

Air Toxics Hot Spots Program

1-Bromopropane Cancer Inhalation Unit Risk Factor

Technical Support Document for
Cancer Potency Factors
Appendix B

Scientific Review Panel Review
Draft

September 2021



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1-Bromopropane

Cancer Inhalation Unit Risk Factor

Technical Support Document for Cancer Potency Factors Appendix B

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List of Acronyms

AIC	Akaike Information Criterion	GSH	Glutathione
ANOVA	Analysis of Variance	GST	Glutathione-S-transferase
1-BP	1-Bromopropane	IL	Interleukin
BMD	Benchmark dose	iNOS	Nitric Oxide Synthetase
BMD05	BMD 5% response rate	IUR	Inhalation unit risk
BMDL05	The 95% lower confidence bound at the 5% response rate	IR	Inhalation rate
BMDS	Benchmark dose modeling software	IARC	International Agency for Research on Cancer
BMR	Benchmark dose response	IV	Intravenous
BR	Breathing rate	NO	Nitric Oxide
BW	Body weight	NTP	National Toxicology Program
CEBS	Chemical Effects Biological Systems	OEHHA	Office of Environmental Health Hazard Assessment
CF	Conversion factor	PBPK	Physiologically-based pharmacokinetic
CO2	Carbon Dioxide	ppm	parts per million
CSF	Cancer slope factor	PrCys	S-propylcysteine
CTI	California Toxics Inventory	TNF	Tumor necrosis factor
CYP	Cytochrome P450	TRI	Toxics Release Inventory
CYP2E1	Cytochrome P450 2E1 isozyme	TWA	Time-weighted average
DBCP	1,2-dibromo-3-chloropropane	US EPA	United States Environmental Protection Agency
DNA	Deoxyribonucleic acid	VOC	Volatile organic compound
FCM	Flavin-containing monooxygenase	WT	Wild-type

1 **Preface**

2 This document summarizes the carcinogenicity data and the derivation of an
3 inhalation cancer unit risk factor for 1-bromopropane (1-BP). Cancer unit risk factors
4 are used to estimate lifetime cancer risks associated with inhalation exposure to a
5 carcinogen. The National Toxicology Program (NTP) conducted chronic inhalation
6 toxicity and carcinogenicity bioassays of 1-BP (Morgan *et al.*, 2011; NTP, 2011) and
7 found evidence of carcinogenicity in rats and mice. Consequently, OEHHA has
8 derived a cancer inhalation unit risk factor (IUR) from the NTP animal data for use in
9 the Hot Spots program.

10 OEHHA is legislatively mandated to develop guidelines for conducting health risk
11 assessments under the Air Toxics Hot Spots Program (Health and Safety Code
12 Section 44360(b)(2)). In implementing this requirement, OEHHA develops IURs for
13 carcinogenic air pollutants listed under the Air Toxics Hot Spots program. The 1-BP
14 IUR was developed using the most recent “Air Toxics Hot Spots Program Technical
15 Support Document for Cancer Potency Factors”, finalized by OEHHA in 2009
16 (OEHHA, 2009). Literature summarized and referenced in this document covers the
17 relevant published reports for 1-BP through spring 2021

18 1-BP has been proposed to be added to the list of substances for which emissions
19 must be quantified under the OEHHA Air Toxics Hot Spots Program in 2021, and is a
20 reportable chemical under the US EPA Toxics Reporting Inventory (TRI) program
21 (TRI, 2015). 1-BP is listed as a chemical known to the State to cause cancer by the
22 California Proposition 65 program (OEHHA, 2016). In addition, the National
23 Toxicology Program (NTP) listed 1-BP in the 13th Report on Carcinogens, which
24 identifies substances that either are known to be human carcinogens or are
25 reasonably anticipated to be human carcinogens, and to which a significant number
26 of persons residing in the United States are exposed (NTP, 2013). 1-BP is also listed
27 by the International Agency for Research on Cancer (IARC) as a Group 2B
28 carcinogen, i.e., possibly carcinogenic to humans (IARC, 2018).

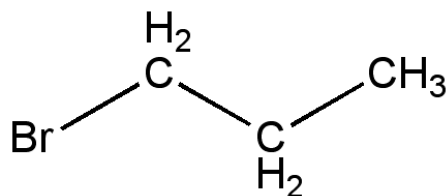
29 1-BP is promoted as an alternative to ozone-depleting chlorofluorocarbons.
30 Exposure to 1-BP may occur from emissions of facilities where 1-BP is used as a
31 solvent vehicle for spray and brush-applied adhesives in laminates and foam
32 products, or as a degreasing/cleaning agent for metals, metal products, plastics,
33 optics, and electronics (TRI, 2015). 1-BP is also listed in California for limited use in
34 dry cleaning technologies, in which it is used as an alternative solvent in modified
35 perchloroethylene dry-cleaning machines (CARB, 2015). Other applications may
36 include use as a chemical intermediate in the production of pharmaceuticals,
37 pesticides, quaternary ammonium compounds, flavors, and fragrances. In California,

38 reduction in chlorinated hydrocarbon use due to phase-out of these compounds has
39 led to the adoption of alternative solvent formulations, such as those including 1-BP,
40 by end-users. A periodic California survey of businesses that conduct solvent
41 cleaning operations noted no use of 1-BP until 2008 (CARB, 2011). In that year, the
42 Statewide Emission Inventory reported a total of 160.7 tons total organic gases/year
43 of 1-BP emissions due to solvent cleaning operations.

44

45 **1-BROMOPROPANE**

46 CAS No: 106-94-5



47

48 **I. PHYSICAL AND CHEMICAL PROPERTIES**

49 (PubChem, 2020)

50	Molecular formula	C ₃ H ₇ Br
51	Molecular weight	122.99 g/mol
52	Synonym	n-Propyl bromide
53	Description	Colorless liquid when fresh
54	Density/Specific gravity	1.353 @ 20°C/20°C
55	Boiling point	71°C at 760 mm Hg (torr)
56	Vapor pressure	110.8 mm Hg (torr) @ 20°C
57	Solubility	Soluble in acetone, ethanol, ether, benzene, chloroform, carbon tetrachloride; slightly soluble in water (2,450 mg/L @ 20°C)
58		
59		
60	Conversion factor	1 ppm = 5.03 mg/m ³

61 **II. HEALTH ASSESSMENT VALUES**

62	Unit Risk Factor:	$3.7 \times 10^{-6} (\mu\text{g}/\text{m}^3)^{-1}$
63	Inhalation Slope Factor:	$1.3 \times 10^{-2} (\text{mg}/\text{kg}\text{-day})^{-1}$
64		

65 **III. CARCINOGENICITY**

66 Carcinogenicity studies for 1-BP have been conducted in rats and mice. There are
67 no human carcinogenicity data.

68 **NTP Cancer Bioassay**

69 The NTP conducted two-year 1-BP inhalation studies in male and female F344/N rats
70 and B6C3F₁/N mice (Morgan *et al.*, 2011; NTP, 2011). 1-BP was chosen for study by
71 NTP due to the potential for increasing widespread use and the lack of
72 carcinogenicity data. Rodents were exposed whole-body in chambers to 0, 62.5
73 (mice only), 125, 250, or 500 (rats only) ppm (314, 629, 1,258 and 2,515 mg/m³)
74 1-BP for 6.17 hours/day, 5 days/week for 105 weeks. The daily exposures included
75 the 6 hour exposure time at a uniform aerosol concentration plus the ramp-up time of

76 10 minutes (0.17 hours/day) to achieve 90% of the target concentration after the
77 beginning of aerosol generation. The decay time to 10% of the target concentration
78 at the end of the exposures was about 10-11 minutes.

79 **F344/N rats**

80 Body weights of male and female rats at all exposure levels were similar to controls,
81 remaining within 8% of controls throughout the 2-year study. Survival was
82 significantly reduced in the 500 ppm males compared to controls ($p = 0.033$, life table
83 pairwise comparison) (NTP, 2011). In this exposure group, 9 of 37 deaths prior to
84 terminal sacrifice were attributed to chronic inflammation in various organs (lung,
85 nose, skin, and bone) that were related to 1-BP exposure, while the remaining early
86 deaths were attributed to various types of neoplasia that were not treatment-related.
87 In females, decreased survival in the 500 ppm group was not significant ($p = 0.054$)
88 compared to the controls. However, the life table trend test indicated decreased
89 survival of the female rats with increasing dose ($p = 0.028$).

90 The statistically significant ($p < 0.05$) or biologically noteworthy tumor incidences in
91 male and female rats are shown in Table 1. The incidence of adenoma of the large
92 intestine (colon or rectum) was significantly increased in 500 ppm female rats and a
93 significant positive trend ($p = 0.004$) for this tumor was observed. In 1-BP treated
94 males, the low incidence of these tumors resulted in no significant difference relative
95 to controls, and no significant positive trend was found. This tumor is rare in F344/N
96 rats. The historical incidence in 2-year inhalation studies with male rat chamber
97 controls is: 0/349; all routes 2/1,398 (0.1% \pm 0.5%), range 0-2%. The incidence of
98 adenoma of the large intestine was exceeded in 250 ppm males (2/50, 4%). The
99 NTP (2011) concluded that the presence of these tumors in exposed females and the
100 low historical incidence in controls indicated the tumors in males were exposure
101 related. Although no carcinomas of the large intestine were found in the 1-BP-
102 exposed rats, adenoma of the large intestine can progress to carcinoma (NTP,
103 2011a).

104 Skin tumors of epithelial origin were increased in exposed male rats (Table 1). The
105 tumor incidence of keratoacanthoma was significantly increased in the 250 and 500
106 ppm groups compared to controls, and a significant positive trend was observed.
107 The tumor incidence of keratoacanthoma or squamous cell carcinoma combined was
108 significantly increased in 500 ppm males and a significant positive trend was
109 observed. Keratoacanthoma is a rapidly growing benign neoplasm of squamous
110 epithelial origin that is considered to progress to squamous cell carcinoma. The
111 historical control range for keratoacanthoma and keratoacanthoma or squamous cell
112 carcinoma (combined) was exceeded in 250 and 500 ppm males.

113 When combining all neoplasms of epithelial origin, the tumor incidence of
114 keratoacanthoma, basal cell adenoma, basal cell carcinoma, or squamous cell
115 carcinoma in males was significantly increased in all exposed groups, and a positive
116 trend was observed (Table 1). The incidence for all epithelial tumors combined in all
117 exposed groups exceeded the historical control range for inhalation studies. The
118 NTP (2011) concluded that the increased incidences of all tumors of epithelial origin
119 were a result of 1-BP exposure.

120 Tumors of the skin were not as prevalent in exposed female rats. A positive trend for
121 tumor incidence was found when squamous cell papilloma, keratoacanthoma, basal
122 cell adenoma, or basal cell carcinoma were combined, but pairwise comparison of
123 1-BP exposed groups with controls did not result in a significant increase in tumors in
124 any group (Table 1). The tumor incidence in the 500 ppm group did exceed the
125 historical incidence for controls for inhalation studies (2/350 – 0.6% ± 1.0%, range 0-
126 2%) and for all routes of exposure (16/1350 – 1.2% ± 1.8%, range 0-6%). The NTP
127 (2011) concluded there was equivocal evidence for these skin tumors in exposed
128 female rats due to the absence of statistically significant pairwise comparisons for
129 keratoacanthoma alone and for all tumors combined, and because there were no
130 observed squamous cell carcinomas.

