CHLOROETHANE A-1

APPENDIX A. ATSDR MINIMAL RISK LEVEL WORKSHEETS

MRLs are derived when reliable and sufficient data exist to identify the target organ(s) of effect or the most sensitive health effect(s) for a specific duration for a given route of exposure. An MRL is an estimate of the daily human exposure to a hazardous substance that is likely to be without appreciable risk of adverse noncancer health effects over a specified route and duration of exposure. MRLs are based on noncancer health effects only; cancer effects are not considered. These substance-specific estimates, which are intended to serve as screening levels, are used by ATSDR health assessors to identify contaminants and potential health effects that may be of concern at hazardous waste sites. It is important to note that MRLs are not intended to define clean-up or action levels.

MRLs are derived for hazardous substances using the NOAEL/uncertainty factor approach. They are below levels that might cause adverse health effects in the people most sensitive to such chemical-induced effects. MRLs are derived for acute (1−14 days), intermediate (15−364 days), and chronic (≥365 days) durations and for the oral and inhalation routes of exposure. Currently, MRLs for the dermal route of exposure are not derived because ATSDR has not yet identified a method suitable for this route of exposure. MRLs are generally based on the most sensitive substance-induced endpoint considered to be of relevance to humans. LOAELs for serious health effects (such as irreparable damage to the liver or kidneys, or serious birth defects) are not used as a basis for establishing MRLs. Exposure to a level above the MRL does not mean that adverse health effects will occur.

MRLs are intended only to serve as a screening tool to help public health professionals decide where to look more closely. They may also be viewed as a mechanism to identify those hazardous waste sites that are not expected to cause adverse health effects. Most MRLs contain a degree of uncertainty because of the lack of precise toxicological information on the people who might be most sensitive (e.g., infants, elderly, nutritionally or immunologically compromised) to the effects of hazardous substances. ATSDR uses a conservative (i.e., protective) approach to address this uncertainty consistent with the public health principle of prevention. Although human data are preferred, MRLs often must be based on animal studies because relevant human studies are lacking. In the absence of evidence to the contrary, ATSDR assumes that humans are more sensitive to the effects of hazardous substance than animals and that certain persons may be particularly sensitive. Thus, the resulting MRL may be as much as 100-fold below levels that have been shown to be nontoxic in laboratory animals.

Proposed MRLs undergo a rigorous review process: Health Effects/MRL Workgroup reviews within the Office of Innovation and Analytics, Toxicology Section, expert panel peer reviews, and agency-wide MRL Workgroup reviews, with participation from other federal agencies and comments from the public. They are subject to change as new information becomes available concomitant with updating the toxicological profiles. Thus, MRLs in the most recent toxicological profiles supersede previously published MRLs. For additional information regarding MRLs, please contact the Office of Innovation and Analytics, Toxicology Section, Agency for Toxic Substances and Disease Registry, 1600 Clifton Road NE, Mailstop S106-5, Atlanta, Georgia 30329-4027.

Chemical Name: Chloroethane CAS Numbers: 75-00-3
Date: January 2025

Profile Status:FinalRoute:InhalationDuration:Acute

MRL: 13 ppm (34 mg/m^3)

Critical Effect: Increased incidence of DFFC of skull bones

Reference: Scortichini et al. 1986

Point of Departure: NOAEL of 1,504 ppm (NOAEL_{HEC} of 376 ppm)

Uncertainty Factor: 30
LSE Graph Key: 13
Species: Mouse

MRL Summary: An acute-duration MRL of 13 ppm was derived for chloroethane based on a NOAEL of 1,504 ppm for developmental effects (increased incidence of DFFC of skull bones) in mouse fetuses exposed for 6 hours/day on GDs 6–15 (Scortichini et al. 1986). The NOAEL was adjusted for continuous exposure and converted to a human equivalent concentration (NOAEL_{HEC}) of 376.0 ppm and divided by a total uncertainty factor of 30 (3 for extrapolation from animals to humans after dosimetric adjustment and 10 for human variability).

Selection of the Critical Effect: Developmental and neurological effects were observed at approximately 5,000 ppm following acute-duration inhalation exposure of chloroethane in animals (Table A-1). Two studies looked at developmental effects of chloroethane. In pregnant mice exposed to chloroethane for 6 hours/day on GDs 6–15, an increase in incidence of DFFC of the skull bones (a developmental delay of ossification of small centers of unossified bone of the skull) was seen in the fetuses at 4,946 ppm (Scortichini et al. 1986). Incidence based on number of fetuses affected was significant for dose-response trend, but not in a pairwise comparison to the control group. Incidence data for number of litters affected were not statistically different from controls (by pairwise or trend tests); however, this effect was considered biologically relevant. The second developmental study reported that mouse fetuses exposed to 15,000 ppm for 6 hours/day on GDs 6–15 appeared normal; however, fetuses were not examined for skeletal or visceral alterations (Dow 1985). This study was not included in Table A-1 because no developmental effects occurred in the only treatment group.

Neurological effects have been reported in humans and several animal species. These effects consisted of a feeling of dizziness and intoxication in humans, hyperactivity in mice and dogs, lethargy in rats, and unsteadiness in guinea pigs; however, the study with the lowest LOAEL for neurological effects (Dow 1985) did not report a NOAEL value. Developmental toxicity (increased incidence of DFFC of the skull in fetuses) was selected as the critical effect because Scortichini et al. (1986) provided the lowest LOAEL with an accompanying NOAEL value.

Table A-1. Selected NOAEL and LOAEL Values in Animals Following Acute-
Duration Inhalation Exposure to Chloroethane

Species	Duration	NOAEL (ppm)	LOAEL (ppm)	Effect	Reference						
Developme	ntal effects										
Mouse (CF-1)	10 days 6 hours/day GDs 6–15	1,504	4,946	Increased incidence of delayed fetal foramina closure (DFFC) of the skull bones (developmental delay of ossification of bones of the skull of fetuses)	Scortichini et al. 1986						
Neurological effects											
Human	Up to 22 minutes	s ND	13,000	LOAEL: subjective feeling of intoxication, increased reaction times	Davidson 1925						
Fisher-344 rat	2 weeks 5 days/week 6 hours/day	3,980	9,980	Slight lethargy	Landry et al. 1982						
CD-1 mouse	10 days 6 hours/day GDs 6–15	ND	5,000	Increased activity and stereotypic behavior (highly repetitive running patterns) in dams	Dow 1985						
Beagle dog	2 weeks 5 days/week 6 hours/day	3,980	9,980	Hyperactivity during exposure in 1/2 dogs	Landry et al. 1982						

Selected study/endpoint for derivation of acute-duration inhalation MRL.

LOAEL = lowest-observed-adverse-effect level; ND = not determined; NOAEL = no-observed-adverse-effect level; SLOAEL = serious LOAEL

Selection of the Principal Study: Of the two developmental studies, Scortichini et al. (1986) was selected as the principal study because visceral and skeletal examinations were performed. In addition, this study provided a LOAEL with an accompanying NOAEL value.

Summary of the Principal Study:

Scortichini BH, Johnson KA, Momany-Pfruenderd JJ, et al. 1986. Ethyl chloride: Inhalation teratology study in CF-1 mice. Dow Chemical Company. Submitted to the U.S. Environmental Protection Agency under TSCA section FYI. OTS0001135. FYI-OTS-0794-1135.

https://ntrl.ntis.gov/NTRL/dashboard/searchResults/titleDetail/OTS0001135.xhtml. April 12, 2023.

Pregnant CF-1 mice (23–26/group) were exposed to 99.9% pure chloroethane at 0, 491, 1,504, or 4,946 ppm 6 hours/day on GDs 6–15. Body weights were recorded on GDs 6, 9, 12, 15, and 18, and food and water intakes were measured. The animals were sacrificed on GD 18 and the following data were recorded: maternal liver weight; number and position of fetuses *in utero*; number of live and dead fetuses; number and position of resorption sites; weight and sex of each fetus; and gross external alterations. Half of each litter was examined for visceral alterations, and the other half was examined for skeletal alterations.

No maternal toxicity (body weight, food and water intake, liver weight) was observed. There were no effects on pregnancy rate, resorption rate, litter size, sex ratio, or fetal body weight. No changes in gross external or visceral alterations were seen in the exposed fetuses compared to controls. An increase in supernumerary ribs was found, although the effect was not indicated as statistically significant. The incidences of fetuses with supernumerary ribs were 2/257, 1/299, 6/311, and 2/242 at 0, 491, 1,504, and 4,946 ppm, respectively. The incidences in litters were 2/22, 1/25, 5/26, and 4/22 at 0, 491, 1,504, and 4,946 ppm, respectively.

A small increase in the incidence of DFFC of the skull bones (developmental delay of ossification of small centers of unossified bone of the skull) was observed at the high dose. The incidence data based on number of fetuses affected were 1/126, 1/142, 1/147, and 5/116 at 0, 491, 1,504, and 4,946 ppm, respectively. The study authors indicated that the foramina data were statistically significant. ATSDR's analysis of the data determined that there was a significant trend (p=0.0488) for the fetal data, but no significance in a pairwise comparison to the control group for the fetal data. Incidence data for number of litters affected were 1/22, 1/24, 1/25, and 5/22 at 0, 491, 1,504, and 4,946 ppm, respectively. The study authors indicated that the litter data were not significantly different from control. ATSDR's analysis determined that there were no significant differences in the pairwise comparison to control or trend test for the litter data.

Selection of the Point of Departure for the MRL: The NOAEL of 1,504 ppm for developmental effects (increased incidence of DFFC of the skull) in fetuses exposed 6 hours/day on GDs 6–15 (Scortichini et al. 1986) was selected as the point of departure (POD). Benchmark dose (BMD) modeling was not done on litter data due to lack of statistical significance by both pairwise comparison and for trend.

