3248 [206]	2786 [176]
F ₂	↔	↔	↔	↔	↔	↔	↑4 days	6381 [404]	5307 [336]	4367 [277]	3600 [228]
F ₃	↔	↔	↔	↔	↔	↔	↑3.2 days	7444 [471]	6325 [401]	6249 [396]	3198 [203]
Age of preputial separation adjusted for body weight											
F ₁	↔	↔	↔	↔	↔	↔	↑1.7 days	7350 [466]	6485 [411]	2974 [188]	2580 [163]
F ₂	↔	↔	↔	↔	↔	↔	↑7.4 days	4740 [300]	4025 [255]	3809 [241]	3201 [203]
F ₃	↔	↔	↔	↔	↔	↔	↑4 days	8637 [547]	7466 [473]	3503 [222]	2984 [189]

↑,↓ Statistically significant increase, decrease, ↔ no statistically significant effect.

^aBased on target doses provided by the study authors and expressed as an average of the dose for males and females.

^bA significant (~5%) decrease in pup body weights observed only in F₁ and/or F₂ litters was apparently not considered treatment-related by study authors. From Tyl et al. (293).

1
2 [The NTP Statistics Subpanel (295) stated that the study by Tyl et al. (411) apparently lacked a
3 check for outliers, but noted that the study was in draft form at the time of review. The NTP
4 subpanel agreed with most author conclusions but disagreed with a conclusion that relative uterine
5 weights were equivalent across all groups. The unnecessary use of ANOVA with Dunnett test was
6 noted. Some possible outliers and 10-fold errors in data points that could have affected conclusions
7 were observed. Overall, the NTP Subpanel concluded that Tyl et al. (411) study was the most
8 comprehensive of the studies reviewed. They stated that the statistical methods were well thought
9 out and appropriate.]

10
11 **Strengths/Weaknesses:** This assessment of potential bisphenol A-related effects on multiple generations
12 of rats was well-designed and comprehensive. The large number of rats/group (30), the multiple endpoints
13 examined, and the oral route of administration (diet) are strengths. The concentration of bisphenol A in
14 the test diet was verified, and maternal and paternal toxicity was identified. This study explored a wide
15 dose range and demonstrates an absence of adverse effects on reproductive function at very low bisphenol
16 A dose levels.

17
18 **Utility (Adequacy) for CERHR Evaluation Process:** This study is highly valuable for human risk
19 assessment for oral exposure to bisphenol A. This study identified a NOAEL of 75 ppm (for general
20 toxicity) and 750 ppm (for reproductive toxicity).

21 22 4.2.3.2 Mouse

23 **NTP (456, 457)** sponsored a continuous breeding study in CD-1 mice exposed to bisphenol A through sc
24 implants. Mice were fed Purina certified ground rodent chow (#5002) and housed in polypropylene or
25 polycarbonate cages containing Ab-Sorb-Dri bedding. Silastic implants were used for sc dosing of mice
26 with bisphenol A (~95% purity) in corn oil vehicle. Stability and weight of bisphenol A in pumps was
27 verified. In the dose-range finding portion of the study (Task 1), 8 mice/sex/group (8 weeks old) received
28 implants containing vehicle or bisphenol A. Dosages were estimated by determining the difference in
29 bisphenol A weight at the start and end of the 14-day dosing period. It was estimated that mice received 0,
30 6.25, 12.5, 25, 50, or 100 mg bisphenol A. Endpoints examined included body weight changes, survival,
31 and uterine weight. Blood was collected to determine plasma bisphenol A levels. Data were analyzed by
32 ANOVA, Duncan Multiple Range Test, chi-squared test, and Fisher exact test. The goal of Task 2 was to
33 determine a maximum tolerated dose that produced signs of toxicity but did not reduce body weight or
34 increase lethality by more than 10% and to identify a low dose that did not result in toxicity.
35 Concentrations of bisphenol A in plasma were below the detection limit (3 ng/mL) in the 6.25 mg group
36 but were reported at 7.0–7.7 µg/L in the 12.5 mg group, 8.4 µg/L in the 25 mg group, 13.1–18.5 µg/L in
37 the 50 mg group, and 31.5–56.2 µg/L in the 100 mg group. In mice treated with bisphenol A, there were
38 no increases in death or effects on body weight gain. The study authors noted that reproductive tract
39 weight in the high dose group was greater [by 52%] than in the control group but statistical significance
40 was not achieved because of high variability.

41
42 In the continuous breeding portion of the study (Task 2), mice were 11 weeks old at the start of dosing.
43 Forty mice/sex/group received implants containing the vehicle and 20/sex/dose received implants
44 containing bisphenol A at 25, 50, or 100 mg. Over a dosing period of 18 weeks, it was estimated that
45 animals in each treatment group received 11.65, 20.05, and 38.60 mg bisphenol A. [Assuming body
46 weights of ~38 g, as indicated in the study report, doses would have been ~306, 527, and 1015 mg/kg
47 bw over 18 weeks or 2.4, 4.2, and 8.1 mg/kg bw/day.] Mice were eleven weeks old at the start of
48 dosing, which began during a 7-day pre-mating period. The mice were then randomly paired with animals
49 from the same dose group and housed together during a 98-day breeding period. Litters born during the
50 breeding period were examined for viability, weighed, sexed, and discarded. Following the 98-day mating
51 period, mice were separated for 21 days to allow for the birth of the last litter. Dosing was continued

4.0 Reproductive Toxicity Data

1 throughout the breeding and separation periods. However implants were often expelled through cutaneous
2 lesions or the incision site. When animals expelled their implant, a new one was inserted but pregnant
3 mice were allowed to complete their pregnancy before insertion of the new implant. Therefore dosing was
4 not uniform. Endpoints examined in adult mice included body weight, number of litters/pair, and fertility.
5 Following delivery of the final litter, parental animals were killed and animals in the 0 and 100 mg group
6 were necropsied. Liver, brain, and reproductive organs were weighed. Data were analyzed by chi-squared
7 test, Fisher exact test, Kruskal-Wallis test, Jonckheere test, and Mann-Whitney *U* test.

8
9 With the exception of cutaneous lesions at the implantation site, there were no clinical signs of toxicity. In
10 parental mice, there were no effects on body weight, mortality, fertility, or number of litters born. There
11 were no changes in weights of organs including, liver brain, pituitary, the female reproductive tract, testis,
12 epididymis, prostate, or seminal vesicles. Statistically significant effects observed in pups included
13 increased numbers of live male and total pups and increased adjusted (for litter size) pup weight in the
14 mid-dose group. Unadjusted and adjusted male and female pup weights were significantly increased at the
15 high dose. The study authors noted that the effects observed in this study were random and most likely
16 due to chance. They concluded that bisphenol A did not induce adverse effects on fertility in male or
17 female mice. It was noted that further studies using a better route of exposure are needed for bisphenol A.

18
19 **Strengths/Weaknesses:** This study appears to have been well conducted. When compared to studies that
20 used the oral route of exposure, this study provides evidence that the manifestation of maternal toxicity is
21 dependent on the route of administration and that route-dependent metabolism may be important for
22 toxicity. However, the administration of bisphenol A via silastic implants makes the extrapolation for
23 human risk assessment difficult in the absence of an improved pharmacokinetic understanding.

24
25 **Utility (Adequacy) of CERHR Evaluation Process:** This comprehensive study used an irrelevant route
26 of exposure, which makes extrapolation for human risk assessment difficult. The NOAEL for
27 reproductive effects exceeds 8 mg/kg bw.day.

28
29 **NTP (457, 458)** sponsored a continuous breeding study in CD-1 mice exposed to bisphenol A (98%
30 purity). Additional information on ovarian follicle counts in F₀ and F₁ females was published in a report
31 by Bolon et al. (459). In this study, mice were fed NIH-07 open formula rodent chow and housed in
32 polypropylene or polycarbonate cages containing Ab-Sorb-Dri litter. The laboratory at which the study
33 was conducted was stated to be in full compliance with GLP regulations. In the preliminary study (Task
34 1), 8 mice/sex/group (8 weeks old) were fed diet containing bisphenol A at 0, 0.3125, 0.625, 1.25, 2.5, or
35 5.0% for 14 days. By assuming that a 40 g mouse ingests 7 g feed/day, the study authors estimated
36 bisphenol intake at 0, 437.5, 875.0, 1750.0, 4375.0, 8750.0 mg/kg bw/day. The aim of the preliminary
37 study was to determine a maximum tolerated dose that induced significant toxicity but resulted in ≥90%
38 survival and ≤10% decrease in weight gain. Statistical analyses included ANOVA, and chi-squared test.
39 Lethality was significantly increased in the high-dose group. Body weight gain was depressed in groups
40 exposed to ≥1.25% bisphenol A. Clinical signs of toxicity were observed in the 2.5 and 5.0% dose groups
41 and included dehydration, dyspnea, lethargy, tremors, ptosis, piloerection, and diarrhea.

42
43 In the reproduction and fertility study (Task 2), 11-week-old mice were randomly assigned to treatment
44 groups according to body weight. The mice were fed diets containing 0, 0.25, 0.5, or 1.0% bisphenol A.
45 The NTP stated that a 40 g mouse consuming 7 g of feed/day would be exposed to bisphenol A at 437.5,
46 875, and 1750 mg/kg bw/day. **[Based on body weight and feed intake values reported for males at ~3**
47 **week intervals, CERHR estimated mean bisphenol A intake at ~365, 740, and 1630 mg/kg bw/day.**
48 **Feed intakes were reported only at week 1 and 18 for females, and week 18 most likely represented**
49 **the lactation period. For week 1, bisphenol A intake by females was estimated at 410, 890, and 1750**
50 **mg/kg bw/day. At week 18, bisphenol A intake by females was estimated at 1090, 1785, and 3660**
51 **mg/kg bw/day.]** There were 40 mice/sex in the vehicle control group and 20/sex in each bisphenol A

4.0 Reproductive Toxicity Data

1 group. Exposure to bisphenol A began during a 7-day pre-mating period. Following the pre-mating period,
2 males and females from the same treatment group were randomly paired and housed together for 98 days
3 and following the mating period, each male and female was housed separately for 21 days. Bisphenol A
4 dosing was continued throughout the mating and separation period. Concentration and stability of
5 bisphenol A in feed were verified. During the 98-day cohabitation period, pups born were counted, sexed,
6 and weighed. All litters excluding the last one born were killed on the day of birth so that animals could
7 continue mating. The last litter was raised by the dam and weaned on PND 21 (day of birth not defined).
8 Birth weight and weight gain were recorded in the last litter. Reproductive endpoints in parental rats
9 included the number of litters born and fertility. Statistical analyses included Kruskal-Wallis ANOVA on
10 ranks, Mann-Whitney *U* test, chi-squared test, 1-way ANOVA, arcsine square-root transformation, and
11 Duncan multiple range test.