131 A positive trend for the incidence of malignant mesothelioma was observed in male
132 rats, and the incidence in the 500 ppm group was near statistical significance
133 ($p = 0.059$). This neoplasm originated in the epididymis but was also found in other
134 tissues, particularly the testis. The NTP (2011) noted that the historical control
135 incidence was surpassed in the 500 ppm group (inhalation studies: 5/349 – 1.4% ±
136 2.2%, range 0-6%; all routes: 35/1,398 – 2.5% ± 2.3%, range 0-6%). The NTP
137 concluded there was only equivocal evidence for carcinogenicity for this tumor due to
138 its common occurrence in this strain of male rats, lack of a statistically significant
139 increase in exposed groups relative to controls, and because the 500 ppm group
140 tumor incidence was barely above the historical control range.

141 In male rats, a significant increase in the tumor incidence for pancreatic islet cell
142 adenoma occurred in most 1-BP-exposed groups, and a positive trend near statistical
143 significance ($p = 0.056$) was observed (Table 1). However, the historical control
144 range for this neoplasm in inhalation studies (0% to 12%) was not exceeded in any of
145 the exposed groups and the mean incidence in historical control inhalation studies
146 (5.7% ± 3.9%) was greater than that in chamber controls (0%). Thus, the NTP
147 considered the increased incidence of this tumor as equivocal evidence for
148 carcinogenicity. No significant difference from control was observed for the incidence
149 of pancreatic islet cell carcinoma, and no positive trend was observed. The incidence
150 of carcinomas in the 125 ppm group (7/50) was above the historical control range

151 (inhalation studies: 17/349 – 4.9% ± 3.3%, range 2-10%; all routes: 29/1,394 – 2.1%
152 ± 2.6%, range 0-10%). The NTP (2011) concluded that pancreatic islet cell
153 carcinoma demonstrated equivocal evidence of carcinogenicity due to the lack of a
154 significant increase over control incidence.

155 For pancreatic islet cell adenoma or carcinoma (combined), there was a significantly
156 increased tumor incidence in the 125 ppm group, but a significant positive trend was
157 not demonstrated. The historical control range in the 125 ppm group was exceeded
158 (inhalation studies: 37/349 – 10.6% ± 4.8%, range 6-18%; all routes: 119/1,394 –
159 8.6% ± 4.0%, range 0-18%). Although not specifically addressed by the NTP, the
160 combined tumor incidence data was apparently not strong enough to affect the
161 conclusion of equivocal evidence for carcinogenicity based on the individual
162 adenoma and carcinoma incidence rates.

163

164 **Table 1. Unadjusted tumor incidence in rats exposed to 1-BP for two years (NTP, 2011a)^{a,b}**

Sex and Species	Tumor Type	Incidence by concentration				Statistical p-values for pairwise comparison with controls			
		0 ppm, 0 mg/m ³	125 ppm, 629 mg/m ³	250 ppm, 1258 mg/m ³	500 ppm, 2515 mg/m ³	Trend ^c	125 ppm, 629 mg/m ³	250 ppm, 1258 mg/m ³	500 ppm, 2515 mg/m ³
Male Rats	Large Intestine (colon or rectum): Adenoma	0/50	0/50	2/50	1/50	0.140	1.000	0.247	0.500
	Skin: Basal Cell Adenoma	0/50	1/50	2/50	1/50	0.247	0.500	0.247	0.500
	Skin: Basal Cell Carcinoma	0/50	2/50	1/50	2/50	0.160	0.247	0.500	0.247
	Skin: Keratoacanthoma	0/50	3/50	6/50*	6/50*	0.010	0.309	0.013	0.013
	Skin: Squamous Cell Carcinoma	1/50	1/50	0/50	2/50	0.247	0.753	1.000	0.500
	Skin: Keratoacanthoma or Squamous Cell Carcinoma	1/50	4/50	6/50	8/50*	0.008	0.181	0.056	0.015
	Skin: Keratoacanthoma, Basal Cell Adenoma, Basal Cell Carcinoma, or Squamous Cell Carcinoma	1/50	7/50*	9/50**	10/50**	0.007	0.030	0.008	0.004
	Malignant Mesothelioma [†]	0/50	2/50	2/50	4/50	0.026	0.247	0.247	0.059
	Pancreatic Islets: Adenoma	0/50	5/50*	4/50	5/50*	0.056	0.028	0.059	0.028
	Pancreatic Islets: Carcinoma	3/50	7/50	5/50	3/50	0.662	0.159	0.357	0.661
	Pancreatic Islets: Adenoma or Carcinoma	3/50	10/50*	9/50	8/50	0.158	0.036	0.061	0.100
Female Rats	Large Intestine (colon or rectum): Adenoma	0/50	1/50	2/50	5/50*	0.004	0.500	0.247	0.028
	Skin: Squamous Cell Papilloma, Keratoacanthoma, Basal Cell Adenoma, or Basal Cell Carcinoma	1/50	1/50	1/50	4/50	0.040	0.753	0.753	0.181

165 (a) The numerator represents the number of tumor-bearing animals; the denominator represents animals examined.

166 (b) * = $p < 0.05$, ** = $p < 0.01$; p-value indicators are from pairwise comparisons with controls using Fisher exact tests performed by OEHHA

167 (c) p-values in the trend column are for the Cochran-Armitage trend test performed by OEHHA.

168 † = Tumor type and incidence data represents equivocal finding for carcinogenicity by NTP (2011a)

169 Nonneoplastic findings included increased incidence of various upper respiratory
170 lesions in the nose, larynx and trachea in some or all exposed groups of rats,
171 including chronic active inflammation, suppurative chronic inflammation, epithelial
172 hyperplasia in the nose and trachea, and respiratory metaplasia of the nasal olfactory
173 epithelium. Chronic suppurative inflammation was significantly increased in the lung
174 of 500 ppm females. Chronic suppurative inflammation was also present in skin and
175 some other tissues of 500 ppm males and females. These lesions are characterized
176 by the presence of Splendore Hoeppli material, which were not seen in controls. The
177 presence of Splendore Hoeppli material has been associated with diseases that
178 compromise the immune system (NTP, 2011a).

179 **B6C3F1/N mice**

180 Body weights of male and female mice at all exposure levels were similar to controls
181 throughout the 2-year studies (NTP, 2011). Survival of the mice was unaffected by 1-
182 BP exposure.

183 There was no evidence of carcinogenic activity of 1-BP in male mice. However, an
184 increased incidence of lung tumors was observed in 1-BP-exposed female mice
185 (Table 2). Significantly increased tumor incidences of alveolar/bronchiolar adenomas
186 (250 ppm group), alveolar/bronchiolar carcinomas (62.5 and 125 ppm groups), and
187 combined alveolar/bronchiolar adenoma or carcinoma (all exposed groups) were
188 present, including positive trends for the adenoma and combined adenoma or
189 carcinoma. In addition, multiple adenomas were found in two 250 ppm females and
190 multiple carcinomas were found in two 62.5 ppm females, one 125 ppm female, and
191 one 250 ppm female. The inhalation study historical control range for
192 alveolar/bronchiolar adenoma and for the adenoma or carcinoma (combined) was
193 exceeded by the 250 ppm group and by all exposed groups, respectively.

194

195 **Table 2. Un-adjusted tumor incidence in mice exposed to 1-BP for two years (NTP, 2011a)^{a,b}**

Sex and Species	Tumor Type	Incidence by concentration				Statistical p-values for pairwise comparison with controls			
		0 ppm, 0 mg/m ³	62.5 ppm, 314 mg/m ³	125 ppm, 629 mg/m ³	250 ppm, 1258 mg/m ³	Trend ^c	62.5 ppm, 314 mg/m ³	125 ppm, 629 mg/m ³	250 ppm, 1258 mg/m ³
Female Mice	Lung: Alveolar/Bronchiolar Adenoma	1/50	6/50	4/50	10/50**	0.004	0.056	0.181	0.004
	Lung: Alveolar/Bronchiolar Carcinoma	0/50	7/50**	5/50*	4/50	0.189	0.006	0.028	0.059
	Lung: Alveolar/Bronchiolar Adenoma or Carcinoma	1/50	9/50**	8/50*	14/50**	<0.001	0.008	0.015	<0.001

196 (a) The numerator represents the number of tumor-bearing animals; the denominator represents animals examined

197 (b) * = $p < 0.05$, ** = $p < 0.01$; p -value indicators are from pairwise comparisons with controls using Fisher exact tests performed by OEHHA;

198 (c) p -values in the trend column are for the Cochran-Armitage trend test performed by OEHHA.

199

200 Increased incidences of nonneoplastic findings were observed in the upper and lower
201 respiratory airways in some or all exposed groups of mice. Bronchiolar regeneration
202 was observed in most exposed male and female mice. This lesion was almost
203 completely absent in control mice. Cytoplasmic vacuolization in the bronchiolar
204 epithelium of the lung, the respiratory epithelium of the nose, and the epithelium of
205 the larynx and trachea was increased in all exposed male groups. Cytoplasmic
206 vacuolization was also increased in upper and lower airways in all exposed female
207 groups, but at lower rates compared to males. In the nose of male and female mice,
208 there was also an increased incidence of hyperplasia of the respiratory epithelium
209 and metaplasia of the olfactory epithelium in some or all exposed groups.