Adjustment for Intermittent Exposure: The intermittent NOAEL of 1,504 ppm was adjusted to a 24-hour continuous exposure using the following equation:

$$NOAEL_{ADJ} = NOAEL \times \frac{6 \ hours}{24 \ hours} = 1,504 \ ppm \times \frac{6 \ hours}{24 \ hours} = 376.0 \ ppm$$

Human Equivalent Concentration: The HEC was calculated by multiplying the NOAEL_{ADJ} by the ratio of the chloroethane air:blood partition coefficient for humans and mice. The reported blood:gas (air) partition coefficient ($H_{b/g}$) values for chloroethane are 5.1 for mice and 1.9 or 2.69 for humans (Gargas et al. 1989, 2008; Morgan et al. 1970). Since the ratio of mouse to human blood:gas (air) partition coefficient is >1, a default value of 1 was used. The duration-adjusted LOAEL HEC was 1,237 ppm.

$$NOAEL_{HEC} = NOAEL_{ADJ} \times \frac{\left(H_{b/g}\right)_A}{\left(H_{b/g}\right)_H} = 376.0 \ ppm \times 1 = 376.0 \ ppm$$

Uncertainty Factor: The NOAEL_{HEC} was divided by a total uncertainty factor of 30:

- 3 for extrapolation from animals to humans after dosimetric adjustment
- 10 for human variability

$$MRL = NOAEL_{HEC} \div UFs$$

 $376.0 \ ppm \div 30 = 12.53 \ ppm \approx 13 \ ppm$

Other Additional Studies or Pertinent Information that Lend Support to this MRL: No supporting studies or pertinent information were available.

Chemical Name: Chloroethane CAS Numbers: 75-00-3 Date: January 2025

Final Profile Status: Route: Inhalation Duration: Intermediate

MRL: 13 ppm (34 mg/m^3)

Increased estrous cycle length in mice Critical Effect:

Reference: Bucher et al. 1995

Point of Departure: LOAEL of 15,000 ppm (LOAEL_{HEC} of 3,750 ppm)

Uncertainty Factor: 300 LSE Graph Key: 18 Species: Mouse

MRL Summary: An intermediate-duration inhalation MRL of 13 ppm was derived for chloroethane based on a LOAEL of 15,000 ppm for reproductive effects (increased estrous cycle length) in mice exposed 6 hours/day for 21 days (Bucher et al. 1995). The LOAEL was adjusted to continuous-duration exposure and converted to a human equivalent concentration (LOAEL_{HEC}) of 3,750 ppm and divided by a total uncertainty factor of 300 (10 for use of a LOAEL, 3 for extrapolation from animals to humans after dosimetric adjustment, and 10 for human variability).

Selection of the Critical Effect: The only adverse effect identified in intermediate-duration inhalation studies is related to the estrous cycle. The average duration of the estrous cycle increased significantly by 7% from 5.15 to 5.52 days in mice exposed to 15,000 ppm (only concentration tested) for 21 days, although no consistent effects on estradiol or progesterone were noted (Bucher et al. 1995). Breslin et al. (1988) also studied the length of the estrous cycle in mice after 14 days of exposure to 14,955 ppm for 6 hours/day. No significant increase in the estrous cycle length was seen during exposure compared to pre-exposure (5.0±0.7 days pre-exposure versus 5.6±0.8 days during exposure). The discrepancy between the two studies regarding increased estrous cycle length may be due to duration of exposure (14 versus 21 days) or number of animals studied. Bucher et al. (1995) studied 30 females/group, whereas Breslin et al. (1988) studied 10 females/group. The larger sample size would lend itself to greater statistical power to distinguish differences. In 13-week inhalation studies, no adverse effects (including histopathology on a complete panel of tissues) were observed in rats or mice exposed up to 19,000 ppm (NTP 1989). No adverse effects were observed in rats or rabbits exposed to 10,000 ppm (only concentration tested) for 6.5 months (Dow 1941).

Selection of the Principal Study: The intermediate-duration inhalation study investigating estrous cycle length was selected as the principal study (Bucher et al. 1995). No other intermediate-duration study reported a toxicologically relevant adverse effect.

Summary of the Principal Study:

Bucher JR, Morgan DL, Adkins B, et al. 1995. Early changes in sex hormones are not evident in mice exposed to the uterine carcinogens chloroethane or bromoethane. Toxicol Appl Pharmacol 130:169-173. http://doi.org/10.1006/taap.1995.1022.

Female B6C3F1 mice (30/group) were exposed to chloroethane at 0 or 15,000 ppm 6 hours/day for 21 days. Before the exposures, all the mice were sham-exposed for 21 days, and vaginal cytology studies were completed daily during the sham exposures and during the 21-day exposure period. Body weights

were measured at least weekly. At necropsy, blood was drawn for measurement of serum estradiol and progesterone. The liver and uterus were weighed. The liver, uterus, pituitary gland, adrenal glands, and ovaries were examined microscopically.

No clinical signs of toxicity were observed. Body weights were not different from control (data not shown). In the exposed group, mean estrous cycle length significantly increased from 5.15 ± 0.15 days prior to exposure to 5.52 ± 0.19 days during exposure (a 7% increase). In the control group, mean estrous cycle length was not different between the pre-exposure period and exposure period (5.02 ± 0.2 versus 5.0 ± 0.2 , pre-exposure and exposure, respectively). The proportion of time spent in the stages of the cycle during exposure was significantly different compared to pre-exposure in both the exposed and control group. Mice spent shorter time in metestrus and longer time in the other stages. No significant difference in serum estradiol or progesterone were seen at the end of the exposure. No significant differences in weights of liver, uterus, or ovary were seen compared to control (data not shown). No histological changes were observed in the ovaries, pituitary, uterus, or adrenal glands.

Selection of the Point of Departure for the MRL: The LOAEL of 15,000 ppm for reproductive effects (increased duration of estrous cycle) in mice exposed 6 hours/day for 21 days (Bucher et al. 1995) was selected as the POD.

Adjustment for Intermittent Exposure: The intermittent 6 hours/day LOAEL of 15,000 ppm was adjusted to a 24-hour continuous exposure using the following equation:

$$LOAEL_{ADJ} = LOAEL \times \frac{6 \ hours}{24 \ hours} = 15,000 \ ppm \times \frac{6 \ hours}{24 \ hours} = 3,750 \ ppm$$

Human Equivalent Concentration: The HEC was calculated by multiplying the LOAEL_{ADJ} by the ratio of the chloroethane air:blood partition coefficient for humans and mice. The reported blood:gas (air) partition coefficient ($H_{b/g}$) values for chloroethane are 5.1 for mouse and 1.9 or 2.69 for humans (Gargas et al. 1989, 2008; Morgan et al. 1970). Since the ratio of mouse to human blood:gas (air) partition coefficient is >1, a default value of 1 was used.

$$LOAEL_{HEC} = LOAEL_{ADJ} \times \frac{\left(H_{b/g}\right)_A}{\left(H_{b/g}\right)_H} = 3,750 \ ppm \times 1 = 3,750 \ ppm$$

Uncertainty Factor: The LOAEL_{HEC} was divided by a total uncertainty factor of 300:

- 10 for use of a LOAEL
- 3 for extrapolation from animals to humans after dosimetric adjustment
- 10 for human variability

$$MRL = LOAEL_{HEC} \div UFs$$

 $3,750~ppm~\div 300~= 12.5~ppm~\approx 13~ppm$

Other Additional Studies or Pertinent Information that Lend Support to this MRL: No supporting studies or pertinent information were available.

Chemical Name:ChloroethaneCAS Numbers:75-00-3Date:January 2025

Profile Status:FinalRoute:InhalationDuration:Chronic

MRL Summary: There are insufficient data for derivation of a chronic-duration inhalation MRL for chloroethane.

Rationale for Not Deriving an MRL: Renal and neurological effects were observed in female mice exposed to 15,000 ppm chloroethane for 100 weeks (only concentration tested) (NTP 1989). However, this study also reported decreased survival in these exposed mice, attributed to carcinomas of the uterus. An MRL for renal and/or neurological effects was not derived because serious health effects were seen at the concentration level studied.

Chemical Name: Chloroethane CAS Numbers: 75-00-3
Date: January 2025

Profile Status:FinalRoute:OralDuration:Acute

MRL Summary: The database was not considered adequate for derivation of an acute-duration oral MRL for chloroethane.

Rationale for Not Deriving an MRL: No acute-duration oral MRL was derived due to lack of adequate data regarding the potential effects of chloroethane. No toxicologically relevant effects were seen in rats exposed to chloroethane in drinking water at doses ranging from 297 to 662 mg/kg/day for 7 or 14 days (Dow 1995).

Chemical Name: Chloroethane CAS Numbers: 75-00-3
Date: January 2025

Profile Status: Final **Route:** Oral

Duration: Intermediate

MRL Summary: The database was not considered adequate for derivation of an intermediate-duration oral MRL for chloroethane.

Rationale for Not Deriving an MRL: No intermediate-duration oral studies were located that met ATSDR quality inclusion criteria; therefore, no MRL could be derived.

Chemical Name: Chloroethane CAS Numbers: 75-00-3
Date: January 2025

Profile Status:FinalRoute:OralDuration:Chronic

MRL Summary: The database was not considered adequate for derivation of a chronic-duration oral MRL for chloroethane.

Rationale for Not Deriving an MRL: No studies were located that describe the effects of chronic-duration oral exposure to chloroethane in humans or animals.

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APPENDIX B. LITERATURE SEARCH FRAMEWORK FOR CHLOROETHANE

The objective of the toxicological profile is to evaluate the potential for human exposure and the potential health hazards associated with inhalation, oral, or dermal/ocular exposure to chloroethane.

B.1 LITERATURE SEARCH AND SCREEN

A literature search and screen were conducted to identify studies examining health effects, toxicokinetics, mechanisms of action, susceptible populations, biomarkers, chemical interactions, physical and chemical properties, production, use, environmental fate, environmental releases, and environmental and biological monitoring data for chloroethane. ATSDR primarily focused on peer-reviewed articles without language restrictions. Foreign language studies are reviewed based on available English-language abstracts and/or tables (or summaries in regulatory assessments, such as International Agency for Research on Cancer [IARC] documents). If the study appears critical for hazard identification or MRL derivation, translation into English is requested. Non-peer-reviewed studies that were considered relevant to the assessment of the health effects of chloroethane have undergone peer review by at least three ATSDR-selected experts who have been screened for conflict of interest. The inclusion criteria used to identify relevant studies examining the health effects of chloroethane are presented in Table B-1.