12
13 In the cross-over trial (Task 3), ~20 males and females from the high-dose group were randomly paired
14 with control mice for 7 days in order to determine the affected sex. Twenty control males and females
15 were also paired. The animals were not exposed to bisphenol A during the 1-week mating period, but in
16 animals from the high dose group, dosing with bisphenol A was continued for 21 days upon separation of
17 the mating pairs. Vaginal smears were obtained from females that did not mate or did not appear to be
18 pregnant. Fertility and offspring survival were determined. Parental mice from the control (n = 38/sex)
19 and high-dose groups (n = 19/sex) were necropsied within a week following completion of the cross-over
20 trial. Body, liver, kidney, and reproductive organ weights were obtained, and sperm count, morphology,
21 and motility were determined. Testes, ovaries, and oviducts were fixed in Bouin solution and prostate,
22 seminal vesicles/coagulating glands, uterus, liver, and kidney were fixed in 10% neutral buffered formalin
23 for histopathological evaluation.

24
25 In Task 4 of the study, 20 F₁ mice/sex/group (at least 2/sex from 10 randomly selected litters/group) were
26 mated within dose groups for 7 days and examined for reproductive function. Because fewer F₁ mice in
27 high-dose group were available as a result of increased mortality, only 11 mice/sex were mated. The
28 animals continued to receive the same diet given to their parents. Vaginal smears were obtained from
29 females that did not mate or did not appear to become pregnant. One litter/pair was examined for sex,
30 body weight, and viability. The parental F₁ animals from all dose group were killed and examined as
31 described for Task 3 of the study.

32
33 Treatment-related effects observed in adult rats are summarized in Table 110, and effects occurring in
34 immature rats are summarized in Table 111. Bisphenol A treatment had no effect on mating or fertility
35 index in F₀ or F₁ mice. Postpartum body weights were reduced in F₀ dams of the high-dose group. In F₀
36 mice, the number of litters produced/pair and numbers of live F₁ pups/litter were reduced at the mid- and
37 high-dose level. A decrease in the proportion of pups born alive occurred in F₀ mice of the high-dose
38 group. No effects were observed on sex ratios of F₁ or F₂ pups. Weights of live F₁ pups were increased at
39 the mid and high dose. There were no significant effects when pup weights were adjusted for total
40 numbers of live and dead pups in the litter. Therefore the NTP concluded that the increased pup weights
41 resulted from the smaller litter size. Body weights were evaluated through PND 21 in F₁ pups, and no
42 effects were found on pup body weight gain during the lactation period. Mortality in F₁ offspring during
43 the postnatal period was increased in the high-dose group.

44
45 The cross-over test revealed no effect on mating or fertility in either males or females exposed to
46 bisphenol A. Postpartum body weight was not affected in the treated females. The number of live
47 pups/litter was significantly reduced [by 26%] in the group containing treated males and [by 51%] in the
48 group containing treated females. Live pup weight was increased in the group containing treated females,
49 but there was no significant effect following adjustment for litter size. There were no effects on the
50 proportion of pups born alive or on sex ratio.

4.0 Reproductive Toxicity Data

1 In sperm analyses conducted in high-dose F₀ males and all dose groups of F₁ males, sperm motility was
 2 reduced in high-dose F₀ males and mid-dose F₁ males. There were no effects on sperm count or
 3 morphology in either generation. Effects were observed on organ weights, which were examined in F₀
 4 adults of the high-dose group and F₁ animals from each treatment group. Effects on absolute reproductive
 5 organ weights of F₁ mice included decreased right epididymis weight at all doses, decreased left
 6 testis/epididymis weight at the mid and high dose, and decreased seminal vesicle weight at the high dose.
 7 Significant effects on relative organ weights adjusted for body weight in F₁ rats included decreased right
 8 epididymis weight at all doses, decreased seminal vesicle weight at the low and high dose, and decreased
 9 relative left testis and epididymis weight at the mid and high dose. Reproductive organ weight effects
 10 observed in high-dose F₀ males included decreased absolute and relative seminal vesicle weight. There
 11 were no effects on prostate weight. No effects were reported for estrous cyclicity of F₀ females. There
 12 were no gross or histopathological alterations in F₀ or F₁ reproductive organs including testis, epididymis,
 13 prostate, seminal vesicles, ovary, vagina, and uterus. Effects observed in high-dose F₀ animals were also
 14 summarized in a report by Morrissey et al. (460).

15
 16 Effects were observed on non-reproductive organ weights, which were examined in F₀ adults of the high-
 17 dose group and F₁ animals from each treatment group. In the F₁ mice, dose-related effects on absolute
 18 organ weights included increased kidney/adrenal weight at all doses in both sexes and increased liver
 19 weight in mid- and high-dose females and high-dose males. Significant effects on relative organ weight
 20 adjusted for body weight in F₁ rats included increased liver and kidney/adrenal weights at all doses in
 21 both sexes. Organ weight effects observed in high-dose F₀ males included increased absolute and relative
 22 liver and kidney/adrenal weight. In F₀ female rats of the high-dose group, absolute and relative liver
 23 weight and relative kidney weights were increased. Body weights of high-dose F₀ females were reduced
 24 at necropsy. Histopathology was examined in F₀ rats of the high-dose group and F₁ rats from all dose
 25 groups. Treatment-related hepatic lesions observed in both generations included multifocal necrosis,
 26 multinucleated giant hepatocytes in males and females, and centrilobular hepatocytomegaly in males.
 27 Multifocal mineralization of liver cells was also observed in F₁ females of the high-dose group. Hepatic
 28 lesions were observed at all dose levels for F₁ males and in F₁ females of the mid- and high-dose group.
 29 Treatment-related renal lesions were observed in both generations and described as tubular cell nuclear
 30 variability, increased severity of spontaneous tubular interstitial lesions, cortical tubular dilatation,
 31 mineralization of renal cells, and micro-calculi in tubular epithelium that sometimes occurred with
 32 effaced tubular epithelium, tubular regeneration, and/or dilated tubules containing casts. **[It appears that**
 33 **the incidence of renal lesions was increased at all doses in F₁ rats.]** Renal lesions were stated to
 34 generally be more prominent in females than males. The study authors concluded that exposure of mice to
 35 bisphenol A resulted in toxicity to the reproductive system, kidney, and liver. The possibility was noted
 36 that some or all effects on reproductive performance may have been secondary to the generalized toxicity
 37 of bisphenol A.

38
 39 **Table 110. Effects Observed in Adult Mice Dosed with Bisphenol A in a Continuous Breeding**
 40 **Study.**

Endpoint	Dose, % in diet [mg/kg bw/day]						
	0.25 [437.5]	0.5 [875]	1.0 [1750]	BMD ₁₀	BMDL ₁₀	BMD _{1SD}	BMDL _{1SD}
<i>F₀ males and females</i>							
Litters/pair	↔	↓5%	↓9%	1.0 [1750]	0.74 [1295]	0.96 [1680]	0.66 [1155]
Postpartum dam weight ^a	↔	↔	↓6–9%	1.0 [1750]	0.83 [1452]	0.87 [1522]	0.66 [1155]
Necropsy dam weight	No data	No data	↓4%				
Percent motile sperm	No data	No data	↓39%				
Relative organ weight, males ^b							
Liver	No data	No data	↑29%				
Kidney/adrenal	No data	No data	↑16%				
Seminal vesicle	No data	No data	↓19%				

4.0 Reproductive Toxicity Data

Endpoint	Dose, % in diet [mg/kg bw/day]				BMD ₁₀	BMDL ₁₀	BMD _{1SD}	BMDL _{1SD}
	0.25 [437.5]	0.5 [875]	1.0 [1750]					
Relative organ weight, females ^b								
Liver	No data	No data	↑27%					
Kidney/adrenal	No data	No data	↑10%					
Liver lesions, males and females ^c	No data	No data	↑ ^d					
Kidney lesions, males and females ^c	No data	No data	↑ ^d					
<i>F₁ males and females</i>								
Relative organ weight, males ^b								
Liver	↑7%	↑7%	↑29%	0.62 [1085]	0.42 [735]	0.59 [1032]	0.39 [682]	
Kidney/adrenal ^e	↑16%	↑20%	↑20%	0.18 [315]	0.14 [245]	0.15 [262]	0.12 [210]	
Left testis/epididymis ^e	↔	↓10%	↓9%	0.64 [1120]	0.32 [560]	0.53 [928]	0.27 [472]	
Right testis ^f	↔	↓13%	↔					
Right epididymis ^e	↓11%	↓16%	↓18%	0.24 [420]	0.15 [262]	0.46 [805]	0.25 [438]	
Seminal vesicle	↓11%	↔	↓28%	0.40 [700]	0.29 [508]	0.66 [1155]	0.47 [822]	
Relative organ weight, females ^b								
Liver	↑6%	↑13%	↑20%	0.49 [858]	0.38 [665]	0.45 [788]	0.35 [612]	
Kidney/adrenal ^f	↑13%	↑15%	↑13%					
Percent motile sperm ^f	↔	↓31%	↔					
Liver lesions, males ^c	↑ ^d	↑ ^d	↑ ^d					
Liver lesions, females ^c	↔	↑ ^d	↑ ^d					
Kidney lesions, males and females	↑ ^d	↑ ^d	↑ ^d					

↑,↓ Statistically significant increase, decrease compared to controls; ↔ no statistically significant effects compared to controls.

^aValues were reported following the birth of 5 litters, the benchmark doses are for values reported following the birth of the fifth litter because the greatest magnitude of effect was observed at that time point.

^bRelative organ weights were adjusted for body weight; when absolute and relative organ weights changed in the same direction, only the relative organ weights were listed in this table.

^cSee text for a description of the types of lesions observed

From NTP (458)

^dIt does not appear that statistical analyses were conducted for histopathology data, but incidence was increased compared to controls.

^eBenchmark doses were estimated using a polynomial model.

^fBenchmark doses were not estimated for endpoints without dose-response relationships.

