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U.S. HIGH PRODUCTION VOLUME (HPV)  
CHEMICAL CHALLENGE PROGRAM

ROBUST SUMMARY

2-Propanone, Reaction Products with Phenol (CAS RN 72162-28-8)

Prepared By:  
General Electric Company  
One Plastics Avenue  
Pittsfield, MA 01201

Prepared for:  
U.S. Environmental Protection Agency  
Washington, D.C., USA

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## **ROBUST SUMMARY**

### **PHYSICAL AND CHEMICAL DATA**

#### **1.0 MELTING POINT**

##### 1.0.1

Value: 150-157 °C  
Decomposition: Yes ☐ No ☒ Ambiguous ☐  
Sublimation: Yes ☐ No ☒ Ambiguous ☐  
Method: Not specified  
GLP: Yes ☐ No ☐ ? ☒  
Test Substance: Bisphenol A (CAS RN 80-05-7)  
References: As cited in the EU Risk Assessment:  
Sax, N.I. and Lewis, R.J. 1996. Sax's Dangerous Properties of Industrial Materials, 9th Edition. New York. London. Van Nostrand Reinhold.  
Pohanish, R.P. and Greene, S.A. 1996. Hazardous Materials Handbook. New York. London. Van Nostrand Rheinhold.  
IPCS. 1993. Bisphenol-A; International Chemical Safety Cards, 8th Series prepared by IPCS. International Programme on Chemical Safety (IPCS).  
Merck Index. 1989. The Merck Index. an Encyclopedia of Chemicals Drugs and Biologicals. 11th Edition. Budavari, S. et al. (eds), Rahway, N.J. Merck & Co.  
Hubbard, N.W. et al. 1948. J. Am. Chem. Soc. 70:3259-3261.  
Bayer Leverkusen. 1989.  
Ullmann's Encyclopedia of Industrial Chemistry. 1991. Encyclopedia of Industrial Chemistry. 5th completely revised edition. Gerhartz, W. (ed), Weinheim. VCH.  
Reliability: (Klimisch Code 2) Valid with restrictions.

##### 1.0.2

Value: See Below  
Decomposition: Yes ☐ No ☐ Ambiguous ☐ Not Applicable ☒  
Sublimation: Yes ☐ No ☐ Ambiguous ☐ Not Applicable ☒  
Method: Estimated  
GLP: Yes ☐ No ☒ ? ☐  
Test Substance: See Below  
Remarks: The EPIWIN model was not run with any physico-chemical property input values. The CAS Numbers were entered for the individual chemicals below and the model data used.

CAS No.	Name	Melting Point*
80-05-7	BPA	131.8 °C
13464-24-9	Linear Dimer I	149.8 °C
57244-54-9	Linear Dimer II	151.2 °C
10527-11-4	Cyclic Dimer	164.2 °C
2300-15-4	BPX-I - Linear Trimer of isopropenyl phenol	215.3 °C
287110-79-6	BPX-II	
1568-80-5	Spirobiindane	178.0 °C
837-08-1	o,p-BPA	131.8 °C
472-41-3	Chroman I	141.5 °C

\*Empty cells indicate that data are unavailable for that chemical/endpoint.

Reference: U.S. EPA (Environmental Protection Agency). 2005. EPI Suite™, Version 3.12; MPBPWIN Program, Version 1.41; PC-Computer software developed by EPA's Office of Pollution Prevention Toxics and Syracuse Research Corporation (SRC).

Reliability: (Klimisch Code 2) Valid with restrictions.

## 2.0 BOILING POINT

### 2.0.1

Value: 250-252°C  
 Pressure: 17 kPa  
 Decomposition: Yes [ ] No [ ] Ambiguous [X]  
 Method: Not specified  
 GLP: Yes [ ] No [ ] ? [X]  
 Test Substance: Bisphenol A (CAS RN 80-05-7)  
 Reference: Von Braun. 1925. ANN. 472:65.  
 Reliability: (Klimisch Code 2) Valid with restrictions.

### 2.0.2

Value: 360.5°C  
 Pressure: 101.3 kPa  
 Decomposition: Yes [X] No [ ] Ambiguous [ ]  
 Method: Not specified  
 GLP: Yes [ ] No [ ] ? [X]  
 Test Substance: Bisphenol A (CAS RN 80-05-7)  
 Reference: DIPPR. 1994. Physical and Thermodynamic Properties of Pure Chemicals Data Compilation. Taylor and Francis.  
 Reliability: (Klimisch Code 2) Valid with restrictions.

### 2.0.3

Value: See below  
 Pressure: 760 mm Hg  
 Decomposition: Yes [ ] No [ ] Ambiguous [ ] Not Applicable [X]  
 Method: Estimated  
 GLP: Yes [ ] No [X] ? [ ]  
 Test Substance: See below

Remarks: The EPIWIN model was not run with any physico-chemical property input values. The CAS Numbers were entered for the individual chemicals below and the model data used.

CAS No.	Name	Boiling Point*
80-05-7	BPA	363.5°C
13464-24-9	Linear Dimer I	394.3°C
57244-54-9	Linear Dimer II	398.8°C
10527-11-4	Cyclic Dimer	396.3°C
2300-15-4	BPX-I - Linear Trimer of isopropenyl phenol	505.7°C
287110-79-6	BPX-II	
1568-80-5	Spirobiindane	425.9°C
837-08-1	o,p-BPA	363.5°C
472-41-3	Chroman I	377.3°C

\*Empty cells indicate that data are unavailable for that chemical/endpoint.

Reference: U.S. EPA (Environmental Protection Agency). 2005. EPI Suite™, Version 3.12; MPBPWIN Program, Version 1.41; PC-Computer software developed by EPA's Office of Pollution Prevention Toxics and Syracuse Research Corporation (SRC).

Reliability: (Klimisch Code 2) Valid with restrictions.

### 3.0 VAPOR PRESSURE

#### 3.0.1

Value:  $5.3 \times 10^{-8}$  hPa @ 25°C  
 Decomposition: Yes ☐ No ☐ Ambiguous ☐ Not Applicable ☒  
 Method: Calculated ☒ Measured ☐  
 GLP: Yes ☐ No ☐ ? ☒  
 Test Substance: Bisphenol A (CAS RN 80-05-7)  
 Reference: Bayer AG. 1988. Manufacturers Safety Data Sheet.  
 Reliability: (Klimisch Code 2) Valid with restrictions

#### 3.0.2

Value: See below  
 Decomposition: Yes ☐ No ☐ Ambiguous ☐ Not Applicable ☒  
 Method: Calculated ☒ Measured ☐  
 GLP: Yes ☐ No ☒ ? ☐  
 Test Substance: See below  
 Remarks: The EPIWIN model was not run with any physico-chemical property input values. The CAS Numbers were entered for the individual chemicals below and the model data used.

CAS No.	Name	Vapor Pressure at 25°C*
80-05-7	BPA	$3.0 \times 10^{-7}$ hPa
13464-24-9	Linear Dimer I	$4.45 \times 10^{-8}$ hPa
57244-54-9	Linear Dimer II	$3.21 \times 10^{-8}$ hPa
10527-11-4	Cyclic Dimer	$2.69 \times 10^{-8}$ hPa
2300-15-4	BPX-I - Linear Trimer of isopropenyl phenol	$4.81 \times 10^{-12}$ hPa
287110-79-6	BPX-II	
1568-80-5	Spirobiindane	$2.70 \times 10^{-9}$ hPa
837-08-1	o,p-BPA	$5.08 \times 10^{-7}$ hPa
472-41-3	Chroman I	$5.98 \times 10^{-7}$ hPa

\*Empty cells indicate that data are unavailable for that chemical/endpoint.

Reference: U.S. EPA (Environmental Protection Agency). 2005. EPI Suite™, Version 3.12; MPBPWIN Program, Version 1.41; PC-Computer software developed by EPA's Office of Pollution Prevention Toxics and Syracuse Research Corporation (SRC).  
Reliability: (Klimisch Code 2) Valid with restrictions.