Table B-1. Inclusion Criteria for the Literature Search and Screen

Health Effects

Species

Human

Laboratory mammals

Route of exposure

Inhalation

Oral

Dermal (or ocular)

Parenteral (these studies will be considered supporting data)

Health outcome

Death

Systemic effects

Body weight effects

Respiratory effects

Cardiovascular effects

Gastrointestinal effects

Hematological effects

Musculoskeletal effects

Hepatic effects

Renal effects

Dermal effects

Ocular effects

Endocrine effects

Immunological effects

Neurological effects

Reproductive effects

Table B-1. Inclusion Criteria for the Literature Search and Screen

Developmental effects

Other noncancer effects

Cancer

Toxicokinetics

Absorption

Distribution

Metabolism

Excretion

PBPK models

Biomarkers

Biomarkers of exposure

Biomarkers of effect

Interactions with other chemicals

Potential for human exposure

Releases to the environment

Air

Water

Soil

Environmental fate

Transport and partitioning

Transformation and degradation

Environmental monitoring

Air

Water

Sediment and soil

Other media

Biomonitoring

General populations

Occupation populations

B.1.1 Literature Search

The literature search was conducted to update the Toxicological Profile for Chloroethane released in 1998. All literature cited in the previous (1998) toxicological profile were considered for inclusion in the updated profile. The initial literature search, which was performed in May 2022, was restricted to studies added to databases since January 1996. An updated literature search was performed after the Toxicological Profile for Chloroethane Draft for Public Comment was released in January 2024 to identify any additional studies added to databases between May 2022 and April 2024.

The following main databases were searched in May 2022 and April 2024:

- PubMed
- National Technical Reports Library (NTRL)
- Scientific and Technical Information Network's TOXCENTER

The search strategy used the chemical names, Chemical Abstracts Service (CAS) numbers, synonyms, Medical Subject Headings (MeSH) headings, and keywords for chloroethane. The query strings used for the literature search are presented in Table B-2.

The search was augmented by searching the Toxic Substances Control Act Test Submissions (TSCATS), NTP website, and National Institute of Health Research Portfolio Online Reporting Tools Expenditures and Results (NIH RePORTER) databases using the queries presented in Table B-3. Additional databases were searched in the creation of various tables and figures, such as the TRI Explorer, the Substance Priority List (SPL) resource page, and other items as needed. Regulations applicable to chloroethane were identified by searching international and U.S. agency websites and documents.

Review articles were identified and used for the purpose of providing background information and identifying additional references. ATSDR also identified reports from the grey literature, which included unpublished research reports, technical reports from government agencies, conference proceedings and abstracts, and theses and dissertations.

	Table B-2. Database Query Strings
Database search date	e Query string
PubMed	
04/2024 05/2022	(("Ethyl Chloride"[mh] OR 75-00-3[rn]) AND (2022/05/01:3000[mhda] OR 2021:3000[dp])) OR (("Chloroethane"[tw] OR "Ethane, chloro-"[tw] OR "Ethyl chloride"[tw] OR "ethylchloride"[tw] OR "Freon 160"[tw] OR "Monochlorethane"[tw] OR "Anodynon"[tw] OR "Monochloroethane"[tw] OR "Chlorethane"[tw] OR "Chloryle anesthetic"[tw] OR "Chlorethan"[tw] OR "Chloryle anesthetic"[tw] OR "Cloretilo"[tw] OR "Dublofix"[tw] OR "Ether chloratus"[tw] OR "Ether chloridum"[tw] OR "Ether hydrochloric"[tw] OR "Ether muriatic"[tw] OR "Hydrochloric ether"[tw] OR "Kelene"[tw] OR "Muriatic ether"[tw] OR "Narcotile"[tw] OR "Chelen"[tw] OR "Chlorene"[tw] OR "Chloridum"[tw] OR "Chloryl"[tw] OR "Chlorethyl"[tw]) AND (2022/05/01:3000[edat] OR 2022/05/01:3000[crdt] OR 2021:3000[dp])) ("Ethyl Chloride"[mh] OR 75-00-3[rn] OR "Chloroethane"[tw] OR "Ethane, chloro-"[tw] OR
00/2022	"Ethyl chloride"[tw] OR "ethylchloride"[tw] OR "Freon 160"[tw] OR "Monochlorethane"[tw] OR "Monochloroethane"[tw] OR "Chlorethane"[tw] OR "Chlorethane"[tw] OR "Chloryl anesthetic"[tw] OR "Chloryle anesthetic"[tw] OR "Cloretilo"[tw] OR "Dublofix"[tw] OR "Ether chloratus"[tw] OR "Ether chloridum"[tw] OR "Ether hydrochloric"[tw] OR "Ether muriatic"[tw] OR "Hydrochloric ether"[tw] OR "Kelene"[tw] OR "Muriatic ether"[tw] OR "Narcotile"[tw] OR "Chelen"[tw] OR "Chlorene"[tw] OR "Chloridum"[tw] OR "Chloryl"[tw] OR "Chlorethyl"[tw]) AND 1996:3000[dp]
NTRL	
04/2024	"Chloroethane" OR "Ethane, chloro-" OR "Ethyl chloride" OR "ethylchloride" OR "Freon 160" OR "Monochlorethane" OR "Monochloroethane" OR "chlorethane" OR "Aethylis" OR "Anodynon" OR "Chlorethan" OR "Chloryl anesthetic" OR "Chloryle anesthetic" OR "Cloretilo" OR "Dublofix" OR "Ether chloratus" OR "Ether chloridum" OR "Ether hydrochloric" OR "Ether muriatic" OR "Hydrochloric ether" OR "Kelene" OR "Muriatic ether" OR "Narcotile" OR "Chelen" OR "Chlorene" OR "Chloridum" OR "Chloryl" OR "Chlorethyl" Limited to 2021-2024
05/2022	"Chloroethane" OR "Ethane, chloro-" OR "Ethyl chloride" OR "ethylchloride" OR "Freon 160" OR "Monochlorethane" OR "Monochloroethane" OR "chlorethane" OR "Aethylis" OR

APPENDIX B

Table B-2. Database Query Strings

Database

search date Query string

"Anodynon" OR "Chlorethan" OR "Chloryl anesthetic" OR "Chloryle anesthetic" OR "Cloretilo" OR "Dublofix" OR "Ether chloratus" OR "Ether chloridum" OR "Ether hydrochloric" OR "Ether muriatic" OR "Hydrochloric ether" OR "Kelene" OR "Muriatic ether" OR "Narcotile" OR "Chelen" OR "Chlorene" OR "Chloridum" OR "Chloryl" OR "Chlorethyl"

Limited to 1996-2022

Toxcenter

04/2024

FILE 'TOXCENTER' ENTERED AT 15:38:29 ON 25 APR 2024

CHARGED TO COST=ET027.02.04.LB.02

- L1 2397 SEA FILE=TOXCENTER 75-00-3
- L2 2037 SEA FILE=TOXCENTER L1 NOT PATENT/DT
- L3 128 SEA FILE=TOXCENTER L2 AND ED>=20220501

ACT TOXQUERY/Q

- L4 QUE (CHRONIC OR IMMUNOTOX? OR NEUROTOX? OR TOXICOKIN? OR BIOMARKER? OR NEUROLOG?)
- L5 QUE (PHARMACOKIN? OR SUBCHRONIC OR PBPK OR EPIDEMIOLOGY/ST,CT,

IT)

- L6 QUE (ACUTE OR SUBACUTE OR LD50# OR LD(W)50 OR LC50# OR LC(W)50)
- L7 QUE (TOXICITY OR ADVERSE OR POISONING)/ST,CT,IT
- L8 QUE (INHAL? OR PULMON? OR NASAL? OR LÚNG? OR RESPIR?)
- L9 QUE ((OCCUPATION? OR WORKPLACE? OR WORKER?) AND EXPOS?)
- L10 QUE (ORAL OR ORALLY OR INGEST? OR GAVAGE? OR DIET OR DIETS

OR

DIETARY OR DRINKING(W)WATER?)

- L11 QUE (MAXIMUM AND CONCENTRATION? AND (ALLOWABLE OR PERMISSIBLE))
- L12 QUE (ABORT? OR ABNORMALIT? OR EMBRYO? OR CLEFT? OR FETUS?)
- L13 QUE (FOETUS? OR FETAL? OR FOETAL? OR FERTIL? OR MALFORM?

OR

OVUM?)

- L14 QUE (OVA OR OVARY OR PLACENTA? OR PREGNAN? OR PRENATAL?)
- L15 QUE (PERINATAL? OR POSTNATAL? OR REPRODUC? OR STERIL? OR TERATOGEN?)
- L16 QUE (SPERM OR SPERMAC? OR SPERMAG? OR SPERMATI? OR SPERMAS? OR

SPERMATOB? OR SPERMATOC? OR SPERMATOG?)

L17 QUE (SPERMATOI? OR SPERMATOL? OR SPERMATOR? OR SPERMATOX? OR

SPERMATOZ? OR SPERMATU? OR SPERMI? OR SPERMO?)

L18 QUE (NEONAT? OR NEWBORN? OR DEVELOPMENT OR

DEVELOPMENTAL?)

- L19 QUE (ENDOCRIN? AND DISRUPT?)
- L20 QUE (ZYGOTE? OR CHILD OR CHILDREN OR ADOLESCEN? OR

INFANT?)

- L21 QUE (WEAN? OR OFFSPRING OR AGE(W)FACTOR?)
- L22 QUE (DERMAL? OR DERMIS OR SKIN OR EPIDERM? OR CUTANEOUS?)