210 **Toxicokinetics**

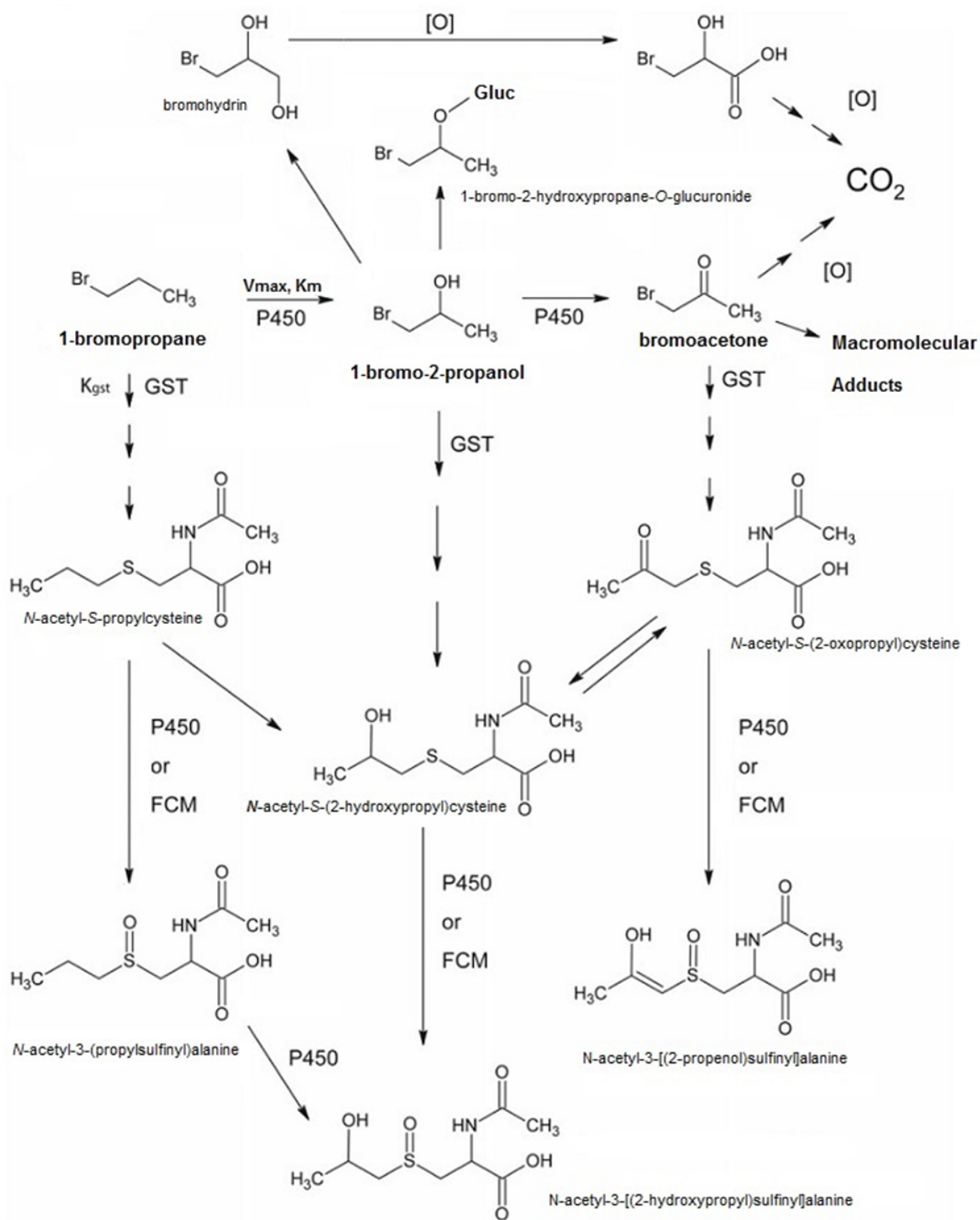
211 The mechanism by which 1-BP causes cancer has not been elucidated, although
212 metabolic activation to reactive metabolites is suspected to be involved (Morgan *et al.*,
213 2011). The metabolism of inhaled and absorbed 1-BP occurs primarily through
214 oxidative metabolism via P450 enzymes, conjugation with glutathione (GSH) and
215 debromination, although the majority of 1-BP can be excreted unchanged in exhaled
216 air. Metabolism of 1-BP has been shown to produce effects that other carcinogens
217 are known for, such as oxidative stress via glutathione depletion and
218 immunomodulation (Lee *et al.*, 2007; Guyton *et al.*, 2009; Liu *et al.*, 2009; Miao *et al.*,
219 2018).

220 **Rodent models**

221 Toxicokinetic studies have been carried out in male F344 rats and B6C3F₁ mice
222 (Garner *et al.*, 2006). The disposition of [1-¹⁴C]-1-BP radioactivity following relatively
223 low doses (3.4 - 5.9 mg/kg) via intravenous (IV) administration was similar in rats and
224 mice. A majority of the radiolabel was exhaled as volatile organic compounds (VOC;
225 40–71%) or as ¹⁴CO₂ (10–31%) within four hours following administration. The
226 radiolabel recovered in urine ranged from 17 to 23%. Roughly 2% and 6% was
227 recovered in feces and carcass, respectively. The radiolabel exhaled as VOC was
228 later identified in Garner *et al.* (2015) as the parent compound, 1-BP.

229 The identification of urinary metabolites was carried out following IV administration
230 and inhalation exposure of [1,2,3-¹³C]-labeled 1-BP in rats (Garner *et al.*, 2006).
231 Similar to the inhalation route, IV administration does not involve hepatic “first pass”
232 metabolism and is more likely to be consistent with metabolism derived from
233 workplace or environment inhalation. As expected, similar results were obtained for
234 both exposure routes. The main urinary metabolites and percent of the total excreted
235 in the urine were: *N*-acetyl-*S*-propylcysteine (37%), *N*-acetyl-3-(propylsulfinyl)alanine

236 (5%), *N*-acetyl-*S*-(2-hydroxypropyl)cysteine (16%), 1-bromo-2-hydroxypropane-*O*-
237 glucuronide (9%), *N*-acetyl-*S*-(2-oxopropyl)cysteine (12%), and *N*-acetyl-3-[(2-
238 oxopropyl)sulfinyl]alanine (% not stated). The authors indicated that many of these
239 metabolites were likely formed after cytochrome P450 (CYP)-catalyzed oxidation of
240 1-BP to 1-bromo-2-propanol and bromoacetone, followed by GSH conjugation with
241 either of those metabolites. Other identified 1-BP metabolites formed by CYP-
242 mediated oxidation in rodents include α -bromohydrin and glycidol, both of which have
243 been shown to be mutagenic (Stolzenberg and Hine, 1979; IARC, 2000; Ishidao *et*
244 *al.*, 2002; Garner *et al.*, 2007). The scheme established in Garner *et al.* (2015) for 1-
245 BP metabolism in the rat is shown in Figure 1.



246

247 **Figure 1. Metabolism of 1-BP in rats: Figure 2 of Garner et al. (2015).**

248 GST = glutathione-S-transferase; FCM = Flavin monooxygenase; Vmax = maximal velocity;

249 Km = Michaelis Constant; Kgst = proportionality constant for linear pathway metabolized by

250 glutathione transferase; →→ multiple steps of reaction

251 When rats were pretreated with 1-aminobenzotriazole, a potent but nonselective CYP
252 inhibitor/inactivator, the only urinary metabolite found was *N*-acetyl-*S*-propylcysteine,
253 which contributed greater than 90% of the urinary radioactivity (Garner *et al.*, 2006).
254 This metabolite is formed by direct conjugation of 1-BP with GSH. The authors
255 concluded that CYP enzymes contribute significantly to the production of the major
256 oxidative metabolites of 1-BP.

257 In a follow-up study, Garner *et al.* (2007) exposed *Cyp2e1*^{-/-} and wild-type (WT) mice
258 to [1,2,3-¹³C]-1-BP to determine the contribution of cytochrome P450 2E1 (CYP2E1)
259 to the metabolism and elimination of the chemical. In *Cyp2e1*^{-/-} mice, which lack the
260 CYP2E1 isozyme, the elimination half-life in gas uptake studies was longer compared
261 to WT mice (3.2 vs. 1.3 hours). The major urinary metabolite, *N*-acetyl-*S*-(2-
262 hydroxypropyl)cysteine, which is derived largely through oxidative metabolism, was
263 reduced about 50% in *Cyp2e1*^{-/-} mice compared to WT mice. In addition, the ratio of
264 products of direct conjugation of 1-BP with GSH to oxidative 2-hydroxylation
265 increased 5-fold in *Cyp2e1*^{-/-} mice relative to WT mice. These data suggested to the
266 authors that CYP2E1 is a major CYP contributor in the oxidative metabolism of 1-BP.

267 Garner and Yu (2014) evaluated the species and sex-dependent factors influencing
268 1-BP toxicokinetics in F-344 rats and B6C3F₁ mice after intravenous and inhalation
269 exposure. Male F-344 rats were given intravenous (iv) bolus injections of 1-BP at 5
270 or 20 mg/kg body weight (BW), and blood levels were determined at time intervals up
271 to 4 hours. Male and female F-344 rats and B6C3F₁ mice were also exposed to
272 initial inhalation concentrations of 70, 240, 800, and 2,700 ppm (0, 350, 1,200, 4,000,
273 and 14,000 mg/m³) 1-BP in a closed gas uptake system, and subsequent 1-BP
274 atmospheric loss rates monitored for 6 hours. Systemic clearance of bolus iv-
275 administered 1-BP in the blood of rats was rapid and decreased with increasing dose.
276 Approximately 99% was eliminated from the body by 3 hours post-exposure. The
277 average elimination half-life was 0.39 ± 0.08 and 0.85 ± 0.09 hour at 5 and 20 mg/kg
278 BW, respectively. However, systemic clearance decreased with increasing iv dose.
279 Plasma bromine levels were measured in the rats after iv administration; bromine is
280 released from 1-BP either by oxidative metabolism or by conjugation with GSH. The
281 bromine levels suggested that approximately 30% of administered 1-BP was
282 metabolized by either route and eliminated in urine. The authors surmised that the
283 remainder was largely lost by exhalation, either as 1-BP or as CO₂.

284 In the gas uptake portion of the study, as the air concentration of 1-BP increased, the
285 terminal air elimination rate decreased suggesting to the authors that one or more
286 routes of elimination became saturated as chamber concentration increased (Garner
287 and Yu, 2014). At a given starting concentration, male rats tended to eliminate 1-BP
288 from the chamber more rapidly than females. Plasma bromide levels were also

289 measured in the rats following gas uptake. The results showed that oxidative
290 metabolism in female rats was lower compared to males, indicating that oxidative
291 metabolism in females may be saturated at lower concentrations. In male and
292 female mice, elimination of inhaled 1-BP occurred at similar rates up to 800 ppm. At
293 higher concentrations, the half-life increased, with male mice eliminating 1-BP from
294 the chamber more slowly than female mice. The data also showed that mice tend to
295 have a higher 1-BP oxidative metabolic capacity relative to rats. Regarding urinary
296 metabolites, the authors noted that rats produced both directly GSH-conjugated
297 parent and oxidative metabolites, while mice only produced a single oxidative
298 metabolite (2-hydroxybromopropane) which was then conjugated with GSH.

299 Rats were also pretreated with chemical inhibitors of CYP (1-aminobenzotriazole)
300 and GSH (D,L-buthionine (S, R)-sulfoximine) synthesis, prior to exposure to 1-BP at
301 800 ppm (4,024 mg/m³) in inhalation chambers (Garner and Yu, 2014). The half-life
302 of 1-BP in rats following inhibition of CYP (9.6 hours) or depletion of GSH (4.1 hours)
303 increased relative to controls (2.0 hours), supporting the authors' position that 1-BP
304 elimination is highly dependent on both CYP and GSH-dependent metabolism.

305 Applying the above gas-uptake experiments in the Fischer 344 rat, a physiologically
306 based pharmacokinetic (PBPK) model was developed by simulating the 1-BP level in
307 a closed chamber (Garner *et al.*, 2015). They tested the hypothesis that metabolism
308 includes both P450 CYP2E1 activity and GSH conjugation. The results showed that
309 two metabolic pathways adequately simulated 1-BP levels in the closed chamber.
310 Furthermore, the model was tested by simulating the gas-uptake data of the female
311 rats pretreated with the P450 inhibitor 1-aminobenzotriazole, or the GSH synthesis
312 inhibitor d,l-buthionine (S,R)-sulfoximine, prior to inhalation of 800 ppm (4,000 mg/m³)
313 1-BP. As in their previous study, pretreatment with either of these inhibitors
314 dramatically prolonged the half-life of 1-BP elimination, and suggested CYP 450 and
315 GSH had major roles for 1-BP metabolism.