#### 4.0 PARTITION COEFFICIENT (LOG10Pow)

##### 4.0.1

Log K<sub>ow</sub>: 3.4  
Temperature: Not reported  
Method: Measured; method not specified  
GLP: Yes ☐ No ☐ ? ☐ Not applicable ☒  
Test Substance: Bisphenol A (CAS RN 80-05-7)  
Reference: Bayer AG. 1993. Material Safety Data Sheet.  
Reliability: (Klimisch Code 2) Valid with restrictions.

##### 4.0.2

Log K<sub>ow</sub>: See below  
Temperature: 25°C  
Method: Estimated  
GLP: Yes ☐ No ☒ ? ☐  
Test Substance: See below  
Remarks: The EPIWIN model was not run with any physico-chemical property input values. The CAS Numbers were entered for the individual chemicals below and the model data used.

CAS No.	Name	Partition Coefficient (Log Kow) at 25°C*
80-05-7	BPA	3.6
13464-24-9	Linear Dimer I	5.6
57244-54-9	Linear Dimer II	5.5
10527-11-4	Cyclic Dimer	5.0
2300-15-4	BPX-I - Linear Trimer of isopropenyl phenol	5.8
287110-79-6	BPX-II	
1568-80-5	Spirobiindane	6.3
837-08-1	o,p-BPA	3.6
472-41-3	Chroman I	5.0

\*Empty cells indicate that data are unavailable for that chemical/endpoint.

Reference: U.S. EPA (Environmental Protection Agency). 2005. EPI Suite™, Version 3.12; KOWWIN Program, Version 1.67; PC-Computer software developed by EPA's Office of Pollution Prevention Toxics and Syracuse Research Corporation (SRC).  
Reliability: (Klimisch Code 2) Valid with restrictions.

## 5.0 WATER SOLUBILITY

### 5.1 SOLUBILITY

5.1.1  
Value: 300 mg/L  
Temperature: Room temperature  
Description: Miscible [ ]; Of very high solubility [ ]; Of high solubility [ ]; Soluble [ ]; Slightly soluble [ ]; Of low solubility [ ]; Of very low solubility [ X ]; Not soluble [ ]  
Method: Not specified  
GLP: Yes [ ] No [ ] ? [X] Not applicable [ ]  
Test Substance: Bisphenol A (CAS RN 80-05-7)  
Reference: Bayer AG. 1988. Manufacturers Safety Data Sheet.  
Reliability: (Klimisch Code 2) Valid with restrictions.

5.1.2  
Value: See below  
Temperature: 25°C  
Description: Miscible [ ]; Of very high solubility [ ]; Of high solubility [ ]; Soluble [ ]; Slightly soluble [ ]; Of low solubility [ ]; Of very low solubility [ ]; Not soluble [ ]; Not Applicable [X]  
Method: Estimated  
GLP: Yes [ ] No [ ] ? [ ] Not applicable [X]  
Test Substance: See below  
Remarks: The EPIWIN model was not run with any physico-chemical property input values. The CAS Numbers were entered for the individual chemicals below and the model data used.



CAS No.	Name	Water Solubility (mg/L)*
80-05-7	BPA	173
13464-24-9	Linear Dimer I	1.3
57244-54-9	Linear Dimer II	1.5
10527-11-4	Cyclic Dimer	4.2
2300-15-4	BPX-I - Linear Trimer of isopropenyl phenol	0.2
1568-80-5	Spirobiindane	0.2
287110-79-6	BPX-II	
837-08-1	o,p-BPA	91.5
472-41-3	Chroman I	4.1

\*Empty cells indicate that data are unavailable for that chemical/endpoint.

Reference: U.S. EPA (Environmental Protection Agency). 2005. EPI Suite™, Version 3.12; WSKOWWIN Program, Version 1.41; PC-Computer software developed by EPA's Office of Pollution Prevention Toxics and Syracuse Research Corporation (SRC).  
Reliability: (Klimisch Code 2) Valid with restrictions.

## 5.2. pH VALUE, pKa VALUE

No studies were found.

## ENVIRONMENTAL FATE AND PATHWAYS

### 6.0 PHOTODEGRADATION

#### 6.0.1

Method: Calculated [X] Measured [ ]  
GLP: Yes [ ] No [X] ? [ ]  
Test Substance: Bisphenol A (CAS RN 80-05-7)  
Concentration: Not applicable  
Temperature °C: Not stated  
Direct photolysis: Not applicable  
Indirect photolysis: Not applicable  
Breakdown products: Not applicable  
Value: Overall OH Rate Constant ( $k_{\text{phot}}$ ) =  $80.577 \times 10^{-12} \text{ cm}^3/\text{molecule} \cdot \text{sec}$   
Half-life ( $t_{1/2}$ ) = 1.593 Hrs (12-hour day;  $1.5 \times 10^6 \text{ OH}/\text{cm}^3$ )  
Remarks: The EPIWIN model was not run with any physico-chemical property input values. The CAS Numbers were entered for the individual chemicals below and the model data used.  
Atmospheric photo-oxidation potential, mediated by reaction with hydroxyl radicals, was estimated using the submodel, Atmospheric Oxidation Potential for Windows (AOPWIN v.1.91, EPIWIN v3.12, US EPA, 2005). The SAR methods rely on structural features of the subject molecule. The model calculates a second-order half-life with units of  $\text{cm}^3/\text{molecules} \cdot \text{sec}$ . A pseudo-first order photo-degradation rate is in

turn based on the second order rate of reaction ( $\text{cm}^3/\text{molecules}\cdot\text{sec}$ ) with hydroxyl radicals ( $\text{HO}\cdot$ ), assuming first-order kinetics and an  $\text{HO}\cdot$  concentration of  $1.5 \times 10^6$  molecules/ $\text{cm}^3$  and 12 hours of daylight. Pseudo-first order half-lives ( $t_{1/2}$ ) were then calculated as follows:  

$$t_{1/2} = 0.693 / [k_{\text{phot}} \times \text{HO}\cdot \times 12\text{-hr} / 24\text{-hr}].$$

Reference: U.S. EPA (Environmental Protection Agency). 2005. EPI Suite™, Version 3.12; AOPWIN Program, Version 1.91; PC-Computer software developed by EPA's Office of Pollution Prevention Toxics and Syracuse Research Corporation (SRC).  
 Reliability: (Klimisch Code 2) Valid with restrictions.

#### 6.0.2

Method: Calculated [X] Measured [ ]  
 GLP: Yes [ ] No [X] ? [ ]  
 Test Substance: See below  
 Concentration: Not applicable  
 Temperature °C: Not stated  
 Direct photolysis: Not applicable  
 Indirect photolysis: Not applicable  
 Breakdown products: Not applicable  
 Value: See below  
 Remarks: The EPIWIN model was not run with any physico-chemical property input values. The CAS Numbers were entered for the individual chemicals below and the model data used.

CAS No.	Name	Overall OH Rate Constant ( $\text{cm}^3/\text{molecule}\cdot\text{sec}$ )*	Half-Life (Hours)*
80-05-7	BPA	$80.58 \times 10^{-12}$	1.59
13464-24-9	Linear Dimer I	$124.48 \times 10^{-12}$	1.03
57244-54-9	Linear Dimer II	$159.00 \times 10^{-12}$	0.81
10527-11-4	Cyclic Dimer	$129.75 \times 10^{-12}$	0.99
2300-15-4	BPX-I - Linear Trimer of isopropenyl phenol	$129.02 \times 10^{-12}$	1.00
287110-79-6	BPX-II		
1568-80-5	Spirobiindane	$178.93 \times 10^{-12}$	0.72
837-08-1	o,p-BPA	$80.58 \times 10^{-12}$	1.59
472-41-3	Chroman I	$71.06 \times 10^{-12}$	1.81

\*Empty cells indicate that data are unavailable for that chemical/endpoint.

Reference: U.S. EPA (Environmental Protection Agency). 2005. EPI Suite™, Version 3.12; AOPWIN Program, Version 1.91; PC-Computer software developed by EPA's Office of Pollution Prevention Toxics and Syracuse Research Corporation (SRC).  
 Reliability: (Klimisch Code 2) Valid with restrictions.