APPENDIX B

Table B-2.	Database	Query Strings
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Database		
search date	Query string	
	L23 QUE (CAF	RCINOG? OR COCARCINOG? OR CANCER? OR PRECANCER?
	OR `	
	NEOPLAS?	
		IOR? OR TUMOUR? OR ONCOGEN? OR LYMPHOMA? OR
	CARCINOM?)	
		IETOX? OR GENOTOX? OR MUTAGEN? OR
	GENETIC(W)TOXIC?)	
		PHROTOX? OR HEPATOTOX?) OCRIN? OR ESTROGEN? OR ANDROGEN? OR HORMON?)
	•	CUPATION? OR WORKER? OR WORKPLACE? OR EPIDEM?)
		R L5 OR L6 OR L7 OR L8 OR L9 OR L10 OR L11 OR L12 OR
		OR L15 OR L16 OR L17 OR L18 OR L19 OR L20 OR L21 OR
		OR L24 OR L25 OR L26 OR L27 OR L28
	L30 QUE (RAT	OR RATS OR MOUSE OR MICE OR GUINEA(W)PIG? OR
	MURIDAE	
		R DOGS OR RABBIT? OR HAMSTER? OR PIG OR PIGS OR
	SWINE	IE OD MONIKEVO OD MA OA ONEOV
		IE OR MONKEY? OR MACAQUE?)
	L31 QUE (MAF LAGOMORPHA	RMOSET? OR FERRET? OR GERBIL? OR RODENT? OR
		N? OR CANINE OR CAT OR CATS OR FELINE OR MURINE)
		OR L30 OR L31
		NHUMAN MAMMALS)/ORGN
	L34 QUE L32 (
		MAN OR HUMANS OR HOMINIDAE OR MAMMALS OR MAMMAL?
	OR	
		OR PRIMATE?)
	L36 QUE L34 (JR L35
	L37 57 SEA FILE	E=TOXCENTER L3 AND L36
		=TOXCENTER L37 AND MEDLINE/FS
		E=TOXCENTER L37 NOT MEDLINE/FS
		M L38 L39 (0 DUPLICATES REMOVED)
	L41 71 SEA FILE	E=TOXCENTER L3 NOT L37
	L42 1 SEA FILE	=TOXCENTER L3 AND MEDLINE/FS
	D SCAN L3	
05/2022		L' ENTERED AT 16:55:39 ON 16 MAY 2022
	CHARGED TO COST:	
		E=TOXCENTER 75-00-3
		E=TOXCENTER L1 NOT TSCATS/FS E=TOXCENTER L2 NOT PATENT/DT
		E=TOXCENTER L2 NOT PATENT/DT E=TOXCENTER L3 AND PY>1995
		OXQUERY/Q
		ongoenna
	L5 QUE (CHR	ONIC OR IMMUNOTOX? OR NEUROTOX? OR TOXICOKIN? OR
		R? OR NEUROLOG?)
		RMACOKIN? OR SUBCHRONIC OR PBPK OR
	EPIDEMIOLOGY/ST,C	CT,
	IT)	TE OD CUDACUTE OD I DE0# OD I D/A/\\co\ OD I OF0# OD
	L7 QUE (ACU	TE OR SUBACUTE OR LD50# OR LD(W)50 OR LC50# OR

APPENDIX B

Table B-2. Database Query Strings

Database		
search date	Query str	ing
		LC(W)50)
	L8	QUE (TOXICITY OR ADVERSE OR POISONING)/ST,CT,IT
	L9	QUE (INHAL? OR PULMON? OR NASAL? OR LÚNG? OR RESPIR?)
	L10	QUE ((OCCUPATION? OR WORKPLACE? OR WORKER?) AND EXPOS?)
	L11	QUE (ORAL OR ORALLY OR INGEST? OR GAVAGE? OR DIET OR DIETS
	OR	
		DIETARY OR DRINKING(W)WATER?)
	L12	QUE (MAXIMUM AND CONCENTRATION? AND (ALLOWABLE OR
	PERMISS	IBLE))
	L13	QUE (ABORT? OR ABNORMALIT? OR EMBRYO? OR CLEFT? OR FETUS?)
	L14	QUE (FOETUS? OR FETAL? OR FOETAL? OR FERTIL? OR MALFORM?
	OR	QUE (1 DE 100. OTT ETTE. OTT DE 17.E. OTT ETTE. OTT MILE OTT
		OVUM?)
	L15	QUE (OVA OR OVARY OR PLACENTA? OR PREGNAN? OR PRENATAL?)
	L16	QUE (PERINATAL? OR POSTNATAL? OR REPRODUC? OR STERIL? OR
	•	TERATÒGEN?)
	L17	QUE (SPERM OR SPERMAC? OR SPERMAG? OR SPERMATI? OR
	SPERMAS	3? OR
	;	SPERMATOB? OR SPERMATOC? OR SPERMATOG?)
	L18	QUE (SPERMATOI? OR SPERMATOL? OR SPERMATOR? OR
	SPERMA1	
		SPERMATOZ? OR SPERMATU? OR SPERMI? OR SPERMO?)
	L19	QUE (NEONAT? OR NEWBORN? OR DEVELOPMENT OR
		PMENTAL?)
	L20 L21	QUE (ENDOCRIN? AND DISRUPT?) QUE (ZYGOTE? OR CHILD OR CHILDREN OR ADOLESCEN? OR
	INFANT?)	
	L22	QUE (WEAN? OR OFFSPRING OR AGE(W)FACTOR?)
	L23	QUE (DERMAL? OR DERMIS OR SKIN OR EPIDERM? OR CUTANEOUS?)
	L24	QUE (CARCINOG? OR COCARCINOG? OR CANCER? OR PRECANCER?
	OR	QUE (OF INTO INTO E. OTT OF INTO ETT. OTT THE OF INTO ETT.
		NEOPLAS?)
	L25	QUE (TUMOR? OR TUMOUR? OR ONCOGEN? OR LYMPHOMA? OR
	CARCINO	OM?)
	L26	QUE (GENETOX? OR GENOTOX? OR MUTAGEN? OR
		(W)TOXIC?)
	L27	QUE (NEPHROTOX? OR HEPATOTOX?)
	L28	QUE (ENDOCRIN? OR ESTROGEN? OR ANDROGEN? OR HORMON?)
	L29	QUE (OCCUPATION? OR WORKER? OR WORKPLACE? OR EPIDEM?)
	L30	QUE L5 OR L6 OR L7 OR L8 OR L9 OR L10 OR L11 OR L12 OR L13 OR
		L14 OR L15 OR L16 OR L17 OR L18 OR L19 OR L20 OR L21 OR L22 OR
		L23 OR L24 OR L25 OR L26 OR L27 OR L28 OR L29
	L31	QUE (RAT OR RATS OR MOUSE OR MICE OR GUINEA(W)PIG? OR
	MURIDAE	
	SWINE	OR DOG OR DOGS OR RABBIT? OR HAMSTER? OR PIG OR PIGS OR
		OR PORCINE OR MONKEY? OR MACAQUE?)
	L32	QUE (MARMOSET? OR FERRET? OR GERBIL? OR RODENT? OR
	LAGOMOI	

Table B-2. Database Query Strings **Database** search date Query string OR BABOON? OR CANINE OR CAT OR CATS OR FELINE OR MURINE) L33 QUE L30 OR L31 OR L32 L34 QUE (NONHUMAN MAMMALS)/ORGN L35 **QUE L33 OR L34** L36 QUE (HUMAN OR HUMANS OR HOMINIDAE OR MAMMALS OR MAMMAL? OR PRIMATES OR PRIMATE?) L37 **QUE L35 OR L36** 414 SEA FILE=TOXCENTER L4 AND L37 L38 L39 52 SEA FILE=TOXCENTER L38 AND MEDLINE/FS L41 362 SEA FILE=TOXCENTER L38 NOT MEDLINE/FS L42 393 DUP REM L39 L41 (21 DUPLICATES REMOVED) L*** DEL 52 S L38 AND MEDLINE/FS L*** DEL 52 S L38 AND MEDLINE/FS 52 SEA FILE=TOXCENTER L42 L*** DEL 362 S L38 NOT MEDLINE/FS L*** DEL 362 S L38 NOT MEDLINE/FS L44 341 SEA FILE=TOXCENTER L42

341 SEA FILE=TOXCENTER (L43 OR L44) NOT MEDLINE/FS

L45

D SCAN L45

Table B-3. Strategies to Augment the Literature Search Source Query and number screened when available TSCATS via ChemView 04/2024; 05/2022 Compounds searched: 75-00-3 NTP 04/2024 75-00-3 chloroethane "ethyl chloride" chlorethane chloryl "Ethane, chloro-" ethylchloride "Freon 160" monochloroethane 05/2022 75-00-3 chloroethane "ethyl chloride" Regulations.gov 04/2024 Limited to 2021 to present chloroethane 75-00-3 "ethyl chloride" 05/2022 chloroethane

7	Table B-3. Strategies to Augment the Literature Search
Source	Query and number screened when available
	75-00-3 "ethyl chloride" Limited to EPA docket or notices
NIH RePORTER	
08/2024	Search Criteria Fiscal Year: Active Projects; Text Search: "Chloroethane" OR "Ethane, chloro-" OR "Ethyl chloride" OR "ethylchloride" OR "Freon 160" OR "Monochlorethane" OR "Monochloroethane" OR "chlorethane" OR "Aethylis" OR "Anodynon" OR "Chlorethan" OR "Chloryl anesthetic" OR "Chloryle anesthetic" OR "Cloretilo" OR "Dublofix" OR "Ether chloratus" OR "Ether chloridum" OR "Ether hydrochloric" OR "Ether muriatic" OR "Hydrochloric ether" OR "Kelene" OR "Muriatic ether" OR "Narcotile" OR "Chelen" OR "Chlorene" OR "Chloridum" OR "Chloryl" OR "Chlorethyl" (advanced) Limit to: Project Title, Project Terms, Project Abstracts
03/2023	Search Criteria Fiscal Year: Active Projects Text Search: "Chloroethane" OR "Ethane, chloro-" OR "Ethyl chloride" OR "ethylchloride" OR "Freon 160" OR "Monochlorethane" OR "Monochloroethane" OR "chlorethane" OR "Aethylis" OR "Anodynon" OR "Chlorethan" OR "Chloryl anesthetic" OR "Chloryle anesthetic" OR "Cloretilo" OR "Dublofix" OR "Ether chloratus" OR "Ether chloridum" OR "Ether hydrochloric" OR "Ether muriatic" OR "Hydrochloric ether" OR "Kelene" OR "Muriatic ether" OR "Narcotile" OR "Chelen" OR "Chlorene" OR "Chloryl" OR "Chlorethyl" (advanced)Limit to: Project Title, Project Terms, Project Abstracts Search Criteria Fiscal Year: Active Projects Text Search: "Chloridum" (advanced)Limit to: Project Title, Project Terms, Project Abstracts
Other	Includes additional reference identified throughout the assessment process, which may include studies found by tree searching; recommended by intraagency, interagency, peer, or public reviewers; or published more recently than the date of literature search(es). Additional references include those for specific regulations or guidelines and publications found by targeted searches for specific information (e.g., searches for reviews of general [not chemical-specific] mechanisms of toxicity).

The 2022 pre-public comment search results were:

- Number of records identified from PubMed, NTRL, and TOXCENTER (after duplicate removal): 688
- Number of records identified from other strategies: 74
- Total number of records to undergo literature screening: 762

The 2024 post-public comment search results were:

- Number of records identified from PubMed, NTRL, and TOXCENTER (after duplicate removal): 184
- Number of records identified from other strategies: 17
- Total number of records to undergo literature screening: 201

B.1.2 Literature Screening

A two-step process was used to screen the literature search to identify relevant studies on chloroethane during the pre- and post-public comment drafts:

- Title and abstract screen
- Full text screen

Pre-Public Comment Title and Abstract Screen. Within the reference library, titles and abstracts were screened manually for relevance. Studies that were considered (see Table B-1 for inclusion criteria) were moved to the second step of the literature screening process. Studies were excluded when the title and abstract clearly indicated that the study was not relevant to the toxicological profile.