316 Based on the closed chamber and gas-uptake data in the female rat, sex-specific
317 metabolic parameters were also estimated and extrapolated into different exposure
318 levels in the PBPK model (Garner *et al.*, 2015). In the model, the metabolic rate
319 V_{max} and K_m were about 1.5 and 2 times larger in the male rat than those in the
320 female. The GSH-related constant (K_{gst}) in the male rat was estimated to be about 2
321 times that of the female constant. After adjusting V_{max} by the rat's body weight
322 (male rat body weight was considerably greater than the female rat body weight), the
323 values were improved and shown to be similar between male and female rats, which
324 indicates body weight as a possible contributor to the sex-specific differences in the
325 toxicokinetics of 1-BP.

326 A human PBPK model for 1-BP was developed by extrapolating the metabolic
327 parameters obtained from the gas-uptake studies in rats, and applying them to a
328 general human PBPK model for volatile compounds (Garner *et al.*, 2015). In a
329 repeated exposure scenario (20 or 200 ppm per day), modeling showed that rats do
330 not accumulate 1-BP in blood, whereas humans show a 20% increase over 5 days of
331 exposure. While 1-BP has a moderate fat:blood partition coefficient (20.2), a higher
332 fat tissue content in humans (21.4%) compared to rats (7%) may explain this
333 increase. However, additional experimental data for specific organ dosimetry and for
334 the metabolites of 1-BP will need to be incorporated into the PBPK model to allow the
335 quantitative extrapolation of animal studies to humans for risk assessment purposes.

336 **Toxicokinetics in children and adults**

337 The urinary mercapturic metabolite, N-acetyl-S-propylcysteine, found in rodents by
338 Garner and associates has also been identified in the urine of 1-BP-exposed workers
339 (Valentine *et al.*, 2007; Hanley *et al.*, 2009). In addition, a urinary metabolite not
340 identified in rodents, N-acetyl-S-(3-hydroxy-n-propyl)cysteine, has been found in
341 workers exposed to 1-BP (Cheever *et al.*, 2009; Hanley *et al.*, 2009). As in rodents,
342 N-acetyl-S-propylcysteine was identified as the predominant urinary metabolite in
343 exposed workers and was proposed as a biomarker of exposure. Although less
344 specific for 1-BP exposure, urinary bromide has also been proposed as a biomarker
345 of 1-BP exposure in workers (Hanley *et al.*, 2010).

346 In a peer-reviewed report, NIOSH investigators obtained 48-hour urine specimens
347 from 30 workers at two factories making polyurethane foam seat cushions and from
348 21 unexposed control subjects (Hanley *et al.*, 2009). The urine was collected into
349 composite samples for three time intervals: at work, after work but before bedtime,
350 and upon awakening. Time-weighted average (TWA) geometric mean breathing
351 zone concentrations of 1-BP were 92.4 ppm (460 mg/m³) for sprayers (n=13) and
352 10.5 ppm (53 mg/m³) for non-spraying jobs (n=17). Urinary N-acetyl-S-
353 propylcysteine in urine showed the same trend as TWA exposures to 1-BP (i.e.,
354 sprayers had higher levels). Associations of N-acetyl-S-propylcysteine
355 concentrations, adjusted for creatinine, with 1-BP TWA exposure were statistically
356 significant for both sprayers (p < 0.05) and non-sprayers (p < 0.01). The study
357 confirmed that urinary N-acetyl-S-propylcysteine is an important 1-BP metabolite and
358 an effective biomarker for highly exposed foam cushion workers.

359 The unmetabolized parent compound has also been identified in end-of-shift urine
360 samples from 1-BP-exposed production workers, and was significantly correlated to
361 the concentration of 1-BP in air (Kawai *et al.*, 2001; Ichihara *et al.*, 2004a).
362 Measurable levels of 1-BP in end-of-shift urine was found when the TWA exposure

363 was >2 ppm (Kawai *et al.*, 2001). Unmetabolized 1-BP has not been detected in the
 364 urine of rats and mice (Garner *et al.*, 2006). Due to potential evaporative loss of
 365 1-BP from urine, the samples need to be immediately placed in sealed head-space
 366 vials with analysis often conducted the next day (Ikeda, 1999; Kawai *et al.*, 2001).

367 CYP2E1 is known to be a major CYP isozyme that metabolizes 1-BP in rodents.
 368 Initial reports in humans did not detect CYP2E1 in fetal liver samples, but CYP2E1
 369 increased rapidly within hours of birth (Vieira *et al.*, 1996; Cresteil, 1998). A more
 370 recent report with 73 fetal samples and 165 postnatal samples found that CYP2E1 is
 371 detectable by immunological techniques at low levels in some (37%) fetuses
 372 beginning in the second trimester, and in the third trimester it is present in most
 373 (80%) fetuses at 10-20% of adult levels (Johnsrud *et al.*, 2003; Hines, 2007). In the
 374 neonatal period (0-29 days) the mean level was about 25% that of adults but the
 375 variability among samples was nearly 80-fold (Johnsrud *et al.*, 2003). From 1 month
 376 to 1 year, the mRNA for CYP2E1 accumulates and CYP2E1 protein increases toward
 377 adult levels (Table 3) (Vieira *et al.*, 1996; Hines, 2007) (Vieira *et al.*, 1996; Hines,
 378 2007). However, considerable interindividual variability is observed in the immediate
 379 postnatal (1–6 months) onset or increase in expression of CYP2E1 and other CYP
 380 enzymes (Johnsrud *et al.*, 2003; Hines, 2007).

381 **Table 3. Increase of CYP2E1 (mean ± SD) with age in human liver (Hines, 2007)**

Age	n	pmol CYP2E1/mg protein
1 st trimester fetus: 8-13.4 weeks	14	- (not detectable)
2 nd trimester fetus: 13.6-25 weeks	45	0.3 ± 0.6
3 rd trimester fetus: 27-40 weeks	14	5.8 ± 4.6
Neonate: 0-29 days	42	13.4 ± 16.0
Infant: 1.1-11.3 months	64	36.2 ± 20.3
Prepubertal: 1.1-10.0 years	41	43.1 ± 20.6
Adolescent: 11.0-17.7 years	20	~68 ^a
Adult	-	~50 ^a

382 ^a Median, in pmol CYP2E1/mg protein

383 OEHHA noted that low levels of CYP2E1 in infants may reduce metabolism of 1-BP
 384 in the infant, leading to possible increased elimination of unchanged 1-BP via
 385 exhalation. Presuming that the parent compound has little or no toxicity, this could
 386 decrease, rather than increase, the sensitivity of the very young to the toxicity of 1-
 387 BP. However, there is currently no evidence to support this concept.

388 In non-occupational settings, surveys of children and pregnant women have found
 389 the 1-BP metabolite, N-acetyl-S-propylcysteine in most urine samples examined.
 390 From 2009 to 2010 the National Children's Vanguard Study collected urine samples
 391 from 488 third trimester pregnant women at in-person study visits (Boyle *et al.*, 2016).

392 Urinary metabolites of 28 VOCs were quantified simultaneously using ultra-high
393 performance liquid chromatography coupled with electrospray ionization tandem
394 mass spectrometry (UPLC-ESI/MSMS). N-acetyl-S-propylcysteine was present in
395 99% of the urine samples. The levels reported were 2.61 ng/mL for the 50th
396 percentile, 9.44 ng/mL for the 75th percentile, and 4,260 ng/mL for the maximum
397 person. The authors did not identify sources of the metabolite, other than to note that
398 dry cleaning and metal cleaning solvents are known sources of 1-BP.

399 Data from the National Health and Nutrition Examination Survey (NHANES) for 2011-
400 2012 were used to evaluate variability in the levels of 20 urinary metabolites of VOCs
401 (including 1-BP) by age, gender, and race/ethnicity (Jain, 2015) Among 417 children
402 ages 6 through 11, the mean levels of N-acetyl-S-propylcysteine were 2.6 (2–3.3)
403 ng/mL in boys and 3.3 (2.5–4.3) ng/mL in girls (adjusted geometric means with 95%
404 confidence intervals). Jain also reported that concentrations of the urinary 1-BP
405 metabolite decreased with the increase in the number of rooms in the child's home
406 ($p = 0.03$). The number of rooms in a child's home is an indicator of socioeconomic
407 status. However, the reason for this correlation was not known. No correlation of the
408 1-BP metabolite was observed with age, poverty income ratio, body mass index, or
409 number of smokers in the house.

410 US EPA (2020a) acknowledged in their review that there may be low-level non-
411 occupational exposure of 1-BP in the general population that resulted in measurable
412 blood levels of N-acetyl-S-propylcysteine found by Boyle et al. (2016), Jain (2015)
413 and others. However, due to the lack of information on the specificity of this
414 biomarker for 1-BP exposure, its use as a biomarker for the general population is
415 uncertain.

416 **Genotoxicity**

417 The genotoxicity and mutagenicity database for 1-BP is not extensive, and the overall
418 results have been mixed. Genotoxicity studies are summarized below, followed by a
419 table summary (Table 4 at the end of the section).

420 **DNA strand-break tests**

421 The comet assay is a commonly used method to identify DNA lesions (e.g., breaks or
422 alkali-labile sites) following exposure of an isolated cell culture with a genotoxin. This
423 assay measures premutagenic lesions, which, in intact cells, can be removed by
424 DNA repair processes, if the repair occurs prior to DNA replication. Thus, positive
425 assay data for a given compound do not necessarily indicate that the compound will
426 induce mutations.

427 Toraason and coworkers used the comet assay to assess DNA damage in human
428 leukocytes exposed *in vitro* to 1-BP, and in peripheral leukocytes *in vivo* from 53
429 workers occupationally exposed at two facilities to 1-BP (Toraason *et al.*, 2006). In
430 the *in vitro* portion of the study, leukocytes were collected from a single non-1-BP-
431 exposed human volunteer and cultured with 1- or 2-BP (0, 0.01, 0.1 or 1 mM). The
432 cells were cultured without metabolic activation. Both 1- and 2-BP induced a
433 significant increase ($p < 0.05$, ANOVA) in comet tail moment at the highest
434 concentration (1 mM). However, 1-BP induced apoptosis at a lower concentration
435 (0.1 mM), which the authors suggested could mean cells with excessive DNA
436 damage may be eliminated and reduce the potential for mutation.

437 In the *in vivo* worker study, 1-BP was used at the facilities as a solvent for spray
438 adhesives in foam cushion fabrication (Toraason *et al.*, 2006). Breathing zone
439 samples, collected with personal air monitors, was assessed in sprayers and non-
440 sprayers for 1 to 3 days. The exposure concentrations ranged from 0.2 to 271 ppm
441 (1 to 1,363 mg/m³). The mean 1-BP time-weighted average (TWA) at Facility A for
442 sprayers (n=3 and 10 for men and women, respectively) and non-sprayers (n=15 and
443 14 for men and women, respectively) was 83 ± 85 ppm and 2 ± 2 ppm, respectively.
444 The mean 1-BP TWA at Facility B for sprayers (n=6, women only) and non-sprayers
445 (n=3 and 13 for men and women, respectively) was 21 ± 5 ppm and 5 ± 1 ppm,
446 respectively. The study lacked a control group with no 1-BP exposure. Internal
447 biomarkers of exposure (serum and urine Br levels) were highly correlated with 1-BP
448 environmental exposure levels.