1

2 **Table 111. Effects in Immature F₁ Mice in a Continuous Breeding Study with Bisphenol A**

Endpoint	Dose, % in diet [mg/kg bw/day]				BMD ₁₀	BMDL ₁₀	BMD _{1SD}	BMDL _{1SD}
	0.25 [437.5]	0.5 [875]	1.0 [1750]					
Live pups/litter	↔	↓20%	↓48%	0.30 [525]	0.20 [350]	0.43 [752]	0.30 [525]	
Proportion pups born alive	↔	↔	↓4%	3.0 [5250]	0.79 [1382]			
Live birth weight ^a	↔	↑5%	↑6%	0.43 [752]		0.34 [595]		
Mortality by PND 21 ^b	↔	↔	↑ to 37.5%	0.48 [840]	0.40 [700]			

↑,↓ Statistically significant increase, decrease compared to controls; ↔ no statistically significant effects compared to controls.

^aHill model used for benchmark dose calculations.

^bControl mortality was 6.3%. Mortality was reported on a per pup basis, which limits the utility of the benchmark dose model.

From NTP (458).

3

4.0 Reproductive Toxicity Data

1 **Strengths/Weaknesses:** This comprehensive toxicology study was well-conducted. General toxicity was
2 clearly demonstrated at all F₁ dose levels, and histopathological findings appear to be a sensitive indicator
3 of effect. In Task 2, a clear effect on fertility was found with a NOAEL of 0.25% bisphenol A in the diet.
4 This study demonstrates changes in F₁ male absolute reproductive weights (seminal vesicle with
5 coagulating gland as well as epididymis; the testis and prostate appear not to have been appreciably
6 affected). This study also suggested that reproductive toxicity and general toxicity occurred at similar
7 dose levels. Bisphenol A-mediated general toxicity may have contributed to the observed female fertility
8 effect, because this effect was noted with dosed females cohabiting with non-dosed males. In the male,
9 however, the effect on motility is likely bisphenol A-related, resulting in the observed fertility deficits. As
10 a limitation of this design, because bisphenol A was in the diet, exposure to bisphenol A did not occur
11 during cohabitation; therefore, direct exposure to bisphenol A was minimal or nonexistent during sperm
12 maturation, capacitation and ovulation.

13
14 **Utility (Adequacy) for CERHR Evaluation Process:** This comprehensive data set examined the
15 reproductive toxicity of bisphenol A in mice with a large number of animals and multiple endpoints.
16 These data are highly useful.

17 **Tyl et al. (376)**, sponsored by the American Plastics Council, conducted a 2-generation study of
18 bisphenol A in mice. The study was conducted according to GLP. CD-1 mice were received in two cohorts
19 approximately 2 weeks apart and data from the 2 cohorts were combined. Mice were fed Purina Certified
20 Ground Rodent Diet No. 5002. The supplier provided information about phytoestrogen content of feed
21 (177–213 ppm genistein, 173–181 ppm daidzein, and 39–55 ppm glycitein). Mice were housed in
22 polypropylene cages with Sani-Chip® bedding. Assignment of F₀ animals to groups involved
23 randomization stratified by weight. F₀ and F₁ mice (28 sex/group/generation) were fed diets containing
24 bisphenol A (99.70–99.76% purity) at 0.018, 0.18, 1.8, 30, 300, or 3500 ppm. Target intakes were 0.003,
25 0.03, 0.3, 5, 50, or 600 mg/kg bw/day, respectively. Based on measured feed intake, the study authors
26 estimated bisphenol A intake in males at 0.0024–0.0038, 0.024–0.037, 0.24–0.37, 3.98–6.13, 39.1–60.8,
27 or 529–782 mg/kg bw/day. Bisphenol A intakes (in mg/kg bw/day) by females were estimated at 0.0030–
28 0.0041, 0.030–0.042, 0.32–0.43, 5.12–7.12, 54.2–67.8, 653–910 during the pre-mating period; 0.0027–
29 0.0029, 0.027–0.028, 0.28–0.29, 4.65–4.80, 47.0–48.6, 552–598 during the gestation period; and 0.0087–
30 0.0063, 0.062–0.091, 0.61–0.89, 10.4–15.1, 103.2–146.4, 1264–1667 during the lactation period. In each
31 generation, there were 2 vehicle controls groups with 28 mice/sex/group. A positive control group was
32 given feed containing 17β-estradiol at 0.5 ppm (target intake of 0.08 mg/kg bw/day). Estimated intakes
33 for 17β-estradiol (in mg/kg bw/day) were 0.074–0.104 in males, 0.093–0.12 in females during the pre-
34 mating period, 0.08–0.081 in females during the gestation period, and 0.160–0.25 in females during the
35 lactation period. Dose selections were based on observations from several studies. Homogeneity, stability,
36 and concentration of bisphenol A in feed were verified. Exposure of F₀ mice began at ~6 weeks of age.
37 Exposure of F₁ animals began at weaning, although it was noted that pups began eating the dosed feed in
38 the late lactation period. F₀ and F₁ mice were fed the bisphenol A-containing diets for a minimum of 8
39 weeks prior to mating and during a 2-week mating period. Exposures of males continued through the
40 gestation period of the litters they sired. Exposures of females continued through the gestation and
41 lactation period. During the study, adult animals were monitored for clinical signs of toxicity, body
42 weight, and food intake.

43
44 Estrous cycles were evaluated in F₀ and F₁ females during the last 3 weeks of the pre-breeding exposure
45 period. Day of vaginal plug was defined as GD 0 and day of birth was considered PND 0. F₁ and F₂ pups
46 were counted, sexed, weighed, and assessed for viability and physical abnormalities at birth and
47 throughout the lactation period. Anogenital distance was measured in F₁ and F₂ pups at birth and on PND
48 21. On PND 4, F₁ and F₂ litters were standardized to 10 pups, with equal numbers per sex when possible.
49 Pups removed on PND 4 were killed and examined for visceral alterations, with a focus on the
50 reproductive system. The remaining pups were maintained and weaned on PND 21. At weaning, 28 F₁

4.0 Reproductive Toxicity Data

1 pups/sex/group (1 per sex per litter) were randomly selected for mating and those animals were referred to
2 as parental mice. An additional F₁ male/litter was selected for a 3 month exposure (referred to as retained
3 males). Two F₁ pups/sex/litter were selected for gross necropsy and organ weight measurement at
4 weaning. Histopathological examination of reproductive organs was conducted in one PND 21
5 pup/sex/litter. Histopathological evaluation of reproductive and systemic organs were conducted in the
6 second F₁ pup from each group at weaning. All F₂ pups were killed at weaning and organ weights were
7 measured. Vaginal opening and preputial separation were monitored in parental and retained F₁ mice.
8 Parental F₀ and F₁ males were killed following delivery of the litters they sired. Retained F₁ males were
9 killed at the same time as the parental F₁ males. Parental F₀ and F₁ females were killed after their pups
10 were weaned. Organs, including those of the reproductive system, were weighed in adult F₀ and F₁
11 animals. Histopathological evaluations were conducted in all animals from the vehicle control group, in
12 10 F₀ and F₁ parental animals from each treatment group, in all F₁ retained males, and 10 animals from
13 the 17 β -estradiol positive control group. Histopathological evaluation of reproductive organs was also
14 conducted in animals with suspected reduced fertility. Testes were preserved in Bouin fixative. Daily
15 sperm production, efficiency of daily sperm production, and epididymal sperm count, motility, and
16 morphology, were evaluated in F₀ and F₁ males. Data from the 2 control groups were analyzed separately
17 and then pooled for statistical analysis of treatment groups. Statistical analyses included ANOVA, Levene
18 test, robust regression methods, Wald chi-squared test, *t*-test, Dunnett test, Fisher exact probability test,
19 and ANCOVA.

20
21 Treatment- or dose-related results and observations in reproductive organs of adult animals are
22 summarized in Table 112. There were no consistent effects on body weight or body weight gain in F₀
23 males. Body weight gain during lactation was increased in F₀ females from the 3500 ppm group. During
24 the pre-mating period, body weights were decreased by $\leq 10\%$ in F₁ parental animals from the 3500 ppm
25 group (study days 0, 7, 49, and 56 in males and study 0 in females). In retained F₁ males from the 3500
26 ppm group, body weights were decreased at most time periods between study days 7 and 84 and at
27 necropsy. No consistent or dose-related changes in feed intake or efficiency were observed throughout the
28 study in F₀ or F₁ animals. There were no clinical signs of toxicity or treatment-related deaths in F₀ or F₁
29 males or females. Increases in absolute and relative to body or brain weights of kidney and liver were
30 consistently observed in F₀ and F₁ adults. Significant and dose-related organ weight changes relative to
31 body weight are summarized in Table 112. Other effects on organ weight (e.g., seminal vesicles,
32 epididymides, coagulating glands, and pituitary) were not considered to be treatment-related by study
33 authors due to factors such as lack of a dose-response relationship, no consistency between absolute and
34 relative weights, no histopathology, or no consistency across generations. Absolute and relative prostate
35 weights were unaffected by bisphenol A exposure. There were no treatment-related gross systemic
36 findings in F₀ or F₁ adults. Incidence of minimal to mild hepatocyte centrilobular hypertrophy was
37 increased in both generations at 300 and/or 3500 ppm (see Table 112). Renal nephropathy incidence was
38 increased in F₀ males and in F₁ males and females of the 3500 ppm group. **[It did not appear that
39 histopathological data were statistically analyzed.]**

40
41 Treatment- or dose-related reproductive effects in adult animals are summarized in Table 112. Bisphenol
42 A exposure had no effect on numbers of implantation sites or resorptions or on mating, fertility, or
43 gestational indices in F₀ or F₁ mice. Gestational length was increased in F₀ and F₁ females from the 3500
44 ppm group; the study authors stated the effect was of unknown biological significance. Epididymal sperm
45 concentration was decreased in F₀ males of the 3500 ppm group but no effect was observed in F₁ parental
46 or retained males. There was no effect on daily sperm production, efficiency of daily sperm production, or
47 sperm motility or morphology in either generation. The study authors did not consider the decrease in
48 sperm concentration in F₀ animals to be treatment-related based on lack of consistency between
49 generations, no effect on any other andrological endpoint, and no effect on fertility. Estrous cyclicity and
50 numbers of ovarian primordial follicle counts were not affected by bisphenol A exposure in F₀ or F₁
51 females. The only gross observation in reproductive organs was a slightly increased incidence of gross

4.0 Reproductive Toxicity Data

1 ovarian cysts in F₀ females from the 3500 ppm group. The incidence of paraovarian cysts was increased
2 in F₀ and F₁ females from the 3500 ppm group. **[It did not appear that histopathological data were**
3 **statistically analyzed.]**
4