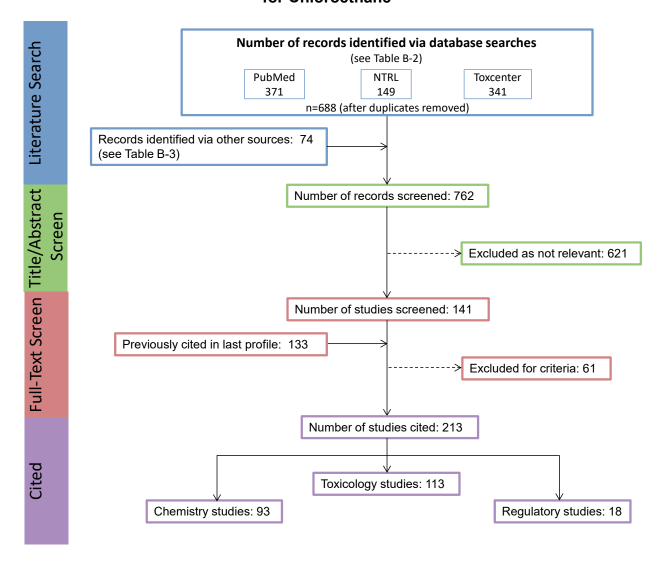
- Number of titles and abstracts screened: 762
- Number of studies considered relevant and moved to the next step: 141

Pre-Public Comment Full Text Screen. The second step in the literature screening process was a full text review of individual studies considered relevant in the title and abstract screen step. Each study was reviewed to determine whether it was relevant for inclusion in the toxicological profile.

- Number of studies undergoing full text review: 141
- Number of studies cited in the previous toxicological profile: 133
- Total number of studies cited in the profile: 213

A summary of the results of the pre-public literature search and screening is presented in Figure B-1.

Figure B-1. May 2022 Pre-Public Comment Literature Search Results and Screen for Chloroethane*



^{*}The chemistry studies category includes studies pertaining to the potential for human exposure (Table B-1). The toxicology studies category includes human and animal studies of health effects as well as studies of toxicokinetics, biomarkers, and interactions with other chemicals (Table B-1). The regulatory studies category includes those studies cited in Chapter 7.

Post-Public Comment Title and Abstract Screen. Within the reference library, titles and abstracts were screened manually for relevance. Studies that were considered relevant (see Table B-1 for inclusion criteria) were moved to the second step of the literature screening process. Studies were excluded when the title and abstract clearly indicated that the study was not relevant to the toxicological profile.

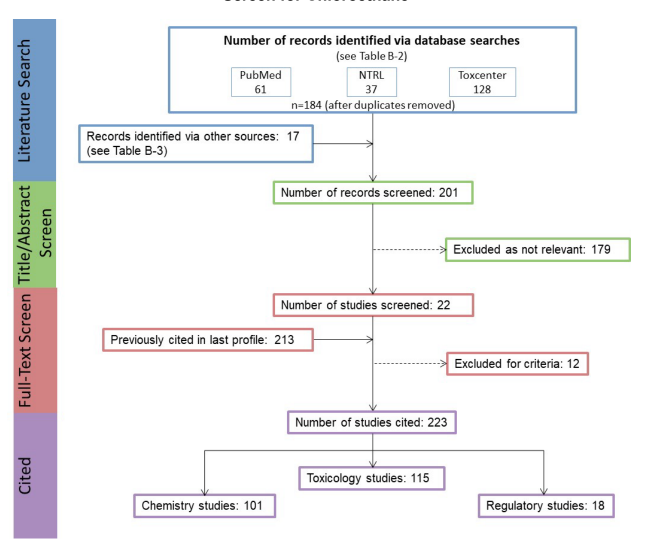
- Number of titles and abstracts screened: 201
- Number of studies considered relevant and moved to the next step: 22

Post-Public Comment Full Text Screen. The second step in the literature screening process was a full text review of individual studies considered relevant in the title and abstract screen step. Each study was reviewed to determine whether it was relevant for inclusion in the toxicological profile.

- Number of studies undergoing full text review: 22
- Number of studies cited in the pre-public draft of the toxicological profile: 213
- Total number of studies cited in the profile: 223

A summary of the results of the post-public comment literature search and screening is presented in Figure B-2.

Figure B-2. April 2024 Post-Public Comment Literature Search Results and Screen for Chloroethane*



^{*}The chemistry studies category includes studies pertaining to the potential for human exposure (Table B-1). The toxicology studies category includes human and animal studies of health effects as well as studies of toxicokinetics, biomarkers, and interactions with other chemicals (Table B-1). The regulatory studies category includes those studies cited in Chapter 7.

CHLOROETHANE C-1

APPENDIX C. FRAMEWORK FOR ATSDR'S SYSTEMATIC REVIEW OF HEALTH EFFECTS DATA FOR CHLOROETHANE

To increase the transparency of ATSDR's process of identifying, evaluating, synthesizing, and interpreting the scientific evidence on the health effects associated with exposure to chloroethane, ATSDR utilized a slight modification of NTP's Office of Health Assessment and Translation (OHAT) systematic review methodology (NTP 2013, 2015; Rooney et al. 2014). ATSDR's framework is an eight-step process for systematic review with the goal of identifying the potential health hazards of exposure to chloroethane:

- Step 1. Problem Formulation
- Step 2. Literature Search and Screen for Health Effects Studies
- Step 3. Extract Data from Health Effects Studies
- Step 4. Identify Potential Health Effect Outcomes of Concern
- Step 5. Assess the Risk of Bias for Individual Studies
- Step 6. Rate the Confidence in the Body of Evidence for Each Relevant Outcome
- Step 7. Translate Confidence Rating into Level of Evidence of Health Effects
- Step 8. Integrate Evidence to Develop Hazard Identification Conclusions

C.1 PROBLEM FORMULATION

The objective of the toxicological profile and this systematic review was to identify the potential health hazards associated with inhalation, oral, or dermal/ocular exposure to chloroethane. The inclusion criteria used to identify relevant studies examining the health effects of chloroethane are presented in Table C-1.

Data from human and laboratory animal studies were considered relevant for addressing this objective. Human studies were divided into two broad categories: observational epidemiology studies and controlled exposure studies. The observational epidemiology studies were further divided: cohort studies (retrospective and prospective studies), population studies (with individual data or aggregate data), and case-control studies.

Table C-1. Inclusion Criteria for Identifying Health Effects Studies

Species

Human

Laboratory mammals

Route of exposure

Inhalation

Oral

Dermal (or ocular)

Parenteral (these studies will be considered supporting data)

Health outcome

Death

Systemic effects

Body weight effects

Respiratory effects

Cardiovascular effects

Table C-1. Inclusion Criteria for Identifying Health Effects Studies

Gastrointestinal effects

Hematological effects

Musculoskeletal effects

Hepatic effects

Renal effects

Dermal effects

Ocular effects

Endocrine effects

Immunological effects

Neurological effects

Reproductive effects

Developmental effects

Other noncancer effects

Cancer

C.2 LITERATURE SEARCH AND SCREEN FOR HEALTH EFFECTS STUDIES

A literature search and screen were conducted to identify studies examining the health effects of chloroethane. The literature search framework for the toxicological profile is discussed in detail in Appendix B.

C.2.1 Literature Search

As noted in Appendix B, the literature searches were intended to update the 1998 Toxicological Profile for Chloroethane. See Appendix B for the databases searched and the search strategy.

A total of 761 and 22 records relevant to all sections of the toxicological profile were identified in the initial and update literature search, respectively.

C.2.2 Literature Screening

As described in Appendix B, a two-step process was used to screen the literature search to identify relevant studies examining the health effects of chloroethane.

Title and Abstract Screen. In the Title and Abstract Screen step, 62 documents (inclusive of both literature searches) were considered to meet the health effects inclusion criteria in Table C-1 and were moved to the next step in the process.

Full Text Screen. In the second step in the literature screening process for the systematic review, a full text review of 62 health effect documents (documents identified in the update literature search and documents cited in older versions of the profile) was performed. From those 62 documents (79 studies), 20 documents (33 studies) were included in the qualitative review.

C.3 EXTRACT DATA FROM HEALTH EFFECTS STUDIES

Relevant data extracted from the individual studies selected for inclusion in the systematic review were collected in customized data forms. A summary of the type of data extracted from each study is presented in Table C-2. For references that included more than one experiment or species, data extraction records were created for each experiment or species.

Table C-2. Data Extracted from Individual Studies

Citation

Chemical form

Route of exposure (e.g., inhalation, oral, dermal)

Specific route (e.g., gavage in oil, drinking water)

Species

Strain

Exposure duration category (e.g., acute, intermediate, chronic)

Exposure duration

Frequency of exposure (e.g., 6 hours/day, 5 days/week)

Exposure length

Number of animals or subjects per sex per group

Dose/exposure levels

Parameters monitored

Description of the study design and method

Summary of calculations used to estimate doses (if applicable)

Summary of the study results

Reviewer's comments on the study

Outcome summary (one entry for each examined outcome)

No-observed-adverse-effect level (NOAEL) value

Lowest-observed-adverse-effect level (LOAEL) value

Effect observed at the LOAEL value

A summary of the extracted data for each study is presented in the Supplemental Document for Chloroethane and overviews of the results of the inhalation, oral, and dermal exposure studies are presented in Sections 2.2–2.18 of the profile and in the Levels Significant Exposures tables in Section 2.1 of the profile (Tables 2-1 and 2-2, respectively).