449 At both facilities, comet tail moments of leukocytes from sprayers were greater than
450 comet tail moments of leukocytes from non-sprayers, but the difference did not reach
451 statistical significance at $p < 0.05$ (Toraason *et al.*, 2006). An increased dispersion
452 coefficient ($p < 0.05$) in sprayers from Facility A was observed at the end-of-week
453 relative to start-of-week. The dispersion coefficient is the tail moment variance
454 divided by the mean, and variance was determined from 100 leukocytes from each
455 sample. The increased dispersion coefficient during the work week occurred in the
456 subgroup with the highest exposure (i.e., sprayers at Facility A), which suggested to
457 the authors that comets in a sub-population of cells were affected by 1-BP. However,
458 confirming this conclusion would require data indicating that dispersion coefficients
459 were not increased during the week in an unexposed control group, which was not
460 included in this study.

461 Using multiple linear regression models, Toraason *et al.*, found that start-of-week tail
462 moment was significantly associated with serum Br quartiles ($p < 0.05$). End-of-week
463 comet tail moment was also significantly associated with 1-BP TWA quartiles and
464 serum Br quartiles ($p < 0.05$). For quartile analysis, all workers were placed into four

465 exposure groups of equal number, (low, medium low, medium high and high
466 exposure). Other positive associations were observed with tail moment (e.g., 1-BP
467 TWA log ppm, serum Br log mg/dl), but did not reach statistical significance. No
468 statistically significant positive associations were found with the dispersion coefficient
469 and exposure. Overall, the authors found the comet assay results inconsistent,
470 providing only limited evidence that 1-BP increased DNA damage in the workers.
471 Possible confounders cited included temporal variation in the comet assay, lack of
472 breathing zone data from some workers, breathing zone concentrations of 1-BP too
473 low to show definitive *in vivo* evidence of DNA damage in leukocytes, and small
474 sample size.

475 The human hepatoma cell line, HepG2, was used to determine if 1-BP can induce
476 DNA single strand breaks in these cells *in vitro* (Hasspieler *et al.*, 2006). In addition,
477 cell viability and altered enzyme activity were measured using the neutral red uptake
478 assay and the ethoxyresorufin *O*-deethylase assay, respectively. The tests were
479 performed at 1-BP concentrations of 0, 100, 200, 250, 300, 400, and 500 ppm on
480 HepG2 cells. 1-BP did not induce an increase in single strand breaks at the
481 concentrations tested. Cell viability was reduced at the highest concentration (500
482 ppm), and no effect on enzyme activity was observed.

483 **DNA Adduct formation in vitro and in vivo**

484 Two studies have demonstrated the formation of N⁷ guanine adducts both *in vitro* and
485 *in vivo* following 1-BP exposure (Thapa, 2016; Nepal *et al.*, 2019). N⁷-guanine
486 adducts have been shown to be excellent biomarkers for internal exposure to direct-
487 acting and metabolically activated carcinogens (Boysen *et al.*, 2009). However, N⁷-
488 guanine adducts themselves generally do not persist, and are not likely to be
489 mutagenic.

490 Thapa *et al.*, (2016) observed the formation of N⁷-guanine adduct (i.e., N-propyl
491 guanine adduct) when 1-BP was incubated with 2'-deoxyguanosine. Subsequently,
492 1-BP was incubated with calf thymus DNA *in vitro* under physiological conditions for
493 18 hr, following which unreacted 1-BP was removed and the reactant subjected to
494 thermal hydrolysis to look for the presence of N-propyl guanine. The adduct was
495 found to be generated in a dose-dependent manner without enzymatic support,
496 suggesting that 1-BP could be a direct-alkylating agent.

497 Adult male Sprague-Dawley rats were injected intraperitoneally with 500 or 1000
498 mg/kg 1-BP once or daily for three days and then necropsied six hours following the
499 last injection to determine the extent of N-propyl guanine adduct formation in DNA of
500 several organs (Nepal *et al.*, 2019). The highest levels of adduct formation (in

501 pmole/g DNA) was found in the liver, followed by spleen and kidney. Smaller
502 amounts were observed in testis and lung, and none was detected in heart tissue.
503 DNA adduct formation in tissues increased in both a time- and dose-dependent
504 manner.

505 In a subsequent study by Nepal et al. (2019), 1-BP was incubated *in vitro* with calf
506 thymus DNA, both with and without liver homogenate. Formation of N-propyl
507 guanine was not affected by the addition of liver homogenate, suggesting to the
508 authors that 1-BP can act as a direct alkylating agent.

509 **Induction of DNA repair**

510 In addition to the DNA single strand break test conducted by Hasspieler et al. (2006),
511 the ability of 1-BP to induce DNA repair in human HepG2 hepatoma cells was
512 investigated over the same concentration levels (0, 100, 200, 250, 300, 400, and 500
513 ppm). Repair of DNA was measured by incorporation of labelled healthy nucleotides
514 ($[^3\text{H}]$ -thymidine) at previously damaged DNA sites. 1-BP did not induce an increase
515 in DNA repair over the range of concentrations tested.

516 **Bacterial mutation tests**

517 Barber and coworkers were able to show mutagenic activity of 1-BP in the Ames
518 Salmonella test when evaporation of 1-BP was prevented by using a closed system
519 (Barber et al., 1981). The plated bacteria were exposed to 1-BP vapor at
520 concentrations of 1.1, 2.3, 4.9, 9.0, and 20.3 $\mu\text{moles/plate}$ (135, 283, 603, 1107, and
521 2497 $\mu\text{g/plate}$, respectively) for a period of 48 hours. Bacterial strains tested
522 included Salmonella typhimurium strains TA1535, TA1537, TA1538, TA98, and
523 TA100. 1-BP was mutagenic only in *S. typhimurium* TA1535 and TA100 strains,
524 showing similar activity in the presence and the absence of induced-rat liver
525 activation enzymes (S9). This finding indicated it is a direct acting mutagen.

526 The mutagenicity of 1-BP was also tested in the five *S. typhimurium* strains (i.e.,
527 TA98, TA100, TA1535, TA1537 and TA1538) and *Escherichia coli* strain WP2 uvrA,
528 with and without S9 mix, by BioReliance (2015). 1-BP concentrations tested were
529 1.5, 5.0, 15, 50, 150, 500, 1500 and 5000 $\mu\text{g/plate}$ in the initial toxicity-mutation
530 assay, and 50, 150, 500, 1500, 2000, 3000, and 5000 $\mu\text{g/plate}$ in the confirmatory
531 mutagenicity assays. The highest concentration of 5000 $\mu\text{g/plate}$ resulted in
532 cytotoxicity. The study also used a closed system to prevent volatilization of 1-BP,
533 but consisted of a preincubation step of mixing bacteria with 1-BP liquid in screw cap
534 tubes for 90 minutes, followed by plating and incubation for 48 to 72 hours. Unlike
535 the results of Barber et al. (1981), no evidence of mutagenicity was observed in any

536 strain, with or without S9 mix. This study has not been published in a peer-reviewed
537 publication, but was extensively summarized by U.S. EPA (2020a).

538 1-BP was not mutagenic in either of two independent bacterial mutagenicity assays,
539 each conducted with and without S9 (NTP, 2011). Bacterial strains tested included
540 *Salmonella typhimurium* strains TA97, TA98, TA100, and TA1535, and *Escherichia*
541 *coli* strain WP2 *uvrA/pKM101*. 1-BP concentrations tested were 33, 100, 333, 1000,
542 3333, and 10,000 µg/plate. The NTP (2011) did not use a closed system to prevent
543 potential 1-BP loss due to volatilization, as Barber *et al.* had used. NTP suggested
544 volatility as a possible cause of the negative results in the study. Cytotoxicity
545 occurred at high treatment doses, but it is unclear what the actual exposure levels
546 were to the bacteria.

547 U.S. EPA (2020a) noted that among the two closed system studies, the BioReliance
548 (2015) study may have had some method limitations that contributed to a negative
549 finding for mutagenicity. The exposure method by Barber *et al.* (1981) consisted of
550 vapor exposure for 48 hours in a fully enclosed chamber while the BioReliance assay
551 employed screw cap tubes with “minimal” headspace for the 90 minute preincubation
552 step. Analytical concentrations of 1-BP in these preincubation tubes (without
553 metabolic activation) during the confirmatory assays were only 4-37% of target
554 concentrations at the beginning of the preincubation period, and 2-5% of target
555 concentrations by the end of the preincubation period.

556 Alternatively, the demonstration of cytotoxicity at the highest dose in the two
557 mutagenicity studies with all negative results suggests that the absence of
558 mutagenicity did not result from lack of 1-BP in the test medium, but rather from lack
559 of mutagenic activity of 1-BP.

560 **Mammalian cell gene mutation tests**

561 1-BP was investigated for the ability to induce mutations *in vitro* at the thymidine
562 kinase (TK) locus in L5178Y mouse lymphoma cells (Elf Atochem, 1996). The test
563 determines if a substance can induce forward mutation from the parental type (TK^{+/-})
564 to the mutant form (TK^{-/-}), which in a specific medium only allows mutant cells to grow
565 and form colonies. The top dose level of 1-BP used was based on cytotoxicity,
566 identified as ≥10-20% relative survival assessed by relative cloning efficiency. Two
567 independent tests were run, each in the presence or absence of S9 mix, resulting in
568 a total of four tests. A positive response was considered to be a dose-related
569 increase in mutant frequency and/or a reproducible increase in the mutant frequency
570 (at least a doubling compared to control) for at least one dose level.

571 Over a dose range of 125 to 1500 µg/ml 1-BP (specific dose levels not provided),
572 without S9 mix, a reproducible and significant increase in the mutation frequency
573 occurred between 1000 and 1500 µg/ml. The relative cloning efficiency at 1500
574 µg/ml was 21-33%, indicating acceptable viability for the tests. A significant increase
575 in the mutation frequency of both large and small colonies was observed. Small,
576 slow growing colonies are mainly produced by chromosome rearrangements and
577 large colonies are mainly produced by point mutations. With S9 over a 1-BP dose
578 range of 125 to 2500 µg/ml, no increase in mutation frequency was observed in the
579 first test. However, a significant increase in the mutation frequency together with an
580 increase in the number of small colonies was observed at 1500 to 2000 µg/ml in the
581 second test. The relative cloning efficiency at 1500 and 2000 µg/ml was 36 and 9%,
582 respectively.

583 Under the experimental conditions, the authors concluded that 1-BP showed
584 mutagenic activity in their mouse lymphoma assay, especially without S9 mix.
585 Although this study has not been published in a peer-reviewed publication, the NTP-
586 CERHR expert panel (NTP, 2003) found that this study was well conducted and
587 without any perceived weaknesses.

588 **Chromosomal damage**

589 The frequency of micronucleated cells in mouse bone marrow cells was examined
590 following intraperitoneal (IP) injection of Swiss OF1/ICO:OF1 mice (at least 5
591 animals/group) to 600 mg/kg (males) or 800 mg/kg (females) 1-BP (Elf Atochem,
592 1995). Micronuclei are biomarkers of induced structural or numerical chromosomal
593 alterations and are formed when acentric fragments or whole chromosomes fail to
594 incorporate into either of two daughter nuclei during cell division. Initial studies found
595 that exposure of male mice to 800 mg/kg 1-BP by IP injection resulted in mortality, so
596 the dose was reduced to 600 mg/kg for male mice. No increase in micronucleated
597 erythrocytes in bone marrow was observed in either male or female mice. A positive
598 control group treated with cyclophosphamide did show a significant increase in
599 micronucleated erythrocytes. This study has not been published in a peer-reviewed
600 publication, but was summarized by the NTP-CERHR expert panel (NTP, 2003). The
601 panel found the study to be well conducted and without any perceived weaknesses.