5 Significant findings in developing mice are summarized in Table 113. Live F₁ and F₂ pups and litters at
6 birth, sex ratio, and survival during the lactation period were not affected and there were no clinical or
7 gross signs of toxicity in F₁ or F₂ offspring. A non-dose-related decrease in PND 21 survival index and
8 lactational index (pups surviving on PND 21/PND 4) was described in F₂ pups of the 300 ppm group.
9 **[The biological significance of the effect was not discussed by the study authors, but because the**
10 **effect was not dose-related it is unlikely to be of biological significance.]** In F₁ pups from the 3500
11 ppm group, body weights were reduced during PND 7, 14, and 21 in F₁ females and both sexes combined
12 and on PND 7 and 21 in F₁ males. Body weight results for both sexes combined are summarized in Table
13 113. An increase in male pup body weight observed on PND 7 in the 1.8 ppm group was not considered
14 to be treatment related by the study authors because no dose-response relationship was observed. There
15 was no effect on anogenital distance in F₁ or F₂ males or females on PND 0. Anogenital distance was also
16 unaffected in F₂ males and F₁ and F₂ females on PND 21. Anogenital distance adjusted for body weight
17 was reduced in F₁ males from the 300 and 3500 ppm groups on PND 21. Based on the lack of effect on
18 anogenital distance at birth and inconsistencies between generations, the study authors did not consider
19 the decreases in anogenital distance in F₁ males to be treatment-related. An increase in anogenital distance
20 in F₂ females from the 0.018 ppm group on PND 0 was not considered to be treatment related by the
21 study authors. Preputial separation (absolute age and adjusted for body weight on day of acquisition) was
22 delayed in parental and retained F₁ males of the 3500 ppm group. When adjusted for PND 30 body
23 weight, preputial separation was delayed in retained but not parental F₁ males from the 3500 ppm group.
24 Data for preputial separation adjusted for body weight on day of acquisition are shown in Table 113.
25 Body weights on day of vaginal opening were lower in F₁ females from the 3500 ppm group. Day of
26 vaginal opening was accelerated in the 3500 ppm group if adjusted for PND 21 body weight, but not body
27 weight on the day of acquisition. Due to the lack of effect when adjusted for body weight on day of
28 acquisition, the study authors did not consider effects on vaginal opening to be treatment related.
29

30 Shown in Table 113 are significant organ weight effects relative to body weight. Dose-related organ
31 weight changes in F₁ weanlings that were considered to be treatment-related by study authors included
32 decreased absolute and relative (to body or brain weight) spleen and paired testes weights at 3500 ppm.
33 Treatment-related absolute organ weight changes in F₂ weanlings included decreased weights of spleen,
34 paired testes, and seminal vesicles with coagulating glands in the 3500 ppm group. Changes in organ
35 weights relative to body weight in F₂ weanlings included decreased spleen weight in males and females
36 and increased relative left kidney weight in 3500 ppm males. Treatment-related changes in organ weight
37 relative to brain weight in F₂ weanlings were decreased spleen weight in both sexes and decreased paired
38 testes weight at 3500 ppm and seminal vesicles with coagulating glands at 300 and 3500 ppm. Other
39 organ weight effects (e.g, affecting epididymides, thymus, brain, ovaries, and/or uterus with cervix and
40 vagina weights) were not considered to be dose-related due to lack of dose-response relationships or no
41 consistent effects across generations. Included in Table 113 are significant organ weight effects relative to
42 body weight. Significant organ weight effects relative to brain weight were included in Table 113 when
43 the organ weight effect was significant only when normalized for brain weight. The study authors
44 reported no gross findings in F₁ or F₂ weanlings. **[Although not clear because the number of animals**
45 **examined for gross testicular effects was not reported in Tables 23 and 49 of the study, it appeared**
46 **that the incidence of undescended bilateral testes may have been increased in F₁ and F₂ weanling**
47 **males of the 3500 ppm group.]** The incidence of hepatic cytoplasm alteration (clear hepatocellular
48 cytoplasm, slightly more basophilic cytoplasm, and/or minute vacuoles) was apparently increased in F₁
49 males from the 300 and 3500 ppm groups and F₁ females and F₂ males from the 3500 ppm group. The
50 incidence of seminiferous tubule hypoplasia was increased in F₁ and F₂ weanlings from the 3500 ppm
51 group. **[Another histopathological finding that appeared to be possibly increased in weanlings from**

4.0 Reproductive Toxicity Data

1 **the 3500 ppm group was unilateral hydronephrosis in F₁ males. It did not appear that**
2 **histopathological data were statistically analyzed.]**
3

4 Effects of 17 β -estradiol in males were delayed preputial separation, reduced anogenital distance at
5 weaning but not at birth, decreased weights of testes, epididymides, and seminal vesicles with
6 coagulating gland, and increased incidence of seminiferous tubule hypoplasia and undescended testis.
7 Effects of 17 β -estradiol in female mice were accelerated vaginal patency, increased uterus with cervix
8 and vagina weight, fluid filled/enlarged uterus, enlarged/thickened vagina, increased vaginal epithelial
9 keratinization, and prolonged gestation. Reproductive effects in the 17 β -estradiol group included
10 decreased fertility, increased stillbirth, reduced live pups per litter, and increased dead pups.
11

12 The study authors identified bisphenol A NOELs of 30 ppm (~5 mg/kg bw/day) for systemic effects, 300
13 ppm (~50 mg/kg bw/day) for developmental toxicity, and 3500 ppm (~600 mg/kg bw/day) for
14 reproductive toxicity.
15

16 **Strengths/Weaknesses:** Strengths include the large number and range of doses examined, the rigor with
17 which the study was performed, the large sample size in each group, the number of additional animals
18 per litter that were retained and examined, the use of a concurrent estrogenic positive control group, and
19 the thoroughness of the histologic evaluation. Weaknesses might include that brain biochemistry and
20 other central nervous system (CNS) metrics were not examined, and that statistics was not performed on
21 some histopathology findings.
22

23 **Utility (Adequacy) for CERHR Evaluation Process:** This exceptional study is very useful for the
24 evaluation process, and will carry significant weight in the evaluation of structural, histogenic, and
25 fertility endpoints.
26

4.0 Reproductive Toxicity Data

1 Table 112. Treatment-Related Effects in Adult Mice Fed Bisphenol A through Diet in a Multigeneration Reproductive Toxicity Study

Endpoint	Dose, ppm diet [mg/kg bw/day based on target intakes provided by study authors]							BMD ₁₀	BMDL ₁₀	BMD _{1SD}	BMDL _{1SD}
	0.018 [0.003]	0.18 [0.03]	1.8 [0.3]	30 [5]	300 [50]	3500 [600]					
Body weight gain during lactation, F ₀	↔	↔	↔	↔	↔	↔	↑2.2-fold	249 [42.2]	150 [25.4]	4258 [722]	2941 [498]
Terminal body weight F ₁ retained males	↔	↔	↔	↔	↔	↔	↓10%	3455 [586]	2388 [405]	3503 [594]	2608 [442]
Relative liver to body weight											
F ₀ males	↔	↔	↔	↔	↔	↔	↑17%	2189 [371]	1820 [308]	2021 [343]	1668 [283]
F ₁ parental males	↔	↔	↔	↔	↔	↑5%	↑22%	1662 [282]	1425 [242]	1637 [277]	1389 [235]
F ₁ retained males	↔	↔	↔	↔	↔	↔	↑23%	1584 [268]	1383 [234]	1685 [286]	1405 [238]
F ₀ females	↔	↔	↔	↔	↔	↔	↑17%	2524 [428]	1595 [270]	3014 [511]	2155 [365]
F ₁ females	↔	↔	↔	↔	↔	↔	↑10%	3424 [580]	2438 [413]	3551 [602]	3024 [513]
Relative right kidney to body weight											
F ₀ males	↔	↔	↔	↔	↔	↑8%	↑20%	1861 [315]	1536 [260]	2100 [356]	1723 [292]
F ₁ parental males	↔	↔	↔	↔	↑11%	↑10%	↑21%	2079 [352]	1913 [324]	862 [146]	773 [131]
F ₁ retained males	↔	↔	↔	↔	↔	↔	↑27%	1501 [254]	1229 [208]	1978 [335]	1610 [273]
F ₀ females	↔	↔	↔	↔	↔	↔	↑13%	3568 [605]	2504 [424]	4326 [733]	3041 [515]
F ₁ females	↔	↔	↔	↔	↔	↔	↑8%	3629 [615]	2976 [504]	3702 [627]	3393 [575]
Relative left kidney to body weight											
F ₀ males	↔	↔	↔	↔	↔	↑9%	↑19%	1899 [322]	1548 [262]	2249 [381]	1825 [309]
F ₁ parental males	↔	↔	↔	↔	↑13%	↑10%	↑22%	2074 [352]	1650 [280]	2547 [432]	2020 [342]
F ₁ retained males	↔	↔	↔	↔	↔	↑11%	↑28%	1466 [248]	1205 [204]	1937 [328]	1582 [268]
F ₀ females	↔	↔	↔	↔	↔	↔	↑11%	3746 [635]	2550 [432]	4773 [809]	3258 [552]
Relative pituitary to body weight											
F ₁ parental males ^a	↔	↔	↔	↔	↔	↔	↑10%	3413 [578]	2087 [554]	3627 [615]	3182 [539]
F ₁ retained males ^a	↔	↔	↔	↔	↔	↔	↑16%	2678 [454]	1934 [328]	3476 [589]	2512 [426]
Relative brain to body weight											
F ₁ retained males ^a	↔	↔	↔	↔	↔	↔	↑9%	2678 [454]	1934 [328]	3476 [589]	2512 [426]
Hepatocyte centrilobular hypertrophy incidence (control incidence in parentheses)											
F ₀ males (6/56)	1/10	2/10	2/10	0/10	4/10	10/10		122 [20.7]	70 [11.8]		
F ₁ parental males (7/55)	0/10	0/10	4/10	2/10	1/10	6/10		879 [149]	578 [98.0]		
F ₁ retained males (4/50)	1/10	3/10	2/10	2/10	5/10	7/10		656 [111]	442 [74.9]		
F ₀ females (1/56)	0/10	0/10	0/10	0/10	1/10	6/10		1348 [228]	947 [161]		
F ₁ females (2/55)	0/10	0/10	0/10	0/10	3/11	7/10		962 [163]	679 [115]		

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Endpoint	Dose, ppm diet [mg/kg bw/day based on target intakes provided by study authors]						BMD ₁₀	BMDL ₁₀	BMD _{1SD}	BMDL _{1SD}
	0.018 [0.003]	0.18 [0.03]	1.8 [0.3]	30 [5]	300 [50]	3500 [600]				
Renal nephropathy incidence (control incidence in parentheses)										
F ₀ males (12/56)	0/10	3/10	2/10	2/10	1/10	4/10	1556 [264]	750 [127]		
F ₁ parental males (6/55)	2/10	0/10	1/10	2/10	0/10	4/10	1418 [240]	838 [142]		
F ₁ retained males (8/50)	1/10	0/10	0/10	2/10	0/10	3/10	1991 [337]	992 [168]		
F ₁ females (10/55)	1/10	3/10	0/10	1/10	1/11	4/11	1646 [279]	847 [144]		
Paraovarian cysts (control incidence in parentheses)										
F ₀ (9/56)	1/11	2/12	1/11	1/12	3/14	7/17	1328 [225]	833 [141]		
F ₁ (14/55)	1/11	1/11	1/10	2/10	2/11	7/15	1193 [202]	708 [120]		
Epididymal sperm concentration, F ₀ ^a	↔	↔	↔	↔	↔	↓15%	3343 [567]	1884 [319]	3581 [607]	3241 [549]
Gestational length										
F ₀	↔	↔	↔	↔	↔	↑2%	21,351 [3619]	3770 [639]	6749 [1144]	3536 [599]
F ₁	↔	↔	↔	↔	↔	↑2%	17,820 [3020]	3784 [641]	4552 [772]	3134 [531]

↑,↓ Statistically significant increase, decrease; ↔ no statistically significant effect compared to controls

^aNot considered a treatment-related effect by study authors.