C.4 IDENTIFY POTENTIAL HEALTH EFFECT OUTCOMES OF CONCERN

Overviews of the potential health effect outcomes for chloroethane identified in human and animal studies are presented in Tables C-3 and C-4, respectively. Human inhalation studies examined a limited number of health outcomes, whereas animal inhalation and oral studies examined a comprehensive set of endpoints. Dermal exposure studies in humans were primarily interested in analgesic effects of chloroethane (discussed in Section 2.11). Both human and animal studies suggest the nervous system is the primary target of chloroethane exposure. Additionally, animal studies suggest the reproductive and developmental effects may also be sensitive targets. Although there are several case reports evaluating neurological effects in humans following inhalation, these studies were not included in this systematic review due to either a lack of estimated exposure or a comparison group (discussed in Section 2.15). The

remaining human (inhalation) and animal (inhalation and oral) studies related to neurological, reproductive, and developmental outcomes were carried through to Steps 4–8 of the systematic review. There were 33 studies (published in 20 documents) examining these potential outcomes carried through to Steps 4–8 of the systematic review.

Table C-3. C	Overv	/iew c	of the	Healt	h Out	tcome	es for	Chlo	oroeth	ane E	valuat	ted in	Huma	an St	udies		
	Body weight	Respiratory	Cardiovascular	Gastrointestinal	Hematological	Musculoskeletal	Hepatic	Renal	Dermal	Ocular	Endocrine	Immunological	Neurological	Reproductive	Developmental	Other Noncancer	Caner
Inhalation studies																	
Cohort																	
Case control																	
Population												1	1 1				
Case series		3	2	2 2	1		2 2						10 10				
Human controlled			1 1					'		1			3				
Oral studies																	
Cohort																	
Case control																	
Population																	
Case series																	
Dermal studies																	
Cohort																	
Case control																	
Population																	
Case series									3	1		3					
Human controlled									18 0				1				
Number of studies examining Number of studies reporting of				0	1	2 2	3	4	5–9 5–9	≥10 ≥10							

Table C-4. Overv	iew of	the H	ealth	Outco	omes	for C	hloroe	ethane	Eval	uated	in Ex	perim	ental	Anim	al S	tudie	S
	Body weight	Respiratory	Cardiovascular	Gastrointestinal	Hematological	Musculoskeletal	Hepatic	Renal	Dermal	Ocular	Endocrine	Immunological ^a	Neurological ^a	Reproductivea	Developmental	Other Noncancer	Caner
Inhalation studies	44	44	40			0	10	40		0	7	7	4.4	40	_		
Acute-duration	11	11	10 4	6 1	4	3	12 2	10	5 0	3	0	7	14 8	10	2		
	5	4	2	2		U	5	4	2	1	5	4	2	3			
Intermediate-duration	0	0	0	0			0	0	0	0	0	0	0	1			
Chronic-duration	2	2	2	2		2	2	2	2		2	2	2	2			2 2
Oral studies	0	0	0	0		0	0	1	0		0	0	1	0			
	2		1		1		1	1			1	1	3	1			
Acute-duration	0		0		0		0	0			0	0	1	0			
Intermediate-duration	0																
Chronic-duration																	
Dermal studies																	
Acute-duration						1			1				1				
Intermediate-duration																	
Chronic-duration																	
Number of studies examini Number of studies reporting				0 0	1	2 2	3	4	5–9 5–9	≥10 ≥10							

^aNumber of studies examining endpoint includes study evaluating histopathology, but not evaluating function.

C.5 ASSESS THE RISK OF BIAS FOR INDIVIDUAL STUDIES

C.5.1 Risk of Bias Assessment

The risk of bias of individual studies was assessed using OHAT's Risk of Bias Tool (NTP 2015). The risk of bias questions for observational epidemiology studies, human-controlled exposure studies, and animal experimental studies are presented in Tables C-5, C-6, and C-7, respectively. Each risk of bias question was answered on a four-point scale:

- Definitely low risk of bias (++)
- Probably low risk of bias (+)
- Probably high risk of bias (-)
- Definitely high risk of bias (--)

In general, "definitely low risk of bias" or "definitely high risk of bias" were used if the question could be answered with information explicitly stated in the study report. If the response to the question could be inferred, then "probably low risk of bias" or "probably high risk of bias" responses were typically used.

Table C-5. Risk of Bias Questionnaire for Observational Epidemiology Studies

Selection bias

Were the comparison groups appropriate?

Confounding bias

Did the study design or analysis account for important confounding and modifying variables?

Attrition/exclusion bias

Were outcome data complete without attrition or exclusion from analysis?

Detection bias

Is there confidence in the exposure characterization?

Is there confidence in outcome assessment?

Selective reporting bias

Were all measured outcomes reported?

Table C-6. Risk of Bias Questionnaire for Human-Controlled Exposure Studies

Selection bias

Was administered dose or exposure level adequately randomized?

Was the allocation to study groups adequately concealed?

Performance bias

Were the research personnel and human subjects blinded to the study group during the study?

Attrition/exclusion bias

Were outcome data complete without attrition or exclusion from analysis?

Detection bias

Is there confidence in the exposure characterization?

Is there confidence in outcome assessment?

Selective reporting bias

Were all measured outcomes reported?

Table C-7. Risk of Bias Questionnaire for Experimental Animal Studies

Selection bias

Was administered dose or exposure level adequately randomized?

Was the allocation to study groups adequately concealed?

Performance bias

Were experimental conditions identical across study groups?

Were the research personnel blinded to the study group during the study?

Attrition/exclusion bias

Were outcome data complete without attrition or exclusion from analysis?

Detection bias

Is there confidence in the exposure characterization?

Is there confidence in outcome assessment?

Selective reporting bias

Were all measured outcomes reported?

After the risk of bias questionnaires were completed for the health effects studies, the studies were assigned to one of three risk of bias tiers based on the responses to the key questions listed below and the responses to the remaining questions.

- Is there confidence in the exposure characterization? (only relevant for observational studies)
- Is there confidence in the outcome assessment?
- Does the study design or analysis account for important confounding and modifying variables? (only relevant for observational studies)

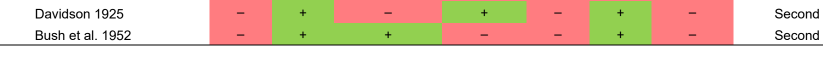
First Tier. Studies placed in the first tier received ratings of "definitely low" or "probably low" risk of bias on the key questions **AND** received a rating of "definitely low" or "probably low" risk of bias on the responses to at least 50% of the other applicable questions.

Second Tier. A study was placed in the second tier if it did not meet the criteria for the first or third tiers.

Third Tier. Studies placed in the third tier received ratings of "definitely high" or "probably high" risk of bias for the key questions **AND** received a rating of "definitely high" or "probably high" risk of bias on the response to at least 50% of the other applicable questions.

The results of the risk of bias assessment for the different types of chloroethane health effects studies (human-controlled exposure and animal experimental studies) are presented in Tables C-8 and C-9, respectively.

	•	Risk of bias criteria and ratings									
	Selecti	Attrition/ Performance exclusion Selection bias bias Detection bias				on bias	Selective reporting bias				
Reference	Was administered dose or exposure level adequately randomized?	Was the allocation to study groups adequately concealed?	Were the research personnel and human subjects blinded to the study group during the study?	Were outcome data complete without attrition or exclusion from analysis?	Is there confidence in the exposure characterization?	Is there confidence in the outcome assessment?*	Were all measured outcomes reported?	Risk of bias tier			
Itcome: Neurological effects Inhalation acute exposure											
USBM 1929	_	+	-	-	-	-	_	Third			
D :1 4005											



+++ = definitely low risk of bias; + = probably low risk of bias; - = probably high risk of bias; - = definitely high risk of bias

^{*}Key question used to assign risk of bias tier.

Table C-9. Summary of Risk of Bias Assessment for Chloroethane—Experimental Animal Studies

	Risk of bias criteria and ratings											
					Attrition/ exclusion			Selective reporting bias	_			
	Selection	n bias	Perform	ance bias	bias	Detecti	Detection bias					
Reference	Was administered dose or exposure level adequately randomized?	Was the allocation to study groups adequately concealed?	Were experimental conditions identical across study groups?	Were the research personnel blinded to the study group during the study?	Were outcome data complete without attrition or exclusion from analysis?	Is the confidence in the exposure characterization?	Is there confidence in the outcome assessment?*	Were all measured outcomes reported?	Risk of bias tier			

Outcome: Neurological Effects

Inhalation acute exposure									_
Bush et al. 1952	-	+	-	+	-		-	-	Third
Dow 1941 (monkey)	_	+	-	+	+	-	+	+	Second
Dow 1992 (mouse)	++	+	++	+	+	++	+	+	First
Dow 1992 (rat, ¹⁴ C-chloroethane)	++	+	++	+	+	++	+	+	First
Dow 1992 (mouse, ¹⁴ C-chloroethane)	++	+	++	+	+	++	+	+	First
Dow 1985	++	+	+	+	+	-	++	+	First
Gohlke and Schmidt 1972; Schmidt et al. 1972	-	+	-	+	-	-	+	+	Second
Landry et al. 1982 (rat)	++	+	+	+	+	++	+	+	First
Landry et al. 1982 (dog)	++	+	+	+	+	++	+	+	First
Landry et al. 1987, 1989	++	+	+	+	++	++	+	++	First
Lazarew 1929	-	+	-	+	-	-	+	+	Second
NTP 1989 (mouse, 2 weeks)	++	+	+	+	+	+	-	+	Second
NTP 1989 (rat, 2 weeks)	++	+	+	+	+	+	-	+	Second
Morris et al. 1953	-	+	-	+	+		+	-	Second
USBM 1929 (810 minutes)	_	+	-	+	++	-	+	+	First

		Risk of bias criteria and ratings							
			Attrition/ Selection report						
	Selection bias		Performance bias		bias	Detection bias		bias	
Reference	Was administered dose or exposure level adequately randomized?	Was the allocation to study groups adequately concealed?	Were experimental conditions identical across study groups?	Were the research personnel blinded to the study group during the study?	Were outcome data complete without attrition or exclusion from analysis?	Is the confidence in the exposure characterization?	Is there confidence in the outcome assessment?*	Were all measured outcomes reported?	Risk of bias tier
Inhalation intermediate exposure									
NTP 1989 (rat, 13 weeks)	++	+	+	+	++	+	+	+	First
NTP 1989 (mouse, 13 weeks) Inhalation chronic exposure	++	+	+	+	++	+	+	+	First
NTP 1989 (rat, 102 weeks)	++	+	+	+	++	+	+	+	First
NTP 1989 (mouse, 100 weeks)	++	+	+	+	++	+	+	+	First
Oral acute exposure									
Dow 1992 (mouse non-labelled)	++	+	-	+	+	++	+	+	First
Dow 1992 (rat, ¹⁴ C-chloroethane)	++	+	_	+	+	++	+	+	First
Dow 1992 (mouse, ¹⁴ C-chloroethane)	++	+	-	+	+	++	+	+	First
Dow 1995	++	+	+	+	++	++	+	++	First
utcome: Reproductive Effects									
Inhalation acute exposure					<u> </u>				
Breslin et al. 1988	++	+	+	+	+	+	+	+	First
Fedtke et al. 1994a, 1994b (rat)	-	+	-	+	+	+	+	-	First
Fedtke et al. 1994a, 1994b (mouse)	-	+	_	+	+	+	+	-	First