602 Mouse peripheral blood was examined for the frequency of micronucleated
603 erythrocytes following 3-month inhalation exposure of male and female B6C3F₁ mice
604 to 62.5, 125, 250, or 500 ppm (314, 629, 1,258, and 2,515 mg/m³, respectively) 1-BP
605 (NTP, 2011). No increases in the frequencies of micronucleated normochromatic
606 erythrocytes were observed in the 1-BP-exposed mice.

607 **Transgenic rodent mutation assay**

608 A gene mutation study was conducted with 1-BP in Big Blue[®] transgenic female
609 B6C3F1 mice to investigate the mutagenic mode of action (Stelljes *et al.*, 2019). The
610 assay measures the mutation frequency in the *c//* gene in any tissue in the body.
611 Female transgenic mice (7 per group) were exposed to 1-BP 6 hours/day, 5
612 days/week, for 28 days at the concentrations used in the NTP (2011) mouse
613 carcinogenicity study – 0, 62.5, 125, and 250 ppm. Another group of female
614 transgenic mice were exposed to N-ethyl-N-nitrosourea, a known mutagen, which
615 acted as a positive control group. At the end of the exposures, the lungs, colon and
616 liver of the mice were collected and analyzed for increased *c//* mutant frequency.
617 1-BP did not induce *c//* mutations different from negative control values in any of the
618 three organs examined, while *c//* mutations were increased in all three tissues of the
619 positive control. The authors indicated this was evidence that 1-BP is not a direct
620 acting genotoxic carcinogen.

621 In their review of 1-BP toxicity, U.S. EPA (2020a) noted some limitations in this study
622 that may have resulted in the negative finding for mutagenicity. The maximum
623 tolerated dose, in the range of 400 to 500 ppm, was not evaluated in the female
624 mice. Also, an exposure time of 28 days, followed by a post-exposure observation
625 period of three days may have been too short to detect mutations in slower dividing
626 tissues. Generally, a post-exposure period of 28 days is recommended to allow
627 fixing of DNA damage into stable mutations in slower dividing tissues. Other
628 limitations included no evaluation of male and female rats, which also exhibited an
629 increase in cancer incidence, or examination of other tissues, such as skin, large
630 intestine and pancreas, which are target sites for tumors in rats. Finally, no
631 carcinogenic/mutagenic structural analogs of 1-BP have been tested with the Big
632 Blue[®] assay. If negative results were found with 1-BP analogs such as bromoethane,
633 it might be concluded that these assays are not suitable for assessing the
634 mutagenicity of 1-BP.

635

636 **Table 4. Genotoxicity and mutagenicity study summaries for 1-BP**

Cell type or species/strain	Description	Metabolic Activation		Reference
		without	with	
DNA strand-break tests (comet assay or other DNA damage assay)				
Human leukocytes (<i>in vitro</i>)	Comet assay	+/-	NA	Toraason <i>et al.</i> , (2006)
Human leukocytes of exposed workers (<i>in vivo</i>)	Comet assay	+/-	NA	Toraason <i>et al.</i> , (2006)
Human HepG2 cells	Hydroxylapatite DNA chromatography	-	NA	Hasspieler <i>et al.</i> , (2006)
DNA adduct formation				
Calf thymus DNA	N-propyl guanine adduct formation	+	NA	Thapa <i>et al.</i> , 2016
Calf thymus DNA	N-propyl guanine adduct formation	+	+	Nepal <i>et al.</i> , 2019
Male rats (<i>in vivo</i>)	N-propyl guanine adducts in tissues	+	NA	Nepal <i>et al.</i> , 2019
Induction of DNA repair (Unscheduled DNA synthesis)				
Human HepG2 cells	[³ H]-thymidine incorporation	-	NA	Hasspieler <i>et al.</i> , (2006)
Bacterial mutation tests				
<i>S. typhimurium</i>	TA98	-	-	Barber <i>et al.</i> , (1981)
	TA100	+	+	
	TA1535	+	+	
	TA1537	-	-	
	TA1538	-	-	
<i>S. typhimurium</i> and <i>E. coli</i>	TA98, TA100, TA1535, TA1537, <i>E. coli</i> WP2 <i>uvrA</i>	-	-	BioReliance, (2015)
<i>S. typhimurium</i> and <i>E. coli</i>	TA97, TA98, TA100, TA1535, and WP2 <i>uvrA/pKM101</i>	-	-	NTP (2011a)
Mammalian cell gene mutation test				
L5178Y mouse lymphoma cells	parental type TK ^{+/-} to mutant form TK ^{-/-} forward mutation	+	+	Elf Atochem (1996)
Chromosomal damage				
Mouse bone marrow cells (<i>in vivo</i>)	Micronuclei after i.p injection	-	NA	Elf Atochem (1995)
Mouse peripheral erythrocytes (<i>in vivo</i>)	Micronuclei after 3-month inhalation exposure	-	NA	NTP (2011a)
Transgenic rodent mutation assay				
Big Blue® transgenic female B6C3F1 mice (<i>in vivo</i>)	<i>cII</i> gene mutation frequency in lung, colon, and liver	-	NA	Stelljes <i>et al.</i> , 2019

637 +/-: equivocal; NA: not applicable

638 **Dominant lethal mutations in rodents**

639 The dominant lethal test identifies germ-cell mutagens by measuring embryonic
640 death of the progeny of treated males caused by an absorbed chemical penetrating
641 gonadal tissue and producing chromosomal breakage in parent germ cells. This test
642 does not detect somatic mutations, so it is not included in Table 4. It also has a low
643 sensitivity for detecting small increases in induced mutation frequency due to a high
644 rate of spontaneous mutations.

645 Dominant lethal studies were conducted in male Sprague Dawley rats (15/chemical)
646 with 5 halogenated 3-carbon compounds (including 1-BP) that were similar in
647 structure to 1,2-dibromo-3-chloropropane (DBCP), a compound that is known to
648 cause dominant lethal mutations (Saito-Suzuki *et al.*, 1982). Treated males were
649 exposed by gastric intubation to 400 mg/kg 1-BP for 5 consecutive days. Males were
650 then mated with untreated females during ten sequential mating periods of a week
651 each. 1,2,3-Tribromopropane (50 mg/kg daily) acted similarly to DBCP (50 mg/kg) in
652 causing dominant lethal mutations based on dead embryonic implants, especially in
653 the early spermatid stage. 1,2-Dibromopropane (200 mg/kg) gave a minimal
654 response. 1-BP (400 mg/kg), 1,2,3-trichloropropane (80 mg/kg) and 1-chloropropane
655 (1,000 mg/kg) were inactive.

656 1-BP was administered orally to ICR male mice (20/group) at 300 or 600 mg/kg for 10
657 days before mating to investigate the potential of 1-BP to induce dominant lethality
658 (Yu *et al.*, 2008). Males were mated with untreated females during six sequential
659 mating periods of a week each. Males were necropsied at the end of mating and the
660 pregnant females on days 15-17 of gestation. A positive control group (40 mg/kg
661 cyclophosphamide administered IP) was included and followed the same mating
662 schedule. There were no treatment-related changes in clinical signs, gross findings,
663 mating index, gestation index, number of corpora lutea and implantations, pre-
664 implantation loss, live fetuses, resorptions, dead fetuses, and post-implantation loss
665 at either 1-BP dose that would indicate dominant lethality. An increase in pre-
666 implantation loss during the fifth week was attributed to treatment-related low sperm
667 quality. In the positive control group mating and gestation indices were normal, but a
668 decrease in the number of implantations and an increase in pre-implantation loss and
669 fetal deaths were observed during the first 2 or 3 weeks, resulting in a markedly
670 increased dominant lethal mutation rate for the first 3 weeks.

671 **Other Supporting Data**

672 **Cancer Bioassays with Structurally Related Compounds**

673 Previous long-term rodent toxicology and carcinogenesis studies with brominated
674 hydrocarbons have been conducted by the NTP, which have resulted in tumors in the
675 same organs and tissues as those following 1-BP exposure (see below).

676 Chronic inhalation exposure of F344 rats and B6C3F₁ mice to 1,2-dibromoethane
677 resulted in significantly increased incidences of alveolar/bronchiolar adenomas and
678 carcinomas in male and female mice and female rats (NTP, 1982a). 1,2-
679 Dibromoethane exposure also led to an increased incidence of mesotheliomas of the
680 tunica vaginalis (epididymis) in male rats. Similarly, long-term inhalation exposure of
681 F344 rats and B6C3F₁ mice to 1,2-dibromo-3-chloropropane led to increased
682 incidences of alveolar/bronchiolar adenomas and alveolar/bronchiolar adenomas or
683 carcinomas in male and female mice (NTP, 1982b).

684 In separate two-year oral gavage studies in F344/N rats and B6C3F₁ mice treated
685 with bromodichloromethane (NTP, 1987) and tribromomethane (NTP, 1989),
686 significantly increased incidences of adenomatous polyp and adenocarcinoma, and
687 adenomatous polyps or adenocarcinomas (combined), respectively, were observed
688 in the large intestine of male and female rats. The occurrence of this rare tumor
689 following exposure to brominated compounds structurally related to 1-BP
690 strengthened NTPs conclusion for “some evidence” for adenoma of the large
691 intestine in male rats resulting from 1-BP exposure (NTP, 2011).

692 **Genotoxicity of 1-BP metabolites**

693 1-BP metabolites formed by CYP-mediated oxidation in rodents include α -
694 bromohydrin and glycidol (Ishidao *et al.*, 2002; Garner *et al.*, 2007). Both are direct-
695 acting mutagens that induce DNA damage in bacteria (Stolzenberg and Hine, 1979;
696 IARC, 2000). In addition, glycidol has been shown to be mutagenic in mammalian
697 cells, and induce DNA damage and chromosomal damage *in vitro* in rodent and
698 human cells. *In vivo* studies in mice indicate that glycidol induces micronucleus
699 formation but not chromosomal aberrations (IARC, 2000).

700 **Immune System and Cancer**

701 Inflammation is a precursor of many diseases including several types of cancer
702 (Coussens and Werb, 2002; Colotta *et al.*, 2009; Korniluk *et al.*, 2017). In the NTP
703 (2011) carcinogenicity study, 1-BP produced an inflammatory reaction in the
704 respiratory system of rats and mice, but only female mice developed tumors in the
705 lung. Chronic suppurative inflammation was significantly increased in rats in the

706 highest 1-BP exposure group. These lesions were characterized by the presence of
707 Splendore Hoeppli (S-H) material, which were primarily found in the nose and skin of
708 affected animals and typically surrounds or is adjacent to the agent causing S-H
709 bodies (i.e., fungi, helminthes or bacteria). Immunosuppression has been suggested
710 as a cause for the development of these lesions (Morgan *et al.*, 2011). S-H bodies
711 following 1-BP exposure in rats and mice have only occurred in rats, although
712 immunotoxicity tests have shown that both rodent species were immunosuppressed
713 after 1-BP exposure (Anderson *et al.*, 2010). Species differences in the presence of
714 opportunistic bacteria, or differences in innate resistance to infection, have been
715 postulated as possible causes.