## 7.0 STABILITY IN WATER

Type: Abiotic (hydrolysis) [X]; biotic (sediment) [ ]  
Half life: Not applicable  
Degradation: Not applicable  
Method: Not applicable  
GLP: Yes [ ] No [X] ? [ ]  
Test Substance: Bisphenol A (CAS RN 80-05-7)  
Remarks: Bisphenol A has no hydrolyzable groups and, therefore, does not undergo hydrolysis.  
Reference: None – conclusion based on chemistry of Bisphenol A.  
Reliability: (Klimisch Code 2) Valid with restrictions.

## 8.0 TRANSPORT AND DISTRIBUTION BETWEEN ENVIRONMENTAL COMPARTMENTS, INCLUDING ESTIMATED ENVIRONMENTAL CONCENTRATIONS AND DISTRIBUTION PATHWAYS

### 8.1 THEORETICAL DISTRIBUTION (FUGACITY CALCULATION)

#### 8.1.1

Type: Fugacity model level III  
Media: Other: air, water, soil, sediment  
Method: Calculated [X] Measured [ ]  
GLP: Yes [ ] No [X] ? [ ]  
Test Substance: Bisphenol A (CAS RN 80-05-7)  
Value: Air < 0.01%  
Water = 11.9%  
Soil = 87.5%  
Sediment = 0.6%  
Remarks: The EPIWIN model was not run with any physico-chemical property input values. The CAS Numbers were entered for the individual chemicals below and the model data used.  
Default values were assumed for environmental compartment descriptions, dimensions, and properties, advective and dispersive properties.

Emissions were assumed to be equally to air, water and soil (Model run with 1000 kg/hr emissions each to air, water and soil.).

Air:  $t_{1/2} = 3.185$  hr  
Water:  $t_{1/2} = 900$  hr  
Soil:  $t_{1/2} = 1800$  hr  
Sediment:  $t_{1/2} = 8100$  hr

Reference: U.S. EPA (Environmental Protection Agency). 2005. EPI Suite™, Version 3.12; Level III Fugacity Model; PC-Computer software developed by EPA's Office of Pollution Prevention Toxics and Syracuse Research Corporation (SRC).

Reliability: (Klimisch Code 2) Valid with restrictions, calculated value.

#### 8.1.2

Type: Fugacity model level III  
Media: Other: air, water, soil, sediment  
Method: Calculated [X] Measured [ ]  
GLP: Yes [ ] No [X] ? [ ]  
Test Substance: See below  
Value: See below  
Remarks: The EPIWIN model was not run with any physico-chemical property input values. The CAS Numbers were entered for the individual chemicals below and the model data used. Model run with 1000 kg/hr emissions each to air, water and soil.

CAS No.	Name	Mass Amount (%)*			
		Air	Water	Soil	Sediment
80-05-7	BPA	<0.01	11.9	87.5	0.57
13464-24-9	Linear Dimer I	<0.01	5.19	58.3	36.5
57244-54-9	Linear Dimer II	<0.01	5.63	60.4	33.9
10527-11-4	Cyclic Dimer	<0.01	8.38	74.4	17.2
2300-15-4	BPX-I - Linear Trimer of isopropenyl phenol	<0.01	3.05	54.6	42.3
287110-79-6	BPX-II				
1568-80-5	Spirobiindane	<0.01	1.9	46.8	51.3
837-08-1	o,p-BPA	<0.01	11.4	87.4	1.15
472-41-3	Chroman I	<0.1	6.87	73.9	19.2

\*Empty cells indicate that data are unavailable for that chemical/endpoint.

Reference: U.S. EPA (Environmental Protection Agency). 2005. EPI Suite™, Version 3.12; Level III Fugacity Model; PC-Computer software developed by EPA's Office of Pollution Prevention Toxics and Syracuse Research Corporation (SRC).

Reliability: (Klimisch Code 2) Valid with restrictions.

#### 8.1.3

Type: Fugacity model level III  
Media: Other: air, water, soil, sediment  
Method: Calculated [X] Measured [ ]  
GLP: Yes [ ] No [X] ? [ ]  
Test Substance: Bisphenol A (CAS RN 80-05-7)  
Value: Air < 0.01%  
Water = 3.06%  
Soil = 96.8%  
Sediment = 0.145  
Remarks: The EPIWIN model was not run with any physico-chemical property input values. The CAS Numbers were entered for the individual chemicals below and the model data used.

Default values were assumed for environmental compartment descriptions, dimensions, and properties, advective and dispersive properties.

Emissions were assumed to be equally to air, water and soil (Model run with 1000 kg/hr emissions to air.).

Air:  $t_{1/2} = 3.185$  hr  
Water:  $t_{1/2} = 900$  hr  
Soil:  $t_{1/2} = 1800$  hr  
Sediment:  $t_{1/2} = 8100$  hr

Reference: U.S. EPA (Environmental Protection Agency). 2005. EPI Suite™, Version 3.12; Level III Fugacity Model; PC-Computer software developed by EPA's Office of Pollution Prevention Toxics and Syracuse Research Corporation (SRC).  
Reliability: (Klimisch Code 2) Valid with restrictions.

#### 8.1.4

Type: Fugacity model level III  
Media: Other: air, water, soil, sediment  
Method: Calculated [X] Measured [ ]  
GLP: Yes [ ] No [X] ? [ ]  
Test Substance: Bisphenol A (CAS RN 80-05-7)  
Value: Air < 0.01%  
Water = 95.5%  
Soil < 0.1%  
Sediment = 4.53

Remarks: The EPIWIN model was not run with any physico-chemical property input values. The CAS Numbers were entered for the individual chemicals below and the model data used. Default values were assumed for environmental compartment descriptions, dimensions, and properties, advective and dispersive properties.

Emissions were assumed to be equally to air, water and soil (Model run with 1000 kg/hr emissions to air.).

Air:  $t_{1/2} = 3.185$  hr  
Water:  $t_{1/2} = 900$  hr  
Soil:  $t_{1/2} = 1800$  hr  
Sediment:  $t_{1/2} = 8100$  hr

Reference: U.S. EPA (Environmental Protection Agency). 2005. EPI Suite™, Version 3.12; Level III Fugacity Model; PC-Computer software developed by EPA's Office of Pollution Prevention Toxics and Syracuse Research Corporation (SRC).  
Reliability: (Klimisch Code 2) Valid with restrictions.

8.1.5

Type: Fugacity model level III  
Media: Other: air, water, soil, sediment  
Method: Calculated [X] Measured [ ]  
GLP: Yes [ ] No [X] ? [ ]  
Test Substance: Bisphenol A (CAS RN 80-05-7)  
Value:

Air < 0.01%  
Water = 0.676%  
Soil < 99.3%  
Sediment = 0.03

Remarks: The EPIWIN model was not run with any physico-chemical property input values. The CAS Numbers were entered for the individual chemicals below and the model data used.

Default values were assumed for environmental compartment descriptions, dimensions, and properties, advective and dispersive properties.

Emissions were assumed to be equally to air, water and soil (Model run with 1000 kg/hr emissions to soil).

Air:  $t_{1/2} = 3.185$  hr  
Water:  $t_{1/2} = 900$  hr  
Soil:  $t_{1/2} = 1800$  hr  
Sediment:  $t_{1/2} = 8100$  hr

Reference: U.S. EPA (Environmental Protection Agency). 2005. EPI Suite™, Version 3.12; Level III Fugacity Model; PC-Computer software developed by EPA's Office of Pollution Prevention Toxics and Syracuse Research Corporation (SRC).

Reliability: (Klimisch Code 2) Valid with restrictions.

8.1.6

Type: Fugacity model level III  
Media: Other: air, water, soil, sediment  
Method: Calculated [X] Measured [ ]  
GLP: Yes [ ] No [X] ? [ ]  
Test Substance: Bisphenol A (CAS RN 80-05-7)  
Value:

Air < 0.01%  
Water = 21.7%  
Soil = 77.3%  
Sediment = 1.03

Remarks: The EPIWIN model was not run with any physico-chemical property input values. The CAS Numbers were entered for the individual chemicals below and the model data used.

Default values were assumed for environmental compartment descriptions, dimensions, and properties, advective and dispersive properties.

Emissions were assumed to be equally to air, water and soil (Model run with 1000 kg/hr emissions to air and water).