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Table C-9. Summary	of Risk o	f Bias Ass	sessment	for Chloro	ethane—	Experime	ental Anir	nal Studies	5
	Risk of bias criteria and ratings								
	Salaati	on bigg	Attrition/ exclusion Performance bias bias Detection bias					Selective reporting bias	-
	Selection bias		•		bias	Detection bias		bias	1
Reference	Was administered dose or exposure level adequately randomized?	Was the allocation to study groups adequately concealed?	Were experimental conditions identical across study groups?	Were the research personnel blinded to the study group during the study?	Were outcome data complete without attrition or exclusion from analysis?	Is the confidence in the exposure characterization?	Is there confidence in the outcome assessment?*	Were all measured outcomes reported?	Risk of bias tier
Gohlke and Schmidt 1972; Schmidt et al. 1972	-	+	-	+	-	-	+	+	Second
Landry et al. 1982 (rat)	++	+	+	+	+	++	+	+	First
Landry et al. 1982 (dog)	++	+	+	+	+	++	+	+	First
Landry et al. 1987, 1989	++	+	+	+	++	++	+	++	First
NTP 1989 (mouse, 2 weeks)	++	+	+	+	+	+	-	+	Second
NTP 1989 (rat, 2 weeks)	++	+	+	+	+	+	-	+	Second
van Liere et al. 1966	_	+	-	+	-		+	-	Second
Inhalation intermediate exposure									
Bucher et al. 1995	+	+	+	+	++	+	+	+	First
NTP 1989 (rat, 13 weeks)	++	+	+	+	++	+	+	+	First
NTP 1989 (mouse, 13 weeks)	++	+	+	+	++	+	+	+	First
Inhalation chronic exposure									
NTP 1989 (rat, 102 weeks)	++	+	+	+	++	+	+	++	First
NTP 1989 (mouse, 100 weeks)	++	+	+	+	++	+	+	++	First
Oral acute exposure									
Dow 1995	++	+	+	+	++	++	+	++	First

Table C-9. Summary of Risk of Bias Assessment for Chloroethane—Experimental Animal Studies Risk of bias criteria and ratings Attrition/ Selective exclusion reporting Selection bias Performance bias bias **Detection bias** bias personnel blinded to the Was administered dose s the confidence in the adequately concealed? study group during the s there confidence in across study groups? Was the allocation to attrition or exclusion outcomes reported? Were outcome data Were experimental conditions identical Were all measured Were the research or exposure level characterization? complete without Risk of bias tier rom analysis? assessment?* study groups randomized? the outcome adequately exposure study? Reference **Outcome: Developmental Effects** Inhalation acute exposure First Dow 1985 First ++ Scortichini et al. 1986 ++

^{🚻 =} definitely low risk of bias; 🛨 = probably low risk of bias; 🗕 = probably high risk of bias; 🗕 = definitely high risk of bias

^{*}Key question used to assign risk of bias tier.

C.6 RATE THE CONFIDENCE IN THE BODY OF EVIDENCE FOR EACH RELEVANT OUTCOME

Confidences in the bodies of human and animal evidence were evaluated independently for each potential outcome. ATSDR did not evaluate the confidence in the body of evidence for carcinogenicity; rather, the Agency defaulted to the cancer weight-of-evidence assessment of other agencies including HHS, EPA, and IARC. The confidence in the body of evidence for an association or no association between exposure to chloroethane and a particular outcome was based on the strengths and weaknesses of individual studies. Four descriptors were used to describe the confidence in the body of evidence for effects or when no effect was found:

- **High confidence:** the true effect is highly likely to be reflected in the apparent relationship
- Moderate confidence: the true effect may be reflected in the apparent relationship
- Low confidence: the true effect may be different from the apparent relationship
- **Very low confidence:** the true effect is highly likely to be different from the apparent relationship

Confidence in the body of evidence for a particular outcome was rated for each type of study: case-control, case series, cohort, population, human-controlled exposure, and experimental animal. In the absence of data to the contrary, data for a particular outcome were collapsed across animal species, routes of exposure, and exposure durations. If species (or strain), route, or exposure duration differences were noted, then the data were treated as separate outcomes.

C.6.1 Initial Confidence Rating

In ATSDR's modification to the OHAT approach, the body of evidence for an association (or no association) between exposure to chloroethane and a particular outcome was given an initial confidence rating based on the key features of the individual studies examining that outcome. The presence of these key features of study design was determined for individual studies using four "yes or no" questions, which were customized for epidemiology, human controlled exposure, or experimental animal study designs. Separate questionnaires were completed for each outcome assessed in a study. The key features for observational epidemiology (cohort, population, and case-control) studies, human controlled exposure, and experimental animal studies are presented in Tables C-10, C-11, and C-12, respectively. The initial confidence in the study was determined based on the number of key features present in the study design:

- **High Initial Confidence:** Studies in which the responses to the four questions were "yes".
- **Moderate Initial Confidence:** Studies in which the responses to only three of the questions were "yes".
- Low Initial Confidence: Studies in which the responses to only two of the questions were "yes".
- Very Low Initial Confidence: Studies in which the response to one or none of the questions was "yes".

Table C-10. Key Features of Study Design for Observational Epidemiology Studies

Exposure was experimentally controlled

Exposure occurred prior to the outcome

Outcome was assessed on individual level rather than at the population level

A comparison group was used

Table C-11. Key Features of Study Design for Human-Controlled Exposure Studies

A comparison group was used or the subjects served as their own control

A sufficient number of subjects were tested

Appropriate methods were used to measure outcomes (i.e., clinically-confirmed outcome versus self-reported)

Appropriate statistical analyses were performed and reported or the data were reported in such a way to allow independent statistical analysis

Table C-12. Key Features of Study Design for Experimental Animal Studies

A concurrent control group was used

A sufficient number of animals per group were tested

Appropriate parameters were used to assess a potential adverse effect

Appropriate statistical analyses were performed and reported or the data were reported in such a way to allow independent statistical analysis

The presence or absence of the key features and the initial confidence levels for studies examining neurological and reproductive outcomes observed in the human-controlled exposure and animal experimental studies are presented in Tables C-13 and C-14, respectively.

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APPENDIX C

Table C-13. Presence of Key Features of Study Design for Chloroethane— Human-Controlled Exposure

			Key Features		
Reference	Comparison group or served as own controls	Sufficient number of subjects tested	Appropriate outcome assessment	Appropriate statistical analysis	Initial study confidence

Outcome: Neurological Effects

Inhalation acute exposure

USBM 1929 No No No No Very Low Davidson 1925 No Yes Very Low No No Bush et al. 1952 Yes Low No Yes No

Table C-14.	Presence of Key Features of Study Design for Chloroethane—
	Experimental Animal Studies

		Key fe	atures		_
Reference	Concurrent control group	Sufficient number of animals per group	Appropriate parameters to assess potential effect	Adequate data for statistical analysis	Initial study confidence

Outcome: Neurological Effects

Inhalation acute exposure

Bush et al. 1952

Dow 1941 (monkey)

Dow 1992 (mouse)

Dow 1992 (rat, ¹⁴C-chloroethane)

Dow 1992 (mouse, ¹⁴C-chloroethane)

Dow 1985 (10 days)

Gohlke and Schmidt 1972; Schmidt et

al. 1972

Landry et al. 1982 (rat)

Landry et al. 1982 (dog)

No	Yes	Yes	No	Low
No	No	No	No	Low
Yes	No	Yes	No	Low
No	Yes	Yes	Yes	Moderate
No	Yes	Yes	No	Low
Yes	Yes	Yes	No	Moderate
Yes	Yes	Yes	No	Moderate
Yes	Yes	Yes	No	Moderate
Yes	Yes	Yes	No	Moderate

Table C-14. Presence of Key Features of Study Design for Chloroethane—

Experimental Animal Studies					
		Key fea	atures		.
Reference	Concurrent control group	Sufficient number of animals per group	Appropriate parameters to assess potential effect	Adequate data for statistical analysis	Initial study confidence
Landry et al. 1987, 1989	Yes	Yes	Yes	Yes	High
Lazarew 1929	No	No	Yes	No	Very Low
NTP 1989 (mouse, 2 weeks)	Yes	Yes	Yes	Yes	High
NTP 1989 (rat, 2 weeks)	Yes	Yes	Yes	Yes	High
Morris et al. 1953	No	Yes	Yes	No	Low
USBM 1929 (540 minutes)	Yes	Yes	Yes	No	Moderate
Inhalation intermediate exposure					
NTP 1989 (rat, 13 weeks)	Yes	Yes	Yes	Yes	High
NTP 1989 (mouse, 13 weeks)	Yes	Yes	Yes	Yes	High
Inhalation chronic exposure					
NTP 1989 (rat, 102 weeks)	Yes	Yes	Yes	No	Moderate
NTP 1989 (mouse, 100 weeks)	Yes	Yes	Yes	No	Moderate
Oral acute exposure					
Dow 1992 (mouse)	No	No	Yes	Yes	Low
Dow 1992 (rat, ¹⁴ C-chloroethane)	No	Yes	Yes	Yes	Moderate
Dow 1992 (mouse, ¹⁴ C-chloroethane)	No	Yes	Yes	Yes	Moderate
Dow 1995	Yes	Yes	Yes	Yes	High
Outcome: Reproductive Effects					
Inhalation acute exposure	V	V	V	V	Littada
Breslin et al. 1988	Yes	Yes	Yes	Yes	High
Fedtke et al. 1994a (rat)	Yes Yes	Yes Yes	No No	No No	Low
Fedtke et al. 1994a (mouse) Gohlke and Schmidt 1972; Schmidt et	res	res	INO	INO	Low
al. 1972	Yes	Yes	Yes	No	Moderate
Landry et al. 1982 (rat)	Yes	Yes	Yes	No	Moderate
Landry et al. 1982 (dog)	Yes	Yes	Yes	No	Moderate
Landry et al. 1987, 1989	Yes	Yes	Yes	Yes	High
NTP 1989 (mouse, 2 weeks)	Yes	Yes	Yes	Yes	High
NTP 1989 (rat, 2 weeks)	Yes	Yes	Yes	Yes	High
van Liere et al. 1966	No	Yes	Yes	No	Low

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Table C-14. Presence of Key Features of Study Design for Chloroethane— **Experimental Animal Studies** Key features Appropriate parameters Sufficient number of Concurrent control o assess potential animals per group statistical analysis Adequate data for Initial study Reference confidence Inhalation intermediate exposure Bucher et al. 1995 Yes Yes Yes Yes High NTP 1989 (rat, 13 weeks) Yes Yes Yes Yes High NTP 1989 (mouse, 13 weeks) Yes Yes Yes Yes High Inhalation chronic exposure NTP 1989 (rat, 102 weeks) Yes Yes Yes Yes High NTP 1989 (mouse, 100 weeks) Yes Yes Yes Yes High Oral acute exposure Dow 1995 Yes Yes No Yes Moderate Outcome: Developmental Effects Inhalation acute exposure Moderate Dow 1985 Yes Yes Yes No Scortichini et al. 1986 Yes Yes Yes Yes High

A summary of the initial confidence ratings for each outcome is presented in Table C-15. If individual studies for a particular outcome and study type had different study quality ratings, then the highest confidence rating for the group of studies was used to determine the initial confidence rating for the body of evidence; any exceptions were noted in Table C-15.