4.0 Reproductive Toxicity Data

1

2 **Table 113. Treatment- or Dose-Related Effects in Developing Mice in a Multigeneration Reproductive Toxicity Study with Bisphenol A.**

Endpoint ^a	Dose, ppm diet [mg/kg bw/day based on target intakes provided by study authors]									
	0.018 [0.003]	0.18 [0.03]	1.8 [0.3]	30 [5]	300 [50]	3500 [600]	BMD ₁₀	BMDL ₁₀	BMD _{1SD}	BMDL _{1SD}
Body weight										
F ₁ , PND 7	↔	↔	↔	↔	↔	↓13%	3304 [560]	1849 [313]	3433 [582]	2403 [407]
F ₁ , PND 14	↔	↔	↔	↔	↔	↓11%	3453 [585]	2256 [382]	3639 [617]	2988 [506]
F ₁ , PND 21	↔	↔	↔	↔	↔	↓17%	3236 [548]	1577 [267]	3421 [580]	2342 [370]
F ₁ male, PND 21 necropsy	↔	↔	↔	↔	↔	↓12%	3325 [564]	1845 [313]	3776 [640]	3536 [599]
F ₁ female, PND 21 necropsy	↔	↔	↔	↔	↔	↓18%	2284 [387]	1501 [254]	4577 [776]	3529 [598]
Lactational survival indices (control index, %, in parentheses)										
F ₂ PND 21 survival (100%) ^c	↔	↔	↔	↔	↓ to 86.6%	↔				
F ₂ Lactational index (97.2%) ^c	↔	↔	↔	↔	↓ to 86.6%	↔				
Relative thymus to body weight, F ₁ male, PND 21 ^b	↔	↔	↔	↔	↑13% ^b	↑10% ^b				
Relative spleen to body weight										
F ₁ male, PND 21	↔	↓12%	↔	↔	↔	↓30%	3123 [529]	1074 [182]	3538 [600]	3148 [534]
F ₂ male, PND 21	↔	↔	↔	↔	↔	↓20%	2148 [364]	1425 [242]	7013 [1189]	3560 [603]
F ₁ female, PND 21	↔	↔	↔	↔	↔	↓23%	3168 [537]	647 [110]	4571 [775]	3677 [623]
F ₂ female, PND 21	↔	↔	↔	↔	↔	↓21%	1787 [303]	1311 [222]	5022 [851]	3517 [596]
Relative paired testes weight to body or brain weight										
F ₁ , PND 21 (body weight)	↔	↔	↔	↔	↔	↓8%	3578 [606]	2720 [461]	3861 [654]	3550 [602]
F ₂ , PND 21 (brain weight)	↔	↔	↔	↔	↔	↓11%	3316 [562]	2003 [339]	5342 [905]	3571 [605]
Relative paired epididymides to body weight, F ₁ ^b	↔	↑18%	↔	↔	↔	↔				
Relative brain to body weight F ₁ female, PND 21 ^b	↔	↔	↔	↔	↔	↑17% ^b	2219 [376]	1415 [240]	3576 [606]	2825 [479]
Relative left kidney to body weight, F ₂ male, PND 21	↔	↔	↔	↔	↔	↑6%	6664 [1129]	3540 [600]	8501 [1441]	3589 [608]
Relative seminal vesicles with coagulating gland to brain weight, F ₂ ^b	↔	↔	↔	↔	↓15%	↓16%	2389 [405]	1315 [223]	11,294 [1914]	3631 [615]

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Endpoint ^a	Dose, ppm diet [mg/kg bw/day based on target intakes provided by study authors]						BMD ₁₀	BMDL ₁₀	BMD _{1SD}	BMDL _{1SD}	
	0.018 [0.003]	0.18 [0.03]	1.8 [0.3]	30 [5]	300 [50]	3500 [600]					
Uterus with cervix and vagina weight relative to bodyweight, F ₂ PND 21 ^b	↔	↔	↔	↔	↓16%	↔					
Relative paired ovary weights, F ₁ ^b	↔	↔	↑	↔	↔	↔					
Hepatic cytoplasm alteration (control incidence in parentheses)											
F ₁ males (6/44)	1/26	0/17	1/22	6/24	10/20	13/20	732 [124]	546 [92.5]			
F ₂ males (6/54)	1/25	1/25	1/25	1/24	2/20	9/23	1442 [244]	1050 [178]			
F ₁ females (2/46)	1/27	2/21	3/24	4/26	8/16	6/22	1966 [333]	1182 [200]			
Unilateral hydronephrosis, F ₁ males (0/44) ^b	0/26	0/17	0/21	0/24	0/21	3/21 ^b					
Seminiferous tubule hypoplasia (control incidence in parentheses)											
F ₁ (1/96)	0/54	0/37	1/45	3/51	2/45	5/43	3485 [591]	2398 [406]			
F ₂ (5/114)	1/53	2/61	2/55	0/51	5/49	20/57	1670 [283]	1377 [233]			
Anogenital distance adjusted for body weight F ₁ male, PND 21 ^b	↔	↔	↔	↔	↓4%	↓5%	8099 [1373]	3582 [607]	10,436 [1769]	3632 [616]	
Age of preputial separation (adjusted per body weight)											
F ₁ parental males	↔	↔	↔	↔	↔	↔	↑2 days	4450 [754]	3397 [576]	3252 [551]	2445 [414]
F ₁ retained males	↔	↔	↓0.6 days	↔	↔	↔	↑1.8 days	4288 [727]	3375 [572]	2897 [491]	2145 [364]
Body weight on day of vaginal opening in F ₁	↔	↔	↔	↔	↔	↔	↓22%	3076 [521]	1281 [217]	3294 [558]	1972 [334]
Age of vaginal opening adjusted for PND 21 body weight ^b	↔	↔	↔	↔	↔	↔	↓2.4	3501 [593]	2953 [501]	3404 [577]	2419 [410]

^aBased on numbers of animals listed in data tables, it appears that statistical analyses in live animals prior to or on PND 21 considered the litter as the statistical unit, but statistical analyses conducted at PND 21 necropsy considered the individual pup as the statistical unit.

^bNot considered treatment related by study authors

^cEffect was not discussed by study authors but it is unlikely related to treatment.

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4.2.3.3 Fish and invertebrates

Kwak et al. (461), supported by the Korean Ministry of the Environment, exposed adult male swordtail fish (*Xiphophorus helleri*) to bisphenol A 0, 0.4, 2, or 10 ppm [mg/L] for 72 hours (n = 20 fish/group). [Nonylphenol was also studied but will not be discussed here.] At the end of the exposure period, the fish were killed and livers were removed for measurement of vitellogenin. Testes of 10 fish/group were processed for flow cytometry by preparation of single cell suspensions stained with annexin V-fluorescein isothiocyanate and propidium iodide to detect necrosis and apoptosis. TUNEL staining was used to confirm apoptosis in testis sections. In a second experiment, juvenile male fish (30 days old) were exposed to bisphenol A in water at 0, 0.2, 2 and 20 ppb [µg/L] for 60 days, after which body length and sword length were measured. [The sword is a portion of the caudal fin that elongates as a secondary sex characteristic.] Statistical analysis used ANOVA followed by least significant difference test. Hepatic vitellogenin was increased by bisphenol A [data were not shown]. Apoptosis was increased in testes from fish exposed to bisphenol A at 10 ppm [mg/L] by TUNEL assay. [Flow cytometry was said to be more sensitive, but data did not appear to have been statistically analyzed.] Sword growth was decreased by bisphenol A exposure in a concentration-dependent manner, with statistically significant decreases from control at 2 and 20 ppb [µg/L]. The authors concluded that bisphenol A at 20 ppb decreases sword growth and that reproductive impairment occurs in a concentration-dependent manner.

Strengths/Weaknesses: This study of bisphenol A is consistent with previous reports on the effects of estrogenic compounds in fish (vitellogenin production and changes secondary sex characteristics). It is unclear exactly how these fish were maintained prior to exposure and during the long-term exposure. Bisphenol A concentrations in the test waters were not determined and only 3 concentrations of bisphenol A were used.

Utility (Adequacy) for CERHR Evaluation Process: Of note is the classic dose response obtained in this apparently sensitive model. Given the absence of confirmation of exposure conditions and that this is a fish species immersed in the test agent, this study is not useful in the evaluation.