716 Han *et al.* (2008) investigated the proinflammatory effects of 1-BP *in vitro* in mouse
717 macrophages. 1-BP induced the production of nitric oxide (NO) and proinflammatory
718 cytokines, including interleukin (IL)-1 β , IL-6, and tumor necrosis factor (TNF)- α in the
719 macrophages. The expression levels of these genes increased in a dose-dependent
720 manner (from 0.5 to 10 micromolar in cell culture). Nuclear transcription factor- κ B
721 (NF- κ B) sites were identified in the promoter of the inducible nitric oxide synthase
722 (iNOS) and proinflammatory cytokine genes. The authors noted that NO synthesized
723 by iNOS is considered an important mediator of carcinogenesis that may elevate
724 cancer progression, and that overexpressed iNOS has been found in human breast
725 and colorectal tumors (Thomsen *et al.*, 1995; Hao *et al.*, 2001).

726 **Induction of cancer stem cells in colorectal cancer**

727 Colorectal cancer has a high relapse rate, attributed to the high proportion of cancer
728 stem cells, or self-renewing cells within tumors. Cho *et al.* (2017) investigated the
729 effects of 1-BP and similar brominated compounds on the “stemness” in human
730 colorectal cancer cell lines [although not explicitly defined by the authors, OEHHA
731 notes that “stemness” generally refers to a state of a cell characterized by a high
732 degree of plasticity, where plasticity is the property of being transmutable into either a
733 less committed or a more committed state]. 1-BP was observed to increase the
734 spheroid formation in colorectal cancer cells (CSC221, DLD1, Caco2, and HT29
735 cells) *in vitro*, which is a measure of the ability to induce cancer cell stemness. 1-BP
736 also induced the expression of cancer stemness markers, including ALDH-1, CD133,
737 Lgr-5, and Msi-1, at both the mRNA and protein levels. Finally, 1-BP was found to
738 increase the transcriptional activity of the Hedgehog, Notch, and Wnt signaling
739 pathways, which supports the hypothesis that induction of cancer cell stemness by
740 1-BP occurs via these signaling pathways. The authors concluded that 1-BP and
741 other related compounds have the potential to promote cancer stemness.

742 **IV. CANCER HAZARD EVALUATION**

743 The chronic toxicity/carcinogenicity studies by NTP in rats and mice (Morgan *et al.*,
744 2011; NTP, 2011) are the only source of carcinogenicity data for 1-BP. Although
745 there is human exposure to 1-BP (Ichihara *et al.*, 2004a; Hanley *et al.*, 2006; Hanley
746 *et al.*, 2009; 2010), widespread exposure has occurred only relatively recently. The
747 initial reports of high occupational exposure were based on workers studied in the
748 1990s (Ichihara *et al.*, 2004a). Human cancer generally has a long latency period, so
749 occupational exposure data of sufficient exposure duration may not yet exist.

750 Lifetime exposure to 1-BP in rodents resulted in tumors in male and female rats, and
751 female mice (Morgan *et al.*, 2011; NTP, 2011). Tumors that the NTP concluded were
752 a result of 1-BP exposure included adenomas of the large intestine in male and
753 female rats, skin tumors of the epithelium in male rats, and alveolar/bronchiolar
754 adenoma or carcinoma of the lungs in female mice. However, the low incidence of
755 adenoma in the large intestine of male rats resulted in no significant difference
756 relative to controls, and no significant positive trend. The low tumor incidence will not
757 contribute to the overall cancer potency, so OEHHA did not use this particular tumor
758 data to derive a cancer potency. The tumors OEHHA identified as being suitable for
759 cancer potency determination were adenomas of the large intestine in female rats,
760 the combined skin neoplasms of epithelial origin in male rats (keratoacanthoma,
761 basal cell adenoma or carcinoma, and squamous cell papilloma or carcinoma), and
762 lung tumors in female mice (alveolar/bronchiolar adenomas or carcinomas
763 combined).

764 Supporting data for the carcinogenicity of 1-BP included some evidence for
765 genotoxicity and mutagenicity in cell culture studies. *In vitro* exposure of cultured
766 human leukocytes to 1-BP resulted in equivocal evidence of increased DNA damage
767 by the comet assay (Toraason *et al.*, 2006). 1-BP was mutagenic in a closed system
768 bacterial Ames assay with and without S9, suggesting 1-BP is a direct acting
769 mutagen (Barber *et al.*, 1981). 1-BP also induced mutations *in vitro* at the thymidine
770 kinase (TK) locus in L5178Y mouse lymphoma cells (Elf Atochem, 1996). Lastly, 1-
771 BP has been shown to produce DNA adducts both *in vitro* and *in vivo* (Thapa, 2016;
772 Nepal *et al.*, 2019)

773 In addition, long-term rodent exposure studies with structurally-related brominated
774 compounds, including 1,2-dibromoethane, 1,2-dibromo-3-chloropropane,
775 bromodichloromethane and tribromomethane, have resulted in similar tumors as that
776 caused by 1-BP. *In vivo* metabolism of 1-BP resulted in the production of direct
777 acting mutagens such as α -bromohydrin and glycidol in rodents (Ishidao *et al.*, 2002;

778 Garner *et al.*, 2007). Finally, 1-BP increased the “stemness” in human colorectal
779 cancer cell lines.

780 **V. QUANTITATIVE CANCER RISK ASSESSMENT**

781 **Effective Tumor Incidences**

782 The effective tumor incidences in rats and mice (Tables 5A and 5B, respectively)
783 were determined from individual animal survival data of the NTP study located in the
784 Chemical Effects in Biological Systems (CEBS) database for rats (NTP-CEBS,
785 2011a) and mice (NTP-CEBS, 2011b). The effective tumor incidence is the number
786 of tumor-bearing animals (numerator) over the number of animals alive at the time of
787 first occurrence of the tumor (denominator). In most cases, the effective tumor
788 incidences were used to calculate the cancer slope factor for 1-BP. This method of
789 tallying tumor incidence removes animals from the assessment that died before they
790 are considered at risk for tumor development. Tables 5A and 5B do not include
791 treatment-related tumors that were of very low incidence (e.g., large intestine tumors
792 in male rats) or tumors that were of equivocal significance (e.g., malignant
793 mesothelioma and pancreatic islet tumors in male rats, and skin tumors in female
794 rats).

795

796 **Table 5A. Adjusted tumor incidence in rats exposed to 1-BP for two years (NTP, 2011b)^{a,b}**

Sex and Species	Tumor Type	Incidence by concentration				Statistical p-values for pairwise comparison with controls			
		0 ppm, 0 mg/m ³	125 ppm, 629 mg/m ³	250 ppm, 1258 mg/m ³	500 ppm, 2515 mg/m ³	Trend ^c	0 ppm, 0 mg/m ³	125 ppm, 629 mg/m ³	250 ppm, 1258 mg/m ³
Male Rats	Skin: Basal Cell Adenoma	0/46	1/42	2/39	1/36	0.191	0.477	0.208	0.439
	Skin: Basal Cell Carcinoma	0/28	2/31	1/26	2/21	0.089	0.272	0.481	0.179
	Skin: Keratoacanthoma	0/49	3/49	6/49*	6/44**	0.006	0.121	0.013	0.009
	Skin: Squamous Cell Carcinoma	1/37	1/34	0/29	2/29	0.881	0.732	1.000	0.408
	Skin: Keratoacanthoma or Squamous Cell Carcinoma	1/49	4/49	6/49	8/44*	0.004	0.181	0.056	0.010
	Skin: Keratoacanthoma, Basal Cell Adenoma, Basal Cell Carcinoma, or Squamous Cell Carcinoma	1/49	7/49*	9/49**	10/44**	0.003	0.030	0.008	0.002
Female Rats	Large Intestine (Colon or Rectum): Adenoma	0/45	1/43	2/41	5/36*	0.001	0.489	0.224	0.015

797
 798 **Table 5B. Adjusted tumor incidence in female mice exposed to 1-BP for two years (NTP, 2011b)^{a,b}**

Sex and Species	Female Mouse	Incidence by concentration				Statistical p-values for pairwise comparison with controls			
		0 ppm, 0 mg/m ³	62.5 ppm, 314 mg/m ³	125 ppm, 629 mg/m ³	250 ppm, 1258 mg/m ³	Trend ^c	62.5 ppm, 314 mg/m ³	125 ppm, 629 mg/m ³	250 ppm, 1258 mg/m ³
Female Mice	Lung: Alveolar/Bronchiolar Adenoma	1/41	6/46	4/42	10/47**	0.006	0.075	0.187	0.007
	Lung: Alveolar/Bronchiolar Carcinoma	0/36	7/42*	5/38*	4/43	0.250	0.010	0.031	0.082
	Lung: Alveolar/Bronchiolar Adenoma or Carcinoma	1/41	9/46*	8/42*	14/47**	0.001	0.012	0.016	<0.001

799 (a) Incidence ratio after adjusting for intercurrent mortality using the effective number adjustment method (i.e., number alive on day of first tumor).
 800 (b) * = $p < 0.05$, ** = $p < 0.01$; p-value indicators are from pairwise comparisons with controls using Fisher exact tests performed by OEHHHA;
 801 (c) p-values in the trend column are for the Cochran-Armitage trend test performed by OEHHHA.

802 **Cancer Slope Factor Derivation**

803 For the derivation of the CSF, 1-BP chamber concentrations of 0, 62.5 (mice only),
804 125, 250 and 500 (rats only) ppm were time-adjusted and converted to mg/m³ (6.17
805 hours/24 hours × 5 days/7 days × 5.03 mg/m³ / ppm) to extrapolate from the
806 intermittent chamber exposure conditions to a continuous exposure over the life span
807 of the animals (*i.e.*, to simulate an annualized average air concentration). The time-
808 adjusted concentrations were 0, 57.73, 115.46, 230.92, and 461.83 mg/m³.

809 The average daily dose, in mg/kg BW-day, is used for calculating the cancer
810 potencies. To calculate the daily dose, the average body weight of the rats and mice
811 over the duration of the study is used to determine the inhalation rate (IR). The
812 weighted average lifetime body weights for control rats of both sexes and female
813 mice were calculated from the NTP (2011) study based on the regular reporting of
814 group mean body weights every 1 to 4 weeks during the 2-year exposure. Body
815 weights and daily dose for male mice were not calculated since no 1-BP-related
816 carcinogenicity was observed in male mice. The average body weights were 440.6,
817 284.9, and 47.4 g for the control male rats, female rats, and female mice,
818 respectively.

819 A comprehensive analysis of rat minute volume data was undertaken by OEHHA
820 (2018) to update the IR equation by Anderson (1983) and is shown below (Eq. 6-1a).
821 The analysis incorporates studies published since 1988 that more accurately reflect
822 true resting IRs of rats. For mice, the IRs were determined using the equation (Eq. 6-
823 1b) by Anderson (1983). These formulas reflect proportional differences of body
824 weight (BW^{2/3}) on the respiratory rate within a species.