Air:  $t_{1/2} = 3.185$  hr  
Water:  $t_{1/2} = 900$  hr  
Soil:  $t_{1/2} = 1800$  hr  
Sediment:  $t_{1/2} = 8100$  hr

Reference: U.S. EPA (Environmental Protection Agency). 2005. EPI Suite™, Version 3.12; Level III Fugacity Model; PC-Computer software developed by EPA's Office of Pollution Prevention Toxics and Syracuse Research Corporation (SRC).

Reliability: (Klimisch Code 2) Valid with restrictions.

#### 8.1.7

Type: Fugacity model level III  
Media: Other: air, water, soil, sediment  
Method: Calculated [X] Measured [ ]  
GLP: Yes [ ] No [X] ? [ ]  
Test Substance: Bisphenol A (CAS RN 80-05-7)  
Value: Air < 0.01%  
Water = 1.82%  
Soil = 98.1%  
Sediment = 0.09

Remarks: The EPIWIN model was not run with any physico-chemical property input values. The CAS Numbers were entered for the individual chemicals below and the model data used. Default values were assumed for environmental compartment descriptions, dimensions, and properties, advective and dispersive properties.

Emissions were assumed to be equally to air, water and soil (Model run with 1000 kg/hr emissions to air and soil).

Air:  $t_{1/2} = 3.185$  hr  
Water:  $t_{1/2} = 900$  hr  
Soil:  $t_{1/2} = 1800$  hr  
Sediment:  $t_{1/2} = 8100$  hr

Reference: U.S. EPA (Environmental Protection Agency). 2005. EPI Suite™, Version 3.12; Level III Fugacity Model; PC-Computer software developed by EPA's Office of Pollution Prevention Toxics and Syracuse Research Corporation (SRC).

Reliability: (Klimisch Code 2) Valid with restrictions.

#### 8.1.8

Type: Fugacity model level III  
Media: Other: air, water, soil, sediment  
Method: Calculated [X] Measured [ ]

GLP: Yes ☐ No ☒ ? ☐  
 Test Substance: Bisphenol A (CAS RN 80-05-7)  
 Value: Air < 0.01%  
 Water = 18.6%  
 Soil = 80.6%  
 Sediment = 0.881  
 Remarks: Default values were assumed for environmental compartment descriptions, dimensions, and properties, advective and dispersive properties.  
 Emissions were assumed to be equally to air, water and soil (Model run with 1000 kg/hr emissions to water and soil).  
 Air:  $t_{1/2} = 3.185$  hr  
 Water:  $t_{1/2} = 900$  hr  
 Soil:  $t_{1/2} = 1800$  hr  
 Sediment:  $t_{1/2} = 8100$  hr  
 Reference: U.S. EPA (Environmental Protection Agency). 2005. EPI Suite™, Version 3.12; Level III Fugacity Model; PC-Computer software developed by EPA's Office of Pollution Prevention Toxics and Syracuse Research Corporation (SRC).  
 Reliability: (Klimisch Code 2) Valid with restrictions.

## 9.0 BIODEGRADATION

Type: Aerobic ☒; Anaerobic ☐  
 Inoculum: Adapted ☐; Non-adapted ☒  
 Concentration of the chemical: 2.52 mg O<sub>2</sub>/mg as ThOD  
 Medium: Water ☒; Water-sediment ☐; Soil ☐; Sewage treatment ☐; Other ☐  
 Contact time: 28 days at 27.1 °C  
 Degradation: 73.0 – 80.1%  
 Result: Readily biodeg. ☒; Inherently biodeg. ☐; Other ☐  
 Kinetic of test substance: See below  
 Reference substance: Information not stated in EU Risk Assessment  
 Degradation Products: Yes ☐ No ☐ Not measured ☒  
 Method (Year): OECD Test Guideline 301F  
 GLP: Yes ☐ No ☐ ? ☒  
 Test Substance: Bisphenol A (99.7% purity)  
 Remarks: The theoretical oxygen demand (ThOD) of bisphenol A was calculated as 2.52 mg O<sub>2</sub>/mg. The inoculum used in the experiment consisted of activated sludge mixed liquor collected from a municipal sewage treatment plant. The experimental details followed the procedures detailed in the OECD 301F test to Good Laboratory Practice (GLP) standards. However, the temperature used in the experiment was 27.1°C, which is 2.1°C above the range of temperatures quoted in the OECD guidelines. The initial concentrations of bisphenol A used in the



	experiment were 7 mg/l and 25 mg/l. Oxygen consumption and CO <sub>2</sub> evolution were measured over 28 days and removal of dissolved organic carbon (DOC) from the biodegradation reactions was determined after 28 days. The time required for achieving 10% degradation for the bisphenol A ranged from 5.6 days (7 mg/l bisphenol A) to 6.1 days (25 mg/l bisphenol A) with biodegradation exceeding the 60% degradation level after an additional 3.5 days (7 mg/l bisphenol A) and 5.0 days (25 mg/l bisphenol A). Ten days following the defined lag periods biodegradation averaged 77.6% and 73.7% for the 7 mg/l and 25 mg/l reactions. The maximum degradation levels averaged 84.6% and 81.7% of the ThOD for the 7 mg/l and 25 mg/l reactions respectively after 28 days. The rate and extent of bisphenol A mineralisation observed indicate that bisphenol A can be classified as “readily biodegradable”. Evolution of CO <sub>2</sub> resulting from mineralisation of bisphenol A closely followed biodegradation of the compound as measured from oxygen consumption. Maximum yields of CO <sub>2</sub> ranged from 73.0% to 80.1% of ThCO <sub>2</sub> indicating nearly complete conversion of the added organic carbon to CO <sub>2</sub> .
Reference:	West, R.J. and Goodwin, P.A. 1997. Evaluation of Ready Biodegradability of Bisphenol A Using the OECD 301F: Manometric Respirometry Test. Dow Company Report. Study ID 971108A. As cited in the EU Risk Assessment.
Reliability:	(Klimisch Code 1) Valid without restrictions.

## **ECOTOXICOLOGICAL DATA**

### **10.0 ACUTE/PROLONGED TOXICITY TO FISH**

Type of Test:	Static <input checked="" type="checkbox"/> Semi-static <input type="checkbox"/> Flow-through <input type="checkbox"/> Other <input type="checkbox"/> Open-system <input type="checkbox"/> Closed-system <input type="checkbox"/>
Species:	Fathead minnow
Exposure Period:	96 hours
Results:	LC <sub>50</sub> (96h) 4.6 mg BPA/L NOAEC = Not reported LOAEC = Not reported
Analytical Monitoring:	Yes <input type="checkbox"/> No <input type="checkbox"/> ? <input checked="" type="checkbox"/>
Method:	ASTM E-35 21 (1992)
GLP:	Yes <input type="checkbox"/> No <input type="checkbox"/> ? <input checked="" type="checkbox"/>
Test Substance:	Bisphenol A (CAS RN 80-05-7)
Remarks:	<u>Test conditions:</u> The reported test conditions are summarized below:

Parameter	Test Condition
Test type	Flow
Duration	96 hours
Test organism	Fathead minnow ( <i>Pimephales promelas</i> )
Average weight $\pm$ SD & range	0.41 g
Average fork length $\pm$ SD & range	3.0 cm
Temperature	16.1 to 17.9 °C
Dissolved oxygen	>81% saturation
pH	7.6-8.0
Criterion for Effect	Death
Toxicity Endpoints	96-hour LD <sub>50</sub>

Results: LC<sub>50</sub> = 4.6 mg BPA/L

Reference: Alexander, H.C., Bartlett, E.A. and Boggs, G.U. 1985. Bisphenol A: Flow-Through Acute Toxicity to the Fathead Minnow. Dow Company Report: ES-764. As cited in the EU Risk Assessment.

Alexander, H.C., Dill, D.C., Smith, L.W., Guiney, P.D. and Dorn, P. 1988. Bisphenol A: Acute aquatic toxicity. Environmental Toxicology and Chemistry 7:19-26. As cited in the EU Risk Assessment.

Reliability: (Klimisch Code 1) Valid without restrictions.