Sohoni et al. (462), supported by the Society of the Plastics Industry, exposed adult (122-day-old) fathead minnows (*Pimephales promelas*) to bisphenol A in water at 0, 1, 16, 160, and 640 µg/L (n = 60/group). Actual concentrations were 70–96% of nominal concentrations. After 42 days of exposure, 15 fish/group were killed for evaluation of somatic growth, relative gonad weight, plasma vitellogenin, and histologic assessment of the testis. Eight breeding pairs/group were segregated for continued exposure for 123 days. Eggs were removed and counted daily. On 2 occasions, eggs were continued in the same bisphenol A concentration as their parents and the percent hatching was assessed 4 days after fertilization. The remaining adult fish were killed after 71 days of exposure for evaluation of somatic growth, relative gonad weight, and histologic assessment of the gonad. Data were analyzed using 2-way ANOVA and Dunnett test or Kruskal-Wallis and Dunn multiple method test. Linear regression was used to evaluate the relationship between bisphenol A concentration and growth. There were no significant long-term effects of treatment on growth of female fish, but male fish showed an inverse relationship between bisphenol A concentration and growth with significant decrements in length and weight on pair-wise comparison at bisphenol A concentrations of 640 and 1280 µg/L. Relative gonad weight was also decreased in males and females at these bisphenol A concentrations. Plasma vitellogenin was increased in females beginning at bisphenol A concentrations of 640 µg/L and in males beginning at 160 µg/L. A delay in spermatogenesis was suggested by an increase in spermatogonia or spermatocytes and a decrease in spermatozoa in testes beginning at a bisphenol A concentration of 16 µg/L. There were no intersex gonads and no treatment-related changes in ovarian histopathology. The number of eggs spawned per female was lower in the control than the treatment groups and attributed by the authors to an unexplained problem in one of the control tanks. The 1280 µg/L bisphenol A concentration resulted in failure of 7 out of 8 females to produce any eggs. Hatching was impaired in eggs exposed to bisphenol A concentrations of 640 and 1280 µg/L. The authors noted that the bisphenol A concentrations resulting in impairment of somatic growth and reproductive success were only 7-fold lower than the 96-hour median

4.0 Reproductive Toxicity Data

1 lethal concentration, and concluded that the reproductive effects may have been the result of sublethal
2 generalized toxicity rather than effects mediated through the endocrine axis.

3
4 **Strengths/Weaknesses:** This study was well-conducted with multiple dose levels and concentrations in the
5 test water were confirmed. “General toxicity” was identified and good histology was used. The conclusions
6 regarding weak estrogenic activity were appropriate at 160 µg/L and higher. Other effects were likely due to
7 general toxicity. A classic dose response was noted.

8
9 **Utility (Adequacy) for CERHR Evaluation Process:** Fish are apparently a sensitive model for assessment
10 of responses to weak estrogenic compounds. Given that this study evaluated a fish species, it is not useful in
11 the evaluation.

12
13 **Kang et al. (463)**, supported by the Japanese Ministry of the Environment, exposed adult (4-month-old)
14 breeding pairs of medaka (*Oryzias latipes*) to bisphenol A (>99% purity) in the water at 0, 1000, or 4000 µg/L
15 for 3 weeks. Bisphenol A concentrations during the exposure period were 78–86% of nominal concentrations.
16 Thirty-two pairs of fish had been selected for exposure during an acclimatization period based on their
17 capacity to spawn daily, with the production of ≥15 eggs/day and 90% fertility. During the exposure period,
18 eggs were collected daily and assessed for fertility. Fertilized eggs collected on the last 3 days of the exposure
19 period were permitted to develop in untreated water, and 60 larvae/group were grown for 60 days after
20 hatching to assess normalcy of development. The parent fish were killed at the end of the treatment period for
21 evaluation of external sex characteristics and for histologic assessment of the gonads. Hepatic vitellogenin
22 was also assessed. Statistical comparisons of egg number were made using ANCOVA with female body
23 weight as a covariate. Fertility, growth endpoints, and hepatic vitellogenin data were analyzed with ANOVA
24 or Kruskal-Wallis test with post hoc Dunnett or Mann-Whitney *U* test. There were no treatment effects on egg
25 number, fertility, mortality, relative gonad weight, or relative liver weight in the adult fish. Ovarian tissue was
26 found in the testis in some males in all bisphenol A-treated groups, although normal testicular tissue with
27 apparently normal spermatogenesis was also found. Hepatic vitellogenin was increased in male fish in the
28 high-dose group to control female levels. There were no treatment-related alterations in hepatic vitellogenin in
29 female fish. Offspring at 60 days of age did not demonstrate treatment-related alterations in survival, growth,
30 or secondary sex characteristics. The sex ratio was not significantly different in offspring of parents exposed
31 to bisphenol A, although the authors noted that the low-dose group had a numerical deficit of males (41%
32 males compared to 50% in the controls). The authors concluded that although bisphenol A increased hepatic
33 vitellogenin in males and produced an intersex gonad, there were no adverse effects on reproductive capacity
34 or the normalcy of offspring.

35
36 **Strengths/Weaknesses:** This appears to have been a well conducted study. The bisphenol A findings are
37 consistent with the work of others, using sensitive endpoints in fish such as vitellogenin production. Given the
38 nature of the intersex gonad observation, it should be considered as adverse even though the severity was not
39 sufficient to induce decreases in reproductive capacity under the conditions tested.

40
41 **Utility (Adequacy) for CERHR Evaluation Process:** This study indicates that bisphenol A is able to induce
42 vitellogenin in male fish and intersex gonads. This study exhibited classic dose responses in the affected
43 endpoints. Because this study was conducted in fish, it is not useful in the evaluation.

44
45 **Lahnsteiner et al. (464)**, supported by the Austrian Federal Ministry of Agriculture, Forestry, Environment,
46 and Water Management, examined the effects of bisphenol A exposure on reproduction of male and female
47 brown trout (*Salmo trutta f. fario*). Fish were caught and acclimated for 2 weeks prior to starting the study.
48 Ten males/group and 6 females/group were exposed in a flow-through system to bisphenol A at 0 (DMSO
49 vehicle), 1.75, 2.4, or 5.00 µg/L beginning in the late prespawning period and continuing through the
50 remainder of the spawning season. The bisphenol A concentrations selected were said to occur in the Austrian
51 water system. Endpoints examined included time point of spawning, sperm count and motility, ability of

4.0 Reproductive Toxicity Data

1 sperm to fertilize eggs from non-treated females, and numbers and viability of eggs produced by treated
2 females. Statistical analyses included ANOVA and Tukey *b* post hoc test.

3
4 Throughout the entire spawning period, only 1 male in the high bisphenol A dose group produced semen and
5 it was of low quality as indicated by significantly reduced sperm density, motility rate, swimming velocity,
6 and fertility. In the low- and mid-dose groups, sperm density was significantly reduced in the early spawning
7 period but was not affected in the mid or end part of the spawning period. Additional significant effects
8 observed in the low-dose group included decreased sperm motility in the early spawning period, reduced
9 swimming velocity in the early and middle spawning period, and increased circular motion and decreased
10 linear motion in the middle of the spawning period. In the mid-dose group, sperm motility and swimming
11 velocity were significantly decreased in the early and mid-spawning period, and a significant increase in
12 circular motion and a decrease in linear motion occurred in the mid and late part of the spawning period. The
13 study authors interpreted the sperm effects as representing a 4-week delay in spawning. Fertility of males in
14 the low- and mid-dose group was not affected by bisphenol A treatment. In females, no eggs were produced
15 by fish in the high-dose group. In all other dose groups, there were no significant effects on egg volume,
16 viability, mass, mass increase during hardening, or on numbers of eggs produced by females. However,
17 ovulation was delayed by 2 weeks in the low-dose group and by 3 weeks in the mid-dose group. The study
18 authors concluded that exposure of trout to bisphenol A resulted in negative effects on semen and egg quality.

19
20 **Strengths/Weaknesses:** In this study of fish, alterations in sperm motility were observed consistent with
21 those observed in mice. Fertility effects in the female were also similar to those observed in other species.
22 Weaknesses include a failure to determine the actual bisphenol A concentrations in the test system, the
23 narrow dose range examined (1.75 to 5 µg/L), and the small number of fish/dose level assessed.

24
25 **Utility (Adequacy) of CERHR Evaluation Process:** This study suggests that fish are sensitive to bisphenol
26 A-induced abnormalities in reproductive endpoints. Because this study was conducted in fish, it is not useful
27 in the evaluation.

28
29 **Ortiz-Zarragoitia and Cajaraville (465),** supported by the European Commission, examined the effects of
30 bisphenol A exposure on the reproductive and digestive systems of adult blue mussels. For a period of 3
31 weeks, mussels were exposed to bisphenol A in acetone vehicle at 0 or 50 ppb [µg/L]. Additional compounds
32 were also tested but will not be discussed. Ten mussels/sex/group were examined at the end of the exposure
33 period. The digestive gland was examined for volume of peroxisomes and peroxisomal proliferation. Gonads
34 were histologically evaluated and assessed for alkali-labile phosphate level, a vitellogenin-like protein that is
35 a possible biomarker of endocrine disruption. Statistical analyses included ANOVA followed by Duncan post
36 hoc test, Kruskal-Wallis, and Mann-Whitney *U* test. Bisphenol A had no effect on gonadal development,
37 gonadal alkali-labile phosphate levels, or digestive gland peroxisomal proliferation or peroxisomal volume.
38 However, observations of follicular brown cell aggregates and gonadal hemocyte infiltration in 35% of male
39 and female mussels indicated severe gamete resorption.

40
41 **Strengths/Weaknesses:** This study evaluated bisphenol A-induced alterations in several reproductive
42 endpoints in adult mussels. Severe gamete resorption was observed. Weaknesses include the failure to
43 confirm bisphenol A concentrations in the test water and the use of only 1 concentration.

44
45 **Utility (Adequacy) for CERHR Evaluation Process:** Because this study was conducted in the mussel, it is
46 not useful in the evaluation.

4.3 Utility of Reproductive Toxicity Data

4.3.1 Human

There are 2 studies that measured serum bisphenol A in healthy women, women with polycystic ovary syndrome, and healthy men and evaluated correlations with serum gonadotropins, prolactin, testosterone, and other androgens. No fertility endpoints were included in these studies. Another study compared serum bisphenol A values in women with a history of recurrent miscarriage and women without a pregnancy history. Due to flaws in design and analysis, these 3 studies were considered to have low utility in the evaluation process. A study of 42 men occupationally exposed to bisphenol A diglycidyl ether and 42 unexposed men evaluated the relationship between urinary levels of bisphenol A and plasma LH, FSH, and free testosterone. No fertility endpoints were evaluated.

4.3.2 Experimental animal

Female reproductive toxicity testing using multiple dose levels has been evaluated in 2 rat, 1 mouse, and 1 gerbil study. Endpoints affected in these studies included brain progesterone receptor, estrous cyclicity, resorptions, and social sniffing. Male reproductive toxicity testing using multiple dose levels has been evaluated in 7 rat and 2 mouse studies. Affected endpoints in males included reproductive organ weight and histology, serum testosterone, daily sperm production, sperm motility, sperm concentration, percent pregnant females after mating, and females with resorptions after mating. There are 3 multigeneration tests, 2 in rats and 1 in mice, involving gavage or dietary treatments with bisphenol A with dose levels as low as 0.0009 mg/kg bw/day. There are also 2 reproductive assessments by continuous breeding, 1 of which involved subcutaneous implants for bisphenol A delivery and 1 of which used dietary administration with a low dose level of ~437.5 mg/kg bw/day.