825 Rats: $IR (m^3/day) = 0.702 m^3/day\text{-kg} \times (BW)^{2/3}$ Eq. 6-1a

826 Mice: $IR (m^3/day) = 0.0345 m^3/day \times (BW / 0.025 \text{ kg})^{2/3}$ Eq. 6-1b

827 The calculated average daily IRs during the 1-BP exposures are 0.406, 0.304, and
828 0.0528 m³/day for male and female rats and female mice, respectively. The average
829 daily doses (shown in Table 6) could then be calculated with the following equation:

830 $Dose (mg/kg \text{ BW-day}) = IR \times C / BW$ Eq. 6-2

831 Where:

832 $C = \text{time-adjusted 1-BP concentration (mg/m}^3\text{)}$

833

834 **Table 6. Calculated average daily exposed dose of 1-BP in rats and mice**

Species sex	1-BP Chamber Concentration (mg/m³)				
	0	314	629	1,258	2,515
Daily Exposed Dose (mg/kg-day)					
<u>Rats</u>					
Males	0	-	106.39	212.78	425.56
Females	0	-	123.20	246.40	492.79
<u>Mice</u>					
Females	0	64.31	128.61	257.22	-

(-): no rat/mouse exposure group at this concentration

835 The United States Environmental Protection Agency’s (US EPA’s) Benchmark dose
 836 (BMD) methodology and Benchmark Dose Modeling Software (BMDS, version 3.2)
 837 were used to perform dose-response extrapolation (US EPA, 2020b). BMD analyses
 838 were run for the tumor data that were identified as treatment-related and showed a
 839 statistically significant increase above control values and a statistically significant
 840 positive trend (See Table 7). Where tumors of the same histological cell type or
 841 tissue type (e.g., skin tumors of epithelial origin; pulmonary alveolar/bronchiolar
 842 adenomas and carcinomas) are observed, the combined incidence is used for dose-
 843 response assessment.

844 The multistage-cancer polynomial model was fit to the female rat and female mouse
 845 data. Survival was unaffected by 1-BP exposure in these groups, so the effective
 846 tumor incidences were used to derive the cancer potencies. The multistage Weibull
 847 model was used for the male rat tumor data due to decreased survival in the 500
 848 ppm group relative to the control group (US EPA, 2017). OEHHA applies this
 849 adjustment in lifetime rodent exposure studies when 1) survival is reduced by about
 850 15% or greater compared to controls before week 85, and 2) less than 85% of these
 851 early deaths occur in animals that have treatment-related tumors. The 500 ppm
 852 group displayed reduced survival of 10-11% between week 70 and week 80. The
 853 difference in survival increased to 22% at week 83, and then varied mostly between
 854 16-22% to the end of the study at week 104. Nine of 37 early deaths were due to
 855 chronic inflammation in various organs (lung, nose, skin, and bone) that was
 856 treatment-related; the remaining early deaths were due to various types of neoplasia
 857 that were not treatment-related. Only seven of the male rats that died early had a
 858 treatment-related tumor (i.e., large intestine or skin tumor), none of which were the
 859 cause of death.

860 For large datasets such as those by the NTP, a Benchmark Dose Response (BMR)
 861 of 5% is recommended by OEHHA (2008) for the BMD, and 95% lower confidence
 862 bound (BMDL). First and 2nd degree polynomial multistage models were run for all

863 suitable tumor data sets, and the most appropriate model fit was chosen based on
864 BMD technical guidance (US EPA, 2012). The degree of polynomial chosen was 1 in
865 all cases.¹ The resulting BMD and BMDL values for each tumor type are shown in
866 Table 7. The rodent CSFs, in units of (mg/kg-day)⁻¹, are calculated as 0.05/BMDL,
867 where 0.05 represents the 5% tumor response, or BMR. The rodent CSFs (CSF(a))
868 were then converted to human equivalents (CSF(h)) using body weight (BW^{3/4})
869 scaling:

$$870 \quad \text{CSF(h)} = \text{CSF(a)} \times (\text{BW(h)} / \text{BW(a)})^{1/4} \quad \text{Eq. 6-3}$$

871 Lifetime body weights for rodents (BW(a)) were calculated from the NTP (2011) study
872 as described above. The default body weight for humans (BW(h)) is 70 kg. The
873 body weight scaling factor assumes that mg/surface area/day is an equivalent dose
874 between species (OEHHA, 2009). Using this interspecies scaling factor is preferred
875 by OEHHA because it is assumed to account not only for pharmacokinetic
876 differences (e.g., breathing rate, metabolism), but also for pharmacodynamic
877 considerations, i.e., tissue responses to chemical exposure (US EPA, 2005).

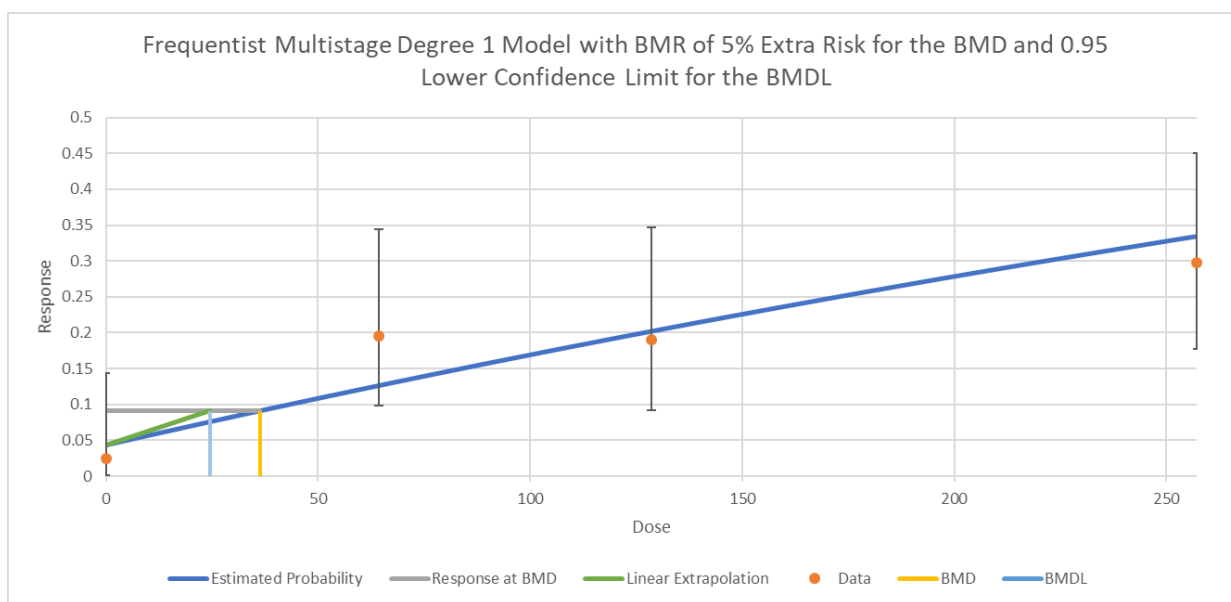
878 When extrapolating to the human equivalent dose using the body weight scaling
879 factor, pulmonary alveolar/bronchiolar adenoma and/or carcinoma combined in
880 female mice provided the highest CSF(h) value of 0.013 (mg/kg-day)⁻¹ (CSFs
881 rounded to two significant figures in the final assessment), establishing this tumor in
882 female mice as the most sensitive endpoint for 1-BP-induced carcinogenicity. The
883 multistage model fit to the female mouse tumor data is shown in Figure 2.

¹ For female mice alveolar/bronchiolar tumors (See Table 7 below), BMD guidance suggested a 2nd degree multistage model provided the best fit to the data. However, this choice was based on an Akaike Information Criterion value that was only 0.0000001 lower than the 1st degree multistage model. Due to nearly identical model fits, OEHHA chose the simpler 1st degree model to calculate the BMDL.

884 **Table 7. BMD5 Modeling Results**

Tumor Type	Sex and Species	Akaike Information Criterion	p-value	BMD (mg/kg-day) ^a	BMDL (mg/kg-day)	CSF - Rodent (mg/kg-day) ⁻¹	CSF - Human (mg/kg-day) ⁻¹
Skin tumors	Male Rats	151.75	NA ^a	57.57	33.43	0.001496	0.0053
Large Intestine	Female Rats	56.84	0.95	202.43	119.07	0.000420	0.0017
Alveolar/ bronchiolar	Female Mice	159.53	0.26	36.34	24.54	0.00204	0.013

885 ^a Not applicable for the multistage Weibull model



886 **Figure 2. Multistage model plot fit to the female mouse lung tumor data for 1-**
 887 **BP. The Multistage polynomial degree 1 model with BMR of 5% extra risk for**
 888 **the BMD and 95% lower confidence bound (BMDL).**
 889

890 **Inhalation Unit Risk Factor**

891 The Inhalation Unit Risk (IUR) describes the excess cancer risk associated with
892 inhalation exposure to a concentration of $1 \mu\text{g}/\text{m}^3$ and is derived from the human
893 CSF(h):

894
$$\text{IUR} = (\text{CSF}(\text{h}) \times \text{BR}) / (\text{BW} \times \text{CF}) \qquad \text{Eq. 6-4}$$

895 Where:

896 BR = mean human breathing rate ($20 \text{ m}^3/\text{day}$)

897 BW = mean human body weight (70 kg)

898 CF = mg to μg conversion factor of 1,000

899 Use of the equation above with the 1-BP CSF of $0.013 (\text{mg}/\text{kg}\text{-day})^{-1}$ results in a
900 calculated IUR of $0.0000037 (\mu\text{g}/\text{m}^3)^{-1}$, which can also be expressed as 3.7×10^{-6}
901 $(\mu\text{g}/\text{m}^3)^{-1}$. Thus, the extra cancer risk associated with continuous lifetime exposure to
902 $1 \mu\text{g}/\text{m}^3$ 1-BP is 3.7 in a million.

903 **VI. CONCLUSIONS**

904 Two-year 1-BP inhalation studies conducted by the NTP established evidence of
905 carcinogenicity in male and female rats, and female mice. Supporting evidence for
906 the carcinogenicity of 1-BP include some positive genotoxic results from *in vitro*
907 studies, a positive *in vivo* study for DNA adduct formation, development of similar
908 tumors in long-term rodent exposure studies by structurally related brominated
909 compounds, and CYP-mediated oxidation of 1-BP to known mutagenic compounds.
910 Rodent CSFs were calculated from the NTP tumor incidence data for each tumor
911 type in each affected species and sex. This was performed by calculating the lower
912 95% confidence limit on the inhalation concentration associated with a 5% tumor
913 response (BMDL) using the multistage or Weibull cancer models in Benchmark Dose
914 Software (BMDS) version 3.1 (US EPA, 2020b). Linear extrapolation from the BMDL
915 to the origin was used to determine the slope of the dose-response curve for low
916 level exposure, the inhalation CSF. The rodent CSFs were then converted to human
917 equivalent exposure levels using body weight scaling to the $3/4$ power. The CSF used
918 for 1-BP, based on the most sensitive species and sex, is $0.013 (\text{mg}/\text{kg}\text{-day})^{-1}$ for
919 pulmonary alveolar/bronchiolar adenomas or carcinomas combined in female mice.
920 An IUR of $3.7 \times 10^{-6} (\mu\text{g}/\text{m}^3)^{-1}$ was calculated from the CSF using the assumption of a
921 human breathing rate of $20 \text{ m}^3/\text{day}$ and an average human body weight of 70 kg.
922

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