#### 11.0 ACUTE TOXICITY TO AQUATIC INVERTEBRATES (E.G. DAPHNIA)

Type of Test: Static ☒ Semi-static ☐ Flow-through ☐ Other ☐

Open-system ☐ Closed-system ☐

Species: *Daphnia magna*

Exposure Period: 48 hours

Results: EC<sub>50</sub> (96h) 10.2 mg/L

NOEC = Not reported

LOEC > Not reported

Analytical Monitoring: Yes ☒ No ☐ ? ☐

Method: ASTM E-35.21

GLP: Yes ☐ No ☐ ? ☒

Test Substance: Bisphenol A (CAS RN 80-05-7)

Remarks: The reported test conditions are summarized below:

Parameter	Test Condition
Test type	Static
Duration of test	48 hours
Test organism	<i>Daphnia magna</i>
Age at Test Initiation	First instar
Test Solution Temperature	19.8-20.4°C
Dissolved Oxygen Concentration	>96%
pH	8.0-8.3
Criterion for effect	Immobilization
Calculated toxicity values	EC <sub>50</sub> at 48-h

Results: 24-hr EC<sub>50</sub>(immobilization) = 15.5 mg BPA/L (measured)  
48-hr EC<sub>50</sub>(immobilization) = 10.2 mg BPA/L (measured)

Reference: Alexander, H.C., Milazzo, D.P. and Boggs, G.U. 1985. Bisphenol A: Daphnid Static Acute Toxicity Test. Dow Company Report. ES-763. As cited in the EU Risk Assessment.  
Alexander, H.C., Dill, D.C., Smith, L.W., Guiney, P.D. and Dorn, P. 1988. Bisphenol A: Acute aquatic toxicity. Environmental Toxicology and Chemistry 7:19-26. As cited in the EU Risk Assessment.

Reliability: (Klimisch Code 1) Valid without restrictions.

## 12.0 TOXICITY TO AQUATIC PLANTS (E.G. ALGAE)

Species: Green algae (*Selenastrum capricornutum*)

End-point: Biomass [ X ] Growth rate [ ] Other [ ]

Exposure Period: 96 hours

Results: Growth:  
EC<sub>50</sub>(96h) = 2.7-3.1 mg/L

Analytical Monitoring: Yes [ ] No [ ] ? [X]

Method: Not reported

GLP: Yes [ ] No [ ] ? [X]

Test Substance: Bisphenol A (CAS RN 80-05-7)

Remarks: Test conditions were not reported. The percentage inhibition of cell count and cell volume was reported for the concentrations tested. From these data, it was possible to derive the EC<sub>1</sub>. using probit analysis.

Results: 96-hr EC<sub>50</sub> = 2.7-3.1 mg BPA/L  
Calculated EC<sub>10</sub> (By the EU RAR) = 1.36 mg BPA/L (cell count) and 1.68 mg BPA/L (cell volume)

Reference: Alexander, H.C., Dill, D.C., Milazzo, D.P. and Boggs, G.U. 1985. Bisphenol A: Algal Toxicity Test. Dow Company Report: ES-762. As cited in the EU Risk Assessment.  
Alexander, H.C., Dill, D.C., Smith, L.W., Guiney, P.D. and Dorn, P. 1988. Bisphenol A: Acute aquatic toxicity. Environmental Toxicology and Chemistry 7:19-26. As cited in the EU Risk Assessment.

Reliability: (Klimisch Code 1) Valid without restrictions.

## **HUMAN-HEALTH RELATED DATA**

### **13.0 ACUTE TOXICITY**

#### **13.1 ACUTE ORAL TOXICITY**

##### **13.1.1**

Type: LD<sub>0</sub> [ ] LD<sub>100</sub> [ ] LD<sub>50</sub> [X] LD<sub>L0</sub> [ ] Other [ ]  
Species/Strain: Rat/F344  
Sex: Male and female  
# Animals: 5/sex  
Vehicle: Not reported  
Value: LD<sub>50</sub> (males) = 4100 mg/kg  
LD<sub>50</sub> (females) = 3300 mg/kg  
Method: Not reported  
GLP: Yes [ ] No [ ] ? [X]  
Test Substance: Bisphenol A (CAS RN 80-05-7)  
Remarks: Rats (5 per sex per group) were given 2,150, 3,160, 4,640 or 6,810 mg/kg of bisphenol A.  
Results:

<b>Dose Level (mg/kg)</b>	<b>Male Mortality</b>	<b>Female Mortality</b>
2150	0/5	1/5
3160	1/5	4/5
4640	3/5	2/5
6810	5/5	5/5

The calculated LD<sub>50</sub> values were 4,100 mg/kg in males and 3,300 mg/kg in females. No other aspects of the study were reported.  
Reference: NTP. 1982. Carcinogenesis Bioassay of Bisphenol A (CAS No. 80-05-7) in F344 Rats B6C3F1 Mice (Feed Study). National Toxicology Program. Technical Report No. 215, Order No. PB82-184060 (NTIS), 1-116. As cited in the EU Risk Assessment.  
Reliability: (Klimisch Code 2) Valid with restrictions.

##### **13.1.2**

Type: LD<sub>0</sub> [ ] LD<sub>100</sub> [ ] LD<sub>50</sub> [X] LD<sub>L0</sub> [ ] Other [ ]  
Species/Strain: Rabbit  
Sex: Not reported  
# Animals: Not reported  
Vehicle: Not reported  
Value: LD<sub>50</sub> = 2230 mg/kg  
Method: Not reported  
GLP: Yes [ ] No [X] ? [ ]  
Test Substance: Bisphenol A (CAS RN 80-05-7)  
Remarks: A rabbit LD<sub>50</sub> value was obtained. No other details are reported.  
Results: LD<sub>50</sub> = 2230 mg/kg

Reference:	Mellon Institute of Industrial Research. 1948. The Acute and Subacute Toxicity of Diphenylol Propane. Study No. 11-13. Union Carbide Corporation Unpublished Report. As cited in the EU Risk Assessment.
Reliability:	(Klimisch Code 2) Valid with restrictions.
13.1.3	
Type:	LD <sub>0</sub> [ ] LD <sub>100</sub> [ ] LD <sub>50</sub> [X] LD <sub>L0</sub> [ ] Other [ ]
Species/Strain:	Rat/Crl:CD®(SD)BR
Sex:	Male and female
# Animals:	5/sex
Vehicle:	Corn oil
Value:	LD <sub>50</sub> > 5000 mg/kg
Method:	Not reported
GLP:	Yes [ ] No [ ] ? [X]
Test Substance:	Spirobiindane (CAS RN 1568-80-5)
Remarks:	Test article was administered once orally via gastric intubation to a single group of 5/sex Crl:CD®(SD)BR rats (221-245g) at a dose of 5000 mg/kg. Mortality, clinical observations, body weight changes and day 14 gross necropsy findings were evaluated.
Results:	There were no deaths, body weight changes or test article-related gross necropsy findings. Seven rats had red and/or yellow external matting. Abnormal excreta was noted for four animals. One female was hypoactive. All animals were normal by day 8 or earlier. The oral LD <sub>50</sub> was found to be >5000 mg/kg.
Reference:	Kern <sup>1</sup> , T., Hurley <sup>1</sup> , J., Wilmot <sup>1</sup> , A. and Dimond, S. 1997. Acute Toxicity Studies with Spirobiindane. General Electric Plastics, Pittsfield, MA, USA and <sup>1</sup> WIL Research Laboratories, Inc. Ashland, OH, USA.
Reliability:	(Klimisch Code 1) Valid without restrictions.

## 13.2 ACUTE INHALATION TOXICITY

Type:	LD <sub>0</sub> [ ]; LD <sub>100</sub> [ ]; LD <sub>50</sub> [X]; LD <sub>L0</sub> [ ]; Other [ ]
Species/strain:	Rat/Fischer F344
Sex:	Males and females
# animals:	10/sex
Value:	>170 mg/L
Method (Year):	Not reported.
GLP:	Yes [ ] No [ ] ? [X]
Test substance:	Bisphenol A (CAS RN 80-05-7)
Remarks:	Groups of 10 male and female Fischer F344 rats were exposed, whole body, to Bisphenol A dust at concentrations of 0 or 170 mg/m <sup>3</sup> .
Results:	The mass median aerodynamic diameter (MMAD) of the test substance was 3.9 micrometers (µm), and the exposure concentration used was the highest attainable by the investigators with the test system used. Half the animals were necropsied on the day following exposure and the rest on day 14. Microscopic examination was limited to the respiratory tract (nasal turbinates, larynx, trachea and lungs) and associated tissues. No

deaths occurred and therefore the LD<sub>50</sub> value for rats is >170 mg/m<sup>3</sup> for 6 hours. No gross signs of toxicity were observed. At necropsy, “slight” inflammation of the epithelium lining of the anterior portion of the nose and “slight ulceration” of the oronasal duct were reported in 5/5 males and 4/5 females exposed to 170 mg/m<sup>3</sup> and sacrificed on day 2. No exposure-related effects were observed in animals necropsied on day 14.