4.4 Summary of Reproductive Toxicity Data

4.4.1 Human

Human reproductive studies are summarized in Table 114. Two papers from Takeuchi et al. (64, 65) suggested a relationship between serum bisphenol A concentration and serum testosterone (total and free). Subjects included women with and without polycystic ovary syndrome and women with and without obesity. One study included men. Although these studies used ELISA, which is inferior to HPLC, in the estimation of serum bisphenol A levels, significant correlations were demonstrated, leading the authors to speculate that androgens may affect bisphenol A metabolism. The authors did explore differences in exposure as a possible alternative explanation for their observations.

A study of 45 women with ≥ 3 consecutive spontaneous abortions found higher mean serum bisphenol A levels than in a group of 32 women without a history of pregnancy (67). The groups were unlikely to have been comparable in occupational and other potentially important factors, and the comparison of non-transformed means was inappropriate due to the skewness of the distribution of bisphenol A values. The study was useful only for showing that a few women with a history of recurrent spontaneous abortion had high bisphenol A levels. Some of the women also had high levels of anti-nuclear antibodies.

A study of 42 men occupationally exposed to an epoxy hardening agent containing bisphenol A diglycidyl ether found higher urinary bisphenol A concentrations, corrected for creatinine, than were found in 42 men who worked in the same factory but did not have known exposure to the hardening agent (82). There were no detected differences between the worker groups in plasma testosterone or LH, but plasma FSH was lower in workers exposed than in workers not exposed to the hardening agent. A significant correlation was noted between urinary bisphenol A concentration and decreased FSH when adjusted for alcohol intake. An association between serum bisphenol A and decreased FSH had also been reported by Takeuchi and Tsumi (65), although that association was not statistically significant.

1 **Table 114. Summary of Human Reproductive Toxicity Studies**

Subjects	Main findings	Reference	
Healthy women (n = 14)	Serum bisphenol A, ng/mL, mean ± SEM 0.64 ± 0.10	Takeuchi and Tsutsumi (65)	
Healthy men (n = 11)	1.49 ± 0.11		
Women with polycystic ovary syndrome	1.04 ± 0.10		
	Serum bisphenol A significantly correlated with total and free testosterone in all subjects and in all women.		
Non-obese cycling women (n=19)	Non-obese women with polycystic ovary syndrome had serum bisphenol A levels 48% higher than those in non-obese cycling women. Bisphenol A levels were 65% higher in obese women with polycystic ovary syndrome and 46% higher in obese cycling women than in non-obese cycling women.	Takeuchi et al. (64).	
Obese cycling women (n = 7)			
Hyperprolactinemic women (n = 7)			
Women with hypothalamic amenorrhea (n = 21)			
Non-obese women with polycystic ovary syndrome (n = 13)	Significant positive correlations were found between serum bisphenol A levels and serum androgens (total and free testosterone, androstenedione, dehydroepiandrosterone sulfate).		
Obese women with polycystic ovary syndrome (n = 6)			
	Serum bisphenol A, ng/mL, mean ± SD		
Women with ≥3 consecutive spontaneous abortions (n = 45)	2.59 ± 5.23	[Medians for the 2 groups were similar and the differences in means may have been due to skewing of the data. The utility of this study was low.]	
Healthy hospital worker with no history of pregnancy (n = 32)	0.77 ± 0.38		
	Urinary bisphenol A, μmol/mol creatinine, median	Plasma FSH, mIU/mL, median	Hanaoka et al. (82)
Men using bisphenol A diglycidyl ether (n = 42)	1.06	5.3	
Men not using bisphenol A diglycidyl ether (n = 42)	0.52	7.6	

2

3 *4.4.2 Experimental animal*

4 Reproductive studies using single dose levels of bisphenol A are summarized in Table 115. The lowest dose
5 level at which an effect was seen in these studies was 0.04 mg/kg/day fed to female rats during pregnancy and
6 lactation and resulting in a decreased duration of licking/grooming pups (420). Female reproductive studies
7 using multiple dose levels are summarized in Table 116. The lowest effect level was 0.002 mg/kg bw/day fed
8 to gerbils during 21 days of cohabitation (425). Limitations of this study included a lack of dose response,
9 which precluded calculation of a benchmark dose, and the absence of supporting data in this species from
10 other laboratories. Male reproductive studies are summarized in Table 117. The lowest effect level was
11 0.0002 mg/kg bw/day given orally (gavage assumed) in olive oil to Wistar rats for 45 days (435, 436).
12 Treated males showed a reduction in relative testis weight, relative epididymis weight, and epididymal sperm
13 motility and an increase in relative ventral prostate weight. Dose-related decreases in activity of superoxide
14 dismutase, catalase, glutathione reductase, and glutathione peroxidase and dose-related increases in hydrogen
15 peroxide generation and lipid peroxidation were seen in sperm at all dose levels. The study authors concluded

4.0 Reproductive Toxicity Data

1 that adverse effects of bisphenol A on the male reproductive system may be due to oxidative stress. The
2 utility of this study was limited by the use of olive oil vehicle without assessment of the stability or reactivity
3 of bisphenol A in the vehicle. In addition, only 6 rats/group were used in this study.
4

5 Multigeneration and continuous breeding studies are summarized in Table 118. The reproductive assessments
6 by continuous breeding included a study using subcutaneous administration (456, 457) and a study using very
7 high dose levels (458), and these studies are not the most informative for reproductive risk assessment. In a
8 multigeneration study, CD rats did not show statistically significant or dose-related reproductive effects over
9 2 generations with bisphenol A gavage doses of 0.0002, 0.002, 0.020, or 0.200 mg/kg bw/day (292). In
10 Sprague Dawley rats treated for 3 generations, adverse reproductive effects consisted of decreased F₁
11 epididymal sperm concentration, decreased F₃ daily sperm production, decreased live pups/litter, decreased
12 pup body weight, and advanced vaginal opening at an average dose level of 475 mg/kg bw/day. Advancement
13 of preputial separation was seen in F₁ males at an average dose level of 47.5 mg/kg bw/day (293, 411). In CD-
14 1 mice given bisphenol A for 2 generations in the diet at dose levels as low as ~0.003 mg/kg bw/day, the most
15 sensitive effect was a reduction in F₂ seminal vesicle weight relative to brain weight at 50 mg/kg bw/day.
16 Effects on F₀ epididymal sperm concentration, gestation length, and relative testis weight occurred at 600
17 mg/kg/day, the next highest dose level (376).
18

19 A summary of LH and testosterone effects observed in humans and in bisphenol A-exposed experimental
20 animals is included in Table 119.
21

22 **Questions for the Expert Panel**

23 Are human data sufficient for an evaluation of the male or female reproductive toxicity of bisphenol A?

24 If so, what are the relevant exposure conditions and endpoints?

25 Are experimental animal data sufficient for an evaluation of the male or female reproductive toxicity of
26 bisphenol A?

27 If so, what are the relevant experimental animal models, exposure conditions, and endpoints?

28 If the experimental animal data are sufficient for an evaluation, are the data assumed relevant, relevant, or not
29 relevant?

30 Note: The definitions of the term sufficient and the terms assumed relevant, relevant, and not relevant are in
31 the CERHR guidelines at <http://cerhr.niehs.nih.gov/news/guidelines.html>.

4.0 Reproductive Toxicity Data

1 **Table 115. Reproductive studies using single dose levels of bisphenol A**

Model	Treatment, mg/kg bw/day	Endpoint	Reference
<i>Female</i>			
Wistar rat, ovariectomized	~40 × 1 sc	Altered progesterone receptor mRNA in different brain regions	Funabashi et al. (416, 419)
Sprague Dawley rat	0.04 fed during pregnancy and lactation	↓Duration of licking/grooming pups	Della Seta et al. (420)
<i>Male</i>			
Wistar or Holtzman SD rat	~200, dietary	No effect on reproductive organ histopathology, daily sperm production, epididymal sperm reserves, or serum testosterone	Takahashi and Oishi (438)
Wistar rat	200, sc	↓Terminal body weight, absolute and relative reproductive organ weight; altered testicular histopathology	Takahashi and Oishi (438)
CD-1 (ICR) mouse	~400, dietary	↑Absolute testis weight, ↓absolute epididymis weight. No effect on testis histopathology, epididymal sperm reserves, daily sperm production, or serum testosterone.	Takahashi and Oishi (438)
C57BL/6CrSlc mouse	~400, dietary	No effect on reproductive organ weights. No effect on testis histopathology, epididymal sperm reserves, daily sperm production, or serum testosterone.	Takahashi and Oishi (438)

2

3 **Table 116. Female reproductive studies using multiple dose levels**

Model and treatment (mg/kg bw/day)	Endpoint	Bisphenol A dose level (mg/kg bw/day)				Reference
		NOAEL	LOAEL	BMD ₁₀	BMDL ₁₀	
Wistar rat, ovariectomized, ~0.004, 0.04, 0.4, or 4 sc, single dose	↑Progesterone receptor in brain regions	0.04	0.4	Data presentation does not permit modeling.		Funabashi et al. (418)
CD rat gavaged with 40, 200, or 600/1000 × 28 days	Altered estrous cycle	Unclear	≤1000/600	Data presentation does not permit modeling.		Yamasaki et al. (128)
Swiss mouse, gavaged with 0.005, 0.025, or 0.1 × 28 days	↓Body weight ↑Resorptions	<0.005 0.005	0.005 0.025	Models unsatisfactory		Al-Hiyasat et al. (422)
Mongolian gerbil, fed 0.002 or 0.02 from 1 st through 21 st day of cohabitation	↑Social sniffing	<0.002	0.002	No dose response		Razzoli et al. (425)

4

4.0 Reproductive Toxicity Data

1 **Table 117. Male reproductive studies using multiple dose levels**

Model and treatment (mg/kg bw/day)	Endpoint	Bisphenol A dose level (mg/kg bw/day)						Reference
		NOAEL	LOAEL	BMD ₁₀	BMDL ₁₀	BMD _{1SD}	BMDL _{1SD}	
Wistar rat, 2 or 20 ip 4 days/week × 1 month	↓ ventral prostate weight	2	20	7	5	9	6	Takahashi and Oishi (438)
	↓Serum testosterone	2	20	3	2	16	9	
CD rat gavaged with 40, 200, or 600/1000 × 28 days	↓Relative ventral prostate weight	200	600/1000	Data presentation does not permit modeling.				Yamasaki et al. (128)
	↑Relative testis weight	200	600/1000	Data presentation does not permit modeling.				
F344 rat, 235, 466, 950 in diet	Histologic alterations in testis	<235	235	No dose response				Takahashi and Oishi (431)
	↓Preputial gland relative weight	<235	235	124	86	171	114	
Sprague Dawley rat, gavaged with 0.020, 0.200, 2, 20, or 200 × 6 days	↓Daily sperm production (absolute and per g testis)	<0.020	0.020	No dose response				Sakaue et al. (432)
Sprague Dawley rat, gavaged with 0.000002, 0.00002, 0.0002, 0.002, 0.020, 0.200, or 2 × 6 days	↓Daily sperm production (absolute and per g testis)	0.002	0.020	Data presentation does not permit modeling.				Sakaue et al. (432)
Sprague Dawley rat, gavaged with 0.020, 0.200, 2, 20, or 200 × 6 days	No effect on daily sperm production, sperm count or reproductive organ weight	200	>200					Ashby et al. (433)
Wistar rat, 0.0002, 0.002 or 0.020 orally × 45 days	↓Relative testis weight	<0.0002	0.0002	0.056	0.021	0.014	0.0087	Chitra et al. (435, 436)
	↓Relative epididymis weight	<0.0002	0.0002	0.011	0.0082	0.0069	0.0050	
	↑Relative ventral prostate weight	<0.0002	0.0002	0.014	0.0083	0.015	0.0089	
	↓Sperm motility	<0.0002	0.0002	Data presentation does not permit modeling.				
	↓Sperm count	0.0002	0.02	Data presentation does not permit modeling.				
Swiss mouse, gavaged with 0.005, 0.025, and 0.1 × 30 days	Sperm/testis	0.005	0.025	0.035	0.029	0.036	0.028	Al-Hiyasat et al. (442)
	Sperm/mg testis	0.005	0.025	0.027	0.023	0.029	0.023	
	Daily sperm production	0.005	0.025	0.035	0.029	0.036	0.028	
	No. sperm/epididymis	<0.005	0.005	0.033	0.026	0.040	0.030	
	Sperm/mg epididymis	0.005	0.025	0.033	0.025	0.053	0.038	
	Percent pregnant females	0.005	0.025	no dose response				
	Females with resorptions	<0.005	0.005	no dose response				
C57BL/6N mouse, gavaged with 0.002, 0.020, or 0.200 × 6 days	No effect on reproductive organ weight or epididymal sperm count	≥0.200	-					Nagao et al. (369)

4.0 Reproductive Toxicity Data

1 **Table 118. Multigeneration and continuous breeding studies**

Model and treatment	Endpoint	Bisphenol A dose level (mg/kg bw/day)						Reference
		NOAEL	LOAEL	BMD ₁₀	BMDL ₁₀	BMD _{1SD}	BMDL _{1SD}	
<i>Multigeneration</i>								
CD rat, gavaged with 0.0002, 0.002, 0.020, or 0.200 prior to mating and × 2 generations.	No significant or dose-related reproductive effects	≥0.200	-					Ema et al. (292)
Sprague Dawley rat, ~0.0009, 0.018, 0.27, 4.5, 45, or 450 (male) and ~0.001, 0.02, 0.3, 5, 50, or 500 (female) in diet × 3 filial generations	↓F ₁ epididymal sperm concentration ↓F ₃ daily sperm production ↓Live pups/litter ^b ↓Pup body weight ^b Advanced vaginal opening ^b Advanced F ₁ preputial separation	47.5 ^a 47.5 47.5 47.5 47.5 4.75	- 475 475 475 475 47.5	317 469 236 183 394 466	216 255 174 163 343 411	700 524 376 177 206 188	469 481 286 153 176 163	Tyl et al. (293, 411)
CD-1 mouse, ~0.003, 0.03, 0.3, 5, 50, or 600 in diet from 6 weeks of age × 2 filial generations	↓F ₀ epididymal sperm concentration ↑Gestation length ^b ↓Relative testis weight ^b ↓Seminal vesicle weight relative to brain weight, F ₂	50 50 50 30	600 600 600 50	567 3619 562 405	319 639 339 223	607 1144 905 1914	549 599 605 615	Tyl et al. (376)
<i>Reproductive assessment by continuous breeding</i>								
CD-1 mouse, ~2.4, 4.2, or 8.1 over 18 week continuous breeding period, sc implant	No adverse effects on fertility	≤8.1	-					NTP (456, 457)
CD-1 mouse, ~437.5, 875, or 1750 in feed over 14-week continuous breeding period	↓Litters/breeding pair ↓Relative epididymis weight ↓Relative seminal vesicle weight	437.5 <437.5 <437.5	875 437.5 437.5	1750 420 700	1295 262 508	1680 805 1155	1155 438 822	NTP (458)

^aDose levels expressed as a mean of the estimated male-female target dose levels

^bBenchmark doses are shown for the generation with the lowest values.

4.0 Reproductive Toxicity Data

Table 119. Summary of Blood LH and Testosterone Changes in Human and Experimental Animal Studies

Endpoints/protocol	LH effects ^a	Testosterone effects ^a	Reference
<i>Human studies</i>			
Blood bisphenol A and hormone levels in healthy men and women and women with polycystic ovaries	↔	Correlation between bisphenol A blood levels and free or total testosterone levels in males and females	Takeuchi and Tsutsumi (65)
Serum sex hormone and bisphenol A concentrations in women with ovarian dysfunction and obesity	↔	Correlation between bisphenol A blood levels and free or total testosterone	Takeuchi et al. (64)
Blood bisphenol A levels in workers compared to controls	↔	↔	Hanaoka et al. (82)
<i>Experimental animal studies with oral exposure</i>			
Adult male and females rats gavaged for 28 days	↔ at 40–1000 mg/kg bw/day	↔ at 40–1000 mg/kg bw/day	Yamasaki et al. (128)
Castrated adult male rats were gavaged for 7 days	↔ at 10–1000 mg/kg bw/day	↔ at 10–1000 mg/kg bw/day	Kim et al. (259)
Four-week-old male rats fed bisphenol A in diet for 44 or 60 days	Not examined	↔ at 235–950 mg/kg bw/day or 200 mg/kg bw/day	Takahashi and Oishi (431, 438)
Male rats were gavaged from PND 21 through 35	↓ at 0.0024 mg/kg bw/day but ↔ at higher doses (0.010–200 mg/kg bw/day)	↓ at 0.0024 mg/kg bw/day but ↔ at higher doses (0.010–200 mg/kg bw/day)	Akingbemi et al. (306)
Male rats gavaged from PND 21 through 90	↑ at 0.0024 mg/kg bw/day	↔ at 0.0024 mg/kg bw/day	Akingbemi et al. (306)
Rats dams given drinking water containing bisphenol A from GD 6 through the lactation period	↓ at 1.2 mg/kg bw/day in adult female offspring (males not examined)	Not examined	Rubin et al. (217)
Rat dams gavaged from GD 12 through PND 21	↔ at 0.0024 mg/kg bw/day in adult males	↔ at 0.0024 mg/kg bw/day in adult males	Akingbemi et al. (306)
Rat dams exposed to bisphenol A [route unclear] throughout gestation and lactation	↔ in adult offspring [dose level uncertain]	↔ in adult offspring [dose level uncertain]	Kubo et al. (310)
Rat dams exposed to bisphenol A [most likely through drinking water during gestation and lactation]	↔ in adult offspring [dose level uncertain]	↔ in adult offspring [dose level uncertain]	Kubo et al. (311)
Rat dams dosed by pipette during gestation and lactation	Not examined	↔ in adult male offspring at 0.040 mg/kg bw/day	Aloisi et al. (314)
Multiple generation gavage dosing study in rats	↓ in F ₀ adult females at 0.0002, 0.002, and 0.020 mg/kg bw/day but not at high dose (0.2 mg/kg bw/day); not considered treatment-related.	↔	Ema et al. (292)

4.0 Reproductive Toxicity Data

Endpoints/protocol	LH effects ^a	Testosterone effects ^a	Reference
Adult male mice given drinking water containing bisphenol A for 4 or 8 weeks	↔ at 0.14–13 mg/kg bw/day	↓ with 8-week exposure to 13 mg/kg bw/day	Takao et al. (441)
Four-week-old mice fed bisphenol A through diet for 2 months	Not examined	↔ at 400 mg/kg bw/day	Takahashi and Oishi (438)
Mouse dams dosed by pipette on GD 11–17	Not examined	↔ in adult offspring at 0.002 or 0.020 mg/kg bw/day	Kawai et al. (358)
<i>Experimental animal studies with parenteral exposure</i>			
Adult male rats sc injected for 2 weeks	↑ at 3 mg/kg bw/day; ↔ following GnRH challenge	↓ blood level and response to hCG challenge at 3 mg/kg bw/day	Tohei et al. (434)
Four-week-old male rats sc dosed on 4 days/week for 1 month.	Not examined	↔ at 200 mg/kg bw	Takahashi and Oishi (438)
Four-week-old male rats ip injected for 1 month.	Not examined	↓ at 20 mg/kg bw	Takahashi and Oishi (438)
Male rats sc injected on PND 2, 4, 6, 8, 10, and 12.	Not examined	↔ with bisphenol A alone at 4–20 mg/kg bw/day but ↓ when administered with GnRH antagonist	Rivas et al. (329)
Male rats sc injected on PND 2–12.	Not examined	↑ on PND 18 at 20–100 mg/kg bw/day but ↔ on PND 25, 35, or 90	Sharpe et al. (330)
Male rats sc injected on PND 0–9.	Not examined	↔ at 0.002–97 mg/kg bw/day on PND 10, 35, or 150	Kato et al. (335)
Female lambs im injected at 4–11 weeks of age; ovariectomy at 9 weeks of age.	↔ on blood levels during treatment; ↓ pulsatile secretion following treatment with 3.5 mg/kg bw biweekly	Not examined	Evans et al. (382)

↑, ↓ Statistically significant increase/decrease compared to controls; ↔ no statistically significant effects compared to controls

^aUnless otherwise stated, animals were examined immediately after the treatment period

5.0 SUMMARIES, CONCLUSIONS, AND CRITICAL DATA NEEDS

To be written at the Expert Panel Meeting

5.1 Summary and Conclusion of Reproductive and Developmental Hazards

5.2 Summary of Human Exposure

5.3 Overall Conclusions

5.4 Critical Data Needs

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