Reference: Nitschke, K.D., Quast, J.F. and Wolfe, E.L. 1985. Bisphenol A: Acute Aerosol Toxicity Study with Fischer 344 Rats. Dow Chemical Company Unpublished Report. As cited in the EU Risk Assessment.

Reliability: (Klimisch Code 2) Valid with restrictions.

### 13.3 ACUTE DERMAL TOXICITY

#### 13.3.1

Type: LD<sub>0</sub> [ ] LD<sub>100</sub> [ ] LD<sub>50</sub> [X] LD<sub>L0</sub> [ ] Other [ ]

Species/Strain: Rats/Crl:CD®(SD)BR

Value: >2000 mg/kg

Method: Not reported

GLP: Yes [ ] No [X] ? [ ]

Test Substance: Spirobiindane (CAS RN 1568-80-5)

Remarks: Test article was administered once dermally at a dose of 2000 mg/kg to the clipped, intact skin of 5/sex Crl: CD®(SD)BR rats (224-247g) for a 24-hour period under semi-occlusive dressing. Mortality, clinical observations, dermal findings, body weight changes and day 14 gross necropsy findings were evaluated.

Results: There were no deaths, test article-related clinical findings, body weight changes or gross necropsy findings. Two and seven rats had very slight to slight erythema and desquamation, respectively. The dermal LD<sub>50</sub> and the No-Observable-Effect Level (NOEL) for systemic toxicity were found to be >2000 mg/kg.

Reference: Kern<sup>1</sup>, T., Hurley<sup>1</sup>, J., Wilmot<sup>1</sup>, A. and Dimond, S. 1997. Acute Toxicity Studies with Spirobiindane. General Electric Plastics, Pittsfield, MA, USA and <sup>1</sup>WIL Research Laboratories, Inc. Ashland, OH, USA.

Reliability: (Klimisch Code 1) Valid without restrictions.

### 14.0 GENETIC TOXICITY *IN VITRO* OR *IN VIVO*

#### 14.1 BACTERIAL *IN VITRO* TEST

##### 14.1.1

Type: Bacterial reverse mutation assay (Ames test)

System of testing: *Salmonella typhimurium* strains TA98, TA100, TA1535, TA1537

Concentrations: 0 to 333 µg/plate

Metabolic activation: With [ ]; Without [ ]; With and Without [X]; No data [ ]

Results: Negative

Cytotoxicity conc.: With and without metabolic activation: 333 µg/plate

Genotoxic effects: With metabolic activation: positive [ ]; ambiguous [ ]

	Negative [X] Without metabolic activation: positive [ ]; ambiguous [ ]; Negative [X]
Year:	1983
GLP:	Yes [ ] No [ ] ? [X]
Test substance:	Bisphenol A (CAS RN 80-05-7)
Remark:	The pre-incubation method with <i>Salmonella typhimurium</i> strains TA98, TA100, TA1535 and TA1537 was used at bisphenol A concentrations up to 333 µg/plate in the presence and absence of metabolic activation (Aroclor induced rat and hamster liver S9). At 333 µg/plate, slight and complete clearance of the background lawn was observed in all strains with (rat and hamster liver S9) and without metabolic activation, respectively. No increase in the number of revertants was seen in any of the tested cultures. Controls gave results that confirmed the validity of this test. The negative result was confirmed by independent experiment.
Conclusion:	The test substance did not exhibit mutagenic activity in this assay.
Reference:	Haworth, S., Lawlor, T., Mortelmans, K., Speck, W. and Zeiger, E. 1983. <i>Salmonella</i> mutagenicity test results for 250 chemicals. <i>Environ. Mutagen.</i> <b>1</b> :3-142. As cited in the EU Risk Assessment.
Reliability:	(Klimisch Code 2) Valid with restrictions.
14.1.2	
Type:	Bacterial reverse mutation assay (Ames test)
System of testing:	<i>S. typhimurium</i> TA 1538 and <i>E. coli</i> strains WP2 and WP2uvrA
Concentrations:	0 to 1.0 mg/mL (concentrations in µg/plate not reported)
Metabolic activation:	With [ ]; Without [ ]; With and Without [X]; No data [ ]
Results:	Negative
Cytotoxicity conc.:	With and without metabolic activation: 1.0 mg/ml
Genotoxic effects:	With metabolic activation: positive [ ]; ambiguous [ ]; Negative [X] Without metabolic activation: positive [ ]; ambiguous [ ]; Negative [X]
Year:	1983
GLP:	Yes [ ] No [ ] ? [X]
Test substance:	Bisphenol A (CAS RN 80-05-7)
Remark:	<i>S. typhimurium</i> TA 1538 and <i>E. coli</i> strains WP2 and WP2uvrA were exposed in a pre-incubation assay to bisphenol A at concentrations up to 1.0 mg/mL in the presence and absence of metabolic activation. Cytotoxicity, as evidenced by a >50% decrease in the number of spontaneous revertants, was observed at 0.5 mg/ml in WP2uvrA with and without metabolic activation and TA 1538 without metabolic activation. No cytotoxicity was observed in WP2 with and without metabolic activation. No increase in the number of revertants was seen in any of the tested cultures. An independent experiment using a plate incorporation protocol and the performance of the controls confirmed the validity of this test.
Conclusion:	The test substance did not exhibit mutagenic activity in this assay.

Reference:	Dean, B.J. and Brooks, T.M. 1978. Toxicity Tests with Diphenylol Propane (DPP): <i>In vitro</i> Mutation Studies. Shell Research Report TLGR.0111.78. As cited in the EU Risk Assessment.
Reliability:	(Klimisch Code 2) Valid with restrictions.
14.1.3	
Type:	Bacterial reverse mutation assay (Ames test)
System of testing:	<i>S. typhimurium</i> TA98, TA100, TA1535 and TA1537 and <i>E. coli</i> strain WP2 <i>uvrA</i>
Concentrations:	0 - 5000 µg/plate
Metabolic activation:	With [ ]; Without [ ]; With and Without [X]; No data [ ]
Results:	Negative
Cytotoxicity conc.:	≥150 µg/plate
Genotoxic effects:	With metabolic activation: positive [ ]; ambiguous [ ]; Negative [X] Without metabolic activation: positive [ ]; ambiguous [ ]; Negative [X]
Year:	1997
GLP:	Yes [ ] No [ ] ? [X]
Test substance:	Spirobiindane (CAS RN 1568-80-5)
Remark:	<i>S. typhimurium</i> tester strains TA98, TA100, TA1535 and TA1537 and <i>E. coli</i> tester strain WP2 <i>uvrA</i> were exposed using the plate incorporation method to Spirobiindane at concentrations up to 5000 µg/plate in the presence and absence of metabolic activation (Aroclor-induced rat liver S9). The test substance was dissolved in DMSO at a concentration of 500 mg/ml.
Results:	In the first phase, the preliminary toxicity assay, the maximum dose tested was 5000 µg/ plate; this dose was achieved using a concentration of 100 mg/ml and a 50 µl plating aliquot. Precipitate was observed at ≥333 µg/plate. Toxicity was observed at ≥267 to ≥333, µg/plate, except for tester strain TA98 in the presence of S9 activation, in which toxicity was not observed. Based on the findings of the toxicity assay, the doses plated in the initial mutagenicity assay were 3.3-450 µg/plate without metabolic activation and 3.3-750 µg/plate with metabolic activation. In the mutagenicity assay, no positive response was observed. Precipitate was observed at ≥150 to 450 µg/plate and toxicity was observed at ≥150 to 750 µg/plate. This result was confirmed in an independent repeat assay.
Conclusion:	The test substance did not exhibit mutagenic activity in this assay.
Reference:	Wagner, V. 1997. Unpublished Report No. G97AH81.5020 entitled "Bacterial Reverse Mutation Assay with an Independent Repeat Assay", dated April 3, 1997 for General Electric Company, Plastics Business Operations, Pittsfield, MA, USA; from BioReliance, Inc., Rockville, MD, USA
Reliability:	(Klimisch Code 1) Valid without restriction



## 14.2 NON-BACTERIAL *IN VITRO* TEST (MAMMALIAN CELLS)

Type:	<i>In vitro</i> mammalian cell gene mutation test (Chromosomal aberration assay)		
System of testing:	CHO cells		
Concentration:	30 - 50 µg/mL with activation 20 - 40 µg/mL without activation		
Metabolic activation:	With [ ]; Without [ ]; With and Without [X]; No data [ ]		
Results:	Negative with and without metabolic activation.		
Cytotoxicity conc.:	With metabolic activation:	50 µg/mL	
	Without metabolic activation:	None reported	
Genotoxic effects:		+	?
	With metabolic activation:	[ ]	[X]
	Without metabolic activation:	[ ]	[X]
GLP:	Yes [ ] No [ ] ? [X]		
Test Substance:	Bisphenol A		
Remarks:	In a chromosome aberration study in Chinese hamster ovary (CHO) cells were exposed in two separate experiments to 30-50 µg/mL bisphenol A for 2 hours with metabolic activation and 20-40 µg/mL for 8 hours without metabolic activation. Cells were harvested at 11 hours with metabolic activation and 21 hours without metabolic activation. In the first test with metabolic activation, an increase in the percentage of metaphases with chromosome aberrations from bisphenol A treated cultures was observed only at the top dose in the presence of cytotoxicity; 14% at 50 µg/mL compared to 3% in controls. In these high-dose cultures, it was stated that cell confluence was reduced by approximately 70%. In the second test, no significant increases were observed in with metabolic activation; only 3% of cells at the highest dose had aberrations. No significant increases in aberrations were observed without metabolic activation with bisphenol A evidently being tested up to "toxic levels." The positive controls produced clear increases in chromosome aberrations.		
Conclusions:	This study was judged to have given a negative result, since the observed increase in the first experiment was not reproducible.		
Reference:	Ivett, J.L., Brown, B.M., Rodgers, C., Anderson, B.E., Ressnick, M.A. and Zeiger, E. 1989. Chromosomal aberrations and sister chromatid exchange tests in Chinese hamster ovary cells <i>in vitro</i> . IV. Results with 15 chemicals. <i>Environ. Mol. Mutagen.</i> <b>14</b> :165-187. As cited in the EU Risk Assessment.		
Reliability:	(Klimisch Code 2) Valid with restrictions.		

## 15.0 REPEATED DOSE TOXICITY

Species/Strain:	Rat/F344
Sex:	Female [ ]; Male [ ]; Male/Female [X]; No data [ ]
Route of Administration:	Oral, Dietary feed
Exposure Period:	103 weeks
Frequency of Treatment:	Daily

Post Exposure  
Observation Period: None  
Dose: 0, 1000 or 2000 ppm (approximately 50 and 100 mg/kg/day)  
Control Group: Yes ☒ No ☐ No data ☐  
Concurrent no treatment ☒ Concurrent vehicle ☐ Historical ☐  
NOAEL: None established  
LOAEL: 1000 ppm  
Method: Chronic study protocol from NTP  
Year: 1982  
GLP: Yes ☐ No ☒ ? ☐  
Test Substance: Bisphenol A  
Remark: The following was summarized for the IRIS Risk Assessment: In this 103-week dietary study, groups of 50 rats/sex were fed diets containing 0, 1000, or 2000 ppm bisphenol A. All treated groups of rats had reduced body weights, compared with controls, evident from the 5th week of exposure. Food consumption was also reduced, compared with controls, but this effect was not observed until the 12th week of treatment. Reduced body weights in rats, therefore, was considered a direct adverse effect of exposure to bisphenol A.  
Results: The LOAEL for BPA was considered to be 50 mg/kg/day. The Oral RfD was established at 0.05 mg/kg/day by IRIS.  
Reference: Integrated Risk Information System (IRIS Document); Bisphenol A. CASRN: 80-05-7. 1993.  
Reliability: (Klimisch Code 2) Valid with restrictions.

## 16.0 REPRODUCTIVE TOXICITY

Type: Fertility ☐ One generation study ☐ Two generation study ☐  
Other ☒

Species/Strain: Rat/CD<sup>®</sup> (Sprague-Dawley)

Sex: Female ☐ Male ☐ Male/Female ☒ No data ☐

Route of Administration: Dietary

Exposure Period: F0 males: 15 weeks; F0 females: 18 weeks  
F1 & F2 males: 18 weeks; F1 & F2 females: 21 weeks  
F3 males & females: 13 weeks

Premating Exposure Period: 10 weeks

Dose: 0, 0.15, 0.3, 4.5, 75, 750 or 7500 ppm (0, 0.001, 0.02, 0.3, 5, 50 or 500 mg/kg/day)

Control Group: Yes ☒; No ☐; No data ☐;  
Concurrent no treatment ☐ Concurrent vehicle ☒ Historical ☐

NOAEL Parental: 75 ppm (5 mg/kg/day)

NOAEL Reproduction: 750 ppm (50 mg/kg/day)

NOAEL F1 Offspring: 750 ppm (50 mg/kg/day)

Method: OECD Test Guideline 416 (modified)

GLP: Yes ☒ No ☐ ? ☐

Test Substance: Bisphenol A (99.5% pure)

Remark: Test procedure: Exposure to bisphenol A began for the F0 generation at about 7 weeks of age and continued throughout a 10-week pre-breed exposure period, a 2-week mating period (when F0 animals were mated [one male and one female] within each dose group) and gestation. F0 males were sacrificed after the F1 delivery period. Exposure of F0 females to bisphenol A continued throughout lactation until weaning (post-natal day 21) when F0 animals were sacrificed. Selected F1 animals were similarly mated to produce the F2 generation and selected F2 animals were mated to produce the F3 generation. The same exposure regime was used for F1 and F2 animals with direct exposure to bisphenol A in the diet commencing approximately at post-natal day 21. Selected F3 animals were exposed to bisphenol A only for a 10-week period from weaning as they were not mated. For the F0 generation and retained F1, F2 and F3 animals, clinical signs of toxicity, body weights and food consumption were reported. Estrous cycles were monitored in the last 3 weeks of the pre-breed exposure period and during the mating period for all generations. At the necropsy of adult animals, sperm samples were taken for analysis of epididymal sperm number, motility (using a computer assisted sperm motion analysis system) and morphology, testicular-resistant spermatid head counts, daily sperm production, and efficiency of daily sperm production, a number of organs were weighed and selected organs were examined histopathologically. Parameters assessed in the young offspring included litter size, body weights, survival, gross appearance, anogenital distance (AGD) (in F2 and F3 offspring only), sexual development and, for animals killed at weaning, gross appearance of organs at necropsy with attention given to reproductive organs.

## Results

**7500 ppm (approximately 500 mg/kg/day).** Adult systemic toxicity (F0, F1, F2, and F3) was expressed as consistent and persistent reduced body weights and weight gains, typically greater than 15%, occasional reductions in feed consumption as g/day, and occasional increases in feed consumption as g/kg/day. Food efficiency was occasionally reduced. For non-reproductive organs, absolute organ weights were consistently reduced in both sexes, while relative organ weights were commonly increased. Histopathologic findings on non-reproductive organs included increased incidence of renal tubular degeneration and chronic hepatic inflammation only in F0, F1, and F2 females (but not F3). There were no treatment- or dose-related direct effects on reproductive organ weights, histopathology, or function. Consistent with the reduced body weights and body weight gains in this group, paired ovarian weights were significantly reduced for F0, F1, and F2 females (absolute and relative weights) and for F3 females (absolute weights only). Paired ovarian follicle counts (ten females each at 0 and 7500 ppm) were significantly increased at 7500 ppm in F0 females but were unaffected in F1, F2, and F3 females. There were no effects of treatment on any andrological parameters at any dose in any generation (except for significantly reduced epididymal sperm counts only in F1 males at 7500 ppm, and for significantly reduced daily sperm production per testis, but not efficiency of daily sperm production only in F3 males at 7500 ppm). The only reproductive effect was a significant decrease in total and live pups per litter (with no significant increase in prenatal postimplantation loss per litter) on pnd 0 and 4 (precull) for F1, F2, and F3 litters at 7500 ppm, associated with reduced uterine weight in parental F0, F1, and F2 females. There were no effects on mating, fertility, or gestational indices, on precoital interval, gestational length, offspring sex ratios, or on nipple and/or areola retention in preweanling F1, F2, and F3 male pups. The only offspring effects at 7500 ppm included statistically significant reductions in pup body weights per litter, beginning on pnd 7 for F1, F2, and F3 offspring, and statistically significant delayed acquisition of both vaginal patency and preputial separation for F1, F2, and F3 postweanling pups. Anogenital distance (measured in F2 and F3 offspring on pnd 0) was unaffected for F2 and F3 males and for F3 females. Only F2 female anogenital distance was significantly increased by 0.03 mm at 0.015, 0.3, and 4.5 ppm, and by 0.04 mm at 750 ppm versus the control value. Although statistically significant, these variations were only 0.03 to 0.04 mm and were considered to be due to the extremely precise measurement technique used and were not considered to be of biological or toxicological significance. These changes (referring to preputial separation, vaginal patency, and anogenital distance) did not affect reproductive function in the animals exhibiting them. The effects observed on reproductive function or postnatal toxicity were present in a clearly toxic dose (adult systemic toxicity) and were not observed at doses below 7500 ppm.

**750 ppm (approximately 50 mg/kg/day).** Adult systemic toxicity was present for F0, F1, F2, and F3 generations, expressed only as reduced

body weights for F0 females (but not males) during prebreed, gestation, and lactation; F1 females (but not males) during gestation and lactation (but not prebreed); F2 males (but not females) only during prebreed, but not for F3 males or females. Feed consumption in g/day or g/kg/day and food efficiency were not consistently or persistently affected. In males, absolute liver weight was significantly reduced in all generations, absolute pituitary weight was significantly reduced in the F1 generation, and absolute kidney and spleen weights were significantly reduced in the F2 generation. In females, absolute and relative non-reproductive organ weights were not affected in any generation. There were no treatment-related gross or histopathologic findings. Reproductive organ weights, histopathology, and reproductive function were all unaffected in both males and females. There were no effects on reproductive, gestational, or offspring parameters, on pre- or postnatal survival, anogenital distance (F2 and F3 offspring, except for a significant increase of 0.04 mm for F2 female anogenital distance), or onset of puberty (except for a small delay in acquisition of preputial separation in F1 males, but not in F2 or F3 males)

Conclusion:	<p><b>75 ppm - 0.015 ppm (approximately 5 - 0.001 mg/kg/day).</b> There were no consistent or persistent treatment-related effects in F0, F1, F2, or F3 animals, nor in F1, F2, or F3 offspring.</p> <p>Exposure to dietary BPA for three generations, one litter per generation, resulted in adult systemic toxicity only at 750 and 7500 ppm. Because of the lack of adverse effects below these doses, the adult systemic toxicity NOAEL (no observable adverse effect level) was established at 75 ppm (equivalent to approximately 5 mg/kg/day) in CD® (Sprague-Dawley) rats, under the conditions of this study. Similarly, because there were no treatment- or dose-related reproductive or postnatal toxicity effects below 7500 ppm, the reproductive and postnatal toxicity NOAELs were concluded to be 750 ppm (equivalent to approximately 50 mg/kg/day) in CD® (Sprague-Dawley) rats, under the conditions of this study. There were no effects of BPA dietary exposure on any adult or offspring parameter at 0.015, 0.3, 4.5, or 75 ppm (resulting in BPA intakes of approximately 0.001, 0.02, 0.3, or 5 mg/kg/day, respectively).</p>
Reference:	<p>R.W. Tyl, Ph.D., DABT, C.B. Myers, M.S., M.C. Marr, B.A., LATG. 2000. Three-Generation Reproductive Toxicity Study of Dietary Bisphenol A in CD Sprague-Dawley Rats. RTI Report #: 65C-07036-000.</p>
Reliability:	<p>(Klimisch Code 1) Valid without restrictions.</p>

## 17.0 DEVELOPMENTAL TOXICITY/TERATOGENICITY

Species/Strain: Rat; Sprague-Dawley CD  
Sex: Female [X]; Male [ ]; Male/Female [ ]; No data [ ]  
Route of Administration: Oral (gavage)  
Duration of Test: 20 days  
Exposure Period: Days 6 through 15 of gestation  
Frequency of Treatment: Daily  
Dose: 0, 160, 320, 640 or 1280 mg/kg/day  
Control group: Yes [X] No [ ] No data [ ]  
Concurrent no treatment [ ] Concurrent vehicle [X] Historical [ ]

NOEL Maternal  
Toxicity: None Established; LOAEL = 160 mg/kg/day.  
NOEL Teratogenicity: > 640 mg/kg/day  
GLP: Yes [X] No [ ] ? [ ]  
Test Substance: Bisphenol A  
Method: Not stated.  
Remark: Range-finding study: groups of 8 time-mated CD rats were gavaged with 0, 500, 1000, 1500, 2000, 2500 or 3000 mg BPA/kg/day in corn oil on days 6 to 15 of gestation. Animals were sacrificed on day 20 of gestation and the fetuses were examined for gross morphological abnormalities only. Deaths were observed in 1, 5, 3 and 5 rats at 1,500, 2,000, 2,500 and 3,000 mg/kg, respectively. Clinical signs of toxicity were observed at all doses and included diarrhea, lacrimation, lethargy, rough coat, wheezing, wet urogenital area, arched back, dyspnea, bloody nose, disorientation, alopecia, piloerection and vaginal bleeding. A statistically significant decrease in maternal body weight gain (24%) was observed at 1000 mg/kg and above during the gestation period, and during the treatment period a decrease in maternal body weight was observed at 1500 mg/kg and above.

In the main study, two tests were performed and the data from both tests combined. Thus in total, groups of 27-29 time-mated CD rats were gavaged with 0, 160, 320, 640 or 1280 mg BPA/kg/day in corn oil on days 6 to 15 of gestation. Animals were sacrificed on day 20 of gestation and the fetuses were subjected to routine external, visceral and skeletal examination.

Results: No deaths were observed in the 0, 160, 320 and 640 mg/kg dose groups. At 1280 mg/kg, deaths were observed in 7/27 females and, because of this high mortality rate, the top dose group was not included in statistical analysis. Outward signs of toxicity observed in BPA treated animals included rough coat, piloerection, alopecia, pica, excess salivation, wet urogenital area, dyspnea, chromodacryorrhea, tremors, red-stained coat, wheezing and vocalization. A statistically significant decrease in mean maternal body weight gain was observed in dams at all dose levels for the treatment period (35-54%) and the gestation period (11-14%). No effect was observed on gravid uterine weights. When maternal body weight gain was corrected for gravid uterine weight, a statistically significant decrease was still apparent at all dose levels (26-34%). At necropsy, no

Conclusion:	effect was observed on mean relative liver weight in dams. Pregnancy rates were not affected by treatment with BPA, nor was there any effect on the number of implantation sites per litter, % resorptions per litter, number of live fetuses per litter, sex ratio, mean fetal body weight per litter, % fetuses malformed per litter and % litters with malformed fetuses. No evidence of developmental toxicity in the rat was evident at exposure levels which are toxic to the mother. A maternal no effect level could not be identified; the LOAEL of 160 mg/kg based on clinical signs of toxicity and a statistically significant decrease (26%) in body weight gain was identified. No fetal effects were seen at the highest dose level evaluated, 640 mg/kg.
Reference:	Morrissey, R.E., George, J.D., Price, C.J., Tyl, R.W., Marr, M.C. and Kimmel, C.A. 1987. The developmental toxicity of bisphenol A in rats and mice. <i>Fundam. Appl. Toxicol.</i> <b>8</b> :571-582. As cited in the EU Risk Assessment.
Reliability:	(Klimisch Code 1) Reliable without restrictions.