Table C-15. Initial Confidence Rating for Chloroethane Health Effects Studies			
	Initial study confidence	Initial confidence rating	
Outcome: Neurological Effects			
Inhalation acute exposure			
Human studies			
USBM 1929	Very Low		
Davidson 1925	Very Low	Low	
Bush et al. 1952	Low		
Animal studies			
Bush et al. 1952	Low		
Dow 1941 (monkey)	Low		

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Table C-15. Initial Confidence Rating for Chloroethane Health Effects Studies

	Initial study confidence	Initial confidence rating	
Dow 1992 (mouse)	Low		
Dow 1992 (rat, ¹⁴ C-chloroethane)	Moderate		
Dow 1992 (mouse, ¹⁴ C-chloroethane)	Low		
Dow 1985 (10 days)	Moderate		
Gohlke and Schmidt 1972; Schmidt et al. 1972	Moderate		
Landry et al. 1982 (rat)	Moderate	1.126	
Landry et al. 1982 (dog)	Moderate	High	
Landry et al. 1987, 1989	High		
Lazarew 1929	Very Low		
NTP 1989 (mouse, 2 weeks)	High		
NTP 1989 (rat, 2 weeks)	High		
Morris et al. 1953	Low		
USBM 1929 (540 minutes)	Moderate		
Inhalation intermediate exposure			
Animal studies			
NTP 1989 (rat, 13 weeks)	High	High	
NTP 1989 (mouse, 13 weeks)	High	піgп	
Inhalation chronic exposure			
Animal studies			
NTP 1989 (rat, 102 weeks)	Moderate	Moderate	
NTP 1989 (mouse, 100 weeks)	Moderate	Moderate	
Oral acute exposure			
Animal studies			
Dow 1992 (mouse)	Low		
Dow 1992 (rat, ¹⁴ C-chloroethane)	Moderate	High	
Dow 1992 (mouse, ¹⁴ C-chloroethane)	Moderate	High	
Dow 1995	High		

Inhalation acute exposure

Animal studies

Breslin et al. 1988

Fedtke et al. 1994a, 1994b (rat)

Fedtke et al. 1994a, 1994b (mouse)

Gohlke and Schmidt 1972; Schmidt et al. 1972

Landry et al. 1982 (rat)

Landry et al. 1982 (dog)

Landry et al. 1987, 1989

NTP 1989 (mouse, 2 weeks

NTP 1989 (rat, 2 weeks)

van Liere et al. 1966

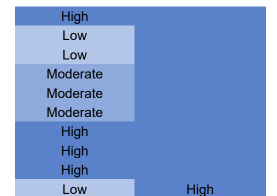


Table C-15. Initial Confidence Rating for Chloroethane Health Effects Studies			
	Initial study confidence	Initial confidence rating	
Inhalation intermediate exposure			
Animal studies			
Bucher et al. 1995	High		
NTP 1989 (rat, 13 weeks)	High	High	
NTP 1989 (mouse, 13 weeks)	High		
Inhalation chronic exposure			
Animal studies			
NTP 1989 (rat, 102 weeks)	High	High	
NTP 1989 (mouse, 100 weeks)	High	High	
Oral acute exposure			
Animal studies			
Dow 1995	Moderate	Moderate	
Outcome: Developmental Effects			
Inhalation acute exposure			
Animal studies			
Dow 1985	Moderate		
Scortichini et al. 1986	High	High	

C.6.2 Adjustment of the Confidence Rating

The initial confidence rating was then downgraded or upgraded depending on whether there were substantial issues that would decrease or increase confidence in the body of evidence. The nine properties of the body of evidence that were considered are listed below. The summaries of the assessment of the confidence in the body of evidence for neurological and reproductive effects are presented in Table C-16. If the confidence ratings for a particular outcome were based on more than one type of human study, then the highest confidence rating was used for subsequent analyses. An overview of the confidence in the body of evidence for all health effects associated with chloroethane exposure is presented in Table C-17.

Table C-16. Adjustments to the Initial Confidence in the Body of Evidence				
	Initial confidence	Adjustments to the initial confidence rating	Final confidence	
Outcome: Neurologic	al Effects			
Human studies	Low	-1 risk of bias +1 consistency	Low	
Animal studies	High	+1 consistency -1 indirectness	High	
Outcome: Reproducti	ve Effects			
Animal studies	High	-1 inconsistency-1 indirectness	Low	

Table C-16. A	djustments to the Init	ial Confidence in the Body	y of Evidence
	Initial confidence	Adjustments to the initial confidence rating	Final confidence
Outcome: Developme	ental Effects		
Animal studies	High	-1 inconsistency-1 indirectness-1 imprecision	Low

Table C-17. Confidence in the Body of Evidence for Chloroethane Confidence in body of evidence Human studies Animal studies Neurological effects Low High Reproductive effects No data Low Developmental effects No data Low

Five properties of the body of evidence were considered to determine whether the confidence rating should be downgraded:

- **Risk of bias.** Evaluation of whether there is substantial risk of bias across most of the studies examining the outcome. This evaluation used the risk of bias tier groupings for individual studies examining a particular outcome (Tables C-8 and C-9). Below are the criteria used to determine whether the initial confidence in the body of evidence for each outcome should be downgraded for risk of bias:
 - o No downgrade if most studies are in the risk of bias first tier
 - Downgrade one confidence level if most studies are in the risk of bias second tier
 - o Downgrade two confidence levels if most studies are in the risk of bias third tier
- Unexplained inconsistency. Evaluation of whether there is inconsistency or large variability in the magnitude or direction of estimates of effect across studies that cannot be explained. Below are the criteria used to determine whether the initial confidence in the body of evidence for each outcome should be downgraded for unexplained inconsistency:
 - No downgrade if there is little inconsistency across studies or if only one study evaluated the outcome
 - Downgrade one confidence level if there is variability across studies in the magnitude or direction of the effect
 - Downgrade two confidence levels if there is substantial variability across studies in the magnitude or direct of the effect
- **Indirectness.** Evaluation of four factors that can affect the applicability, generalizability, and relevance of the studies:
 - Relevance of the animal model to human health—unless otherwise indicated, studies in rats, mice, and other mammalian species are considered relevant to humans
 - O Directness of the endpoints to the primary health outcome—examples of secondary outcomes or nonspecific outcomes include organ weight in the absence of histopathology or clinical chemistry findings in the absence of target tissue effects

- Nature of the exposure in human studies and route of administration in animal studies—inhalation, oral, and dermal exposure routes are considered relevant unless there are compelling data to the contrary
- Duration of treatment in animal studies and length of time between exposure and outcome assessment in animal and prospective human studies—this should be considered on an outcome-specific basis

Below are the criteria used to determine whether the initial confidence in the body of evidence for each outcome should be downgraded for indirectness:

- No downgrade if none of the factors are considered indirect
- o Downgrade one confidence level if one of the factors is considered indirect
- O Downgrade two confidence levels if two or more of the factors are considered indirect
- Imprecision. Evaluation of the narrowness of the effect size estimates and whether the studies have adequate statistical power. Data are considered imprecise when the ratio of the upper to lower 95% Cis for most studies is ≥10 for tests of ratio measures (e.g., odds ratios) and ≥100 for absolute measures (e.g., percent control response). Adequate statistical power is determined if the study can detect a potentially biologically meaningful difference between groups (20% change from control response for categorical data or risk ratio of 1.5 for continuous data). Below are the criteria used to determine whether the initial confidence in the body of evidence for each outcome should be downgraded for imprecision:
 - o No downgrade if there are no serious imprecisions
 - o Downgrade one confidence level for serious imprecisions
 - o Downgrade two confidence levels for very serious imprecisions
- **Publication bias.** Evaluation of the concern that studies with statistically significant results are more likely to be published than studies without statistically significant results.
 - Downgrade one level of confidence for cases where there is serious concern with publication bias

Four properties of the body of evidence were considered to determine whether the confidence rating should be upgraded:

- Large magnitude of effect. Evaluation of whether the magnitude of effect is sufficiently large so that it is unlikely to have occurred as a result of bias from potential confounding factors.
 - O Upgrade one confidence level if there is evidence of a large magnitude of effect in a few studies, provided that the studies have an overall low risk of bias and there is no serious unexplained inconsistency among the studies of similar dose or exposure levels; confidence can also be upgraded if there is one study examining the outcome, provided that the study has an overall low risk of bias
- **Dose response.** Evaluation of the dose-response relationships measured within a study and across studies. Below are the criteria used to determine whether the initial confidence in the body of evidence for each outcome should be upgraded:
 - Upgrade one confidence level for evidence of a monotonic dose-response gradient
 - Upgrade one confidence level for evidence of a non-monotonic dose-response gradient where there is prior knowledge that supports a non-monotonic dose-response and a nonmonotonic dose-response gradient is observed across studies
- Plausible confounding or other residual biases. This factor primarily applies to human studies and is an evaluation of unmeasured determinants of an outcome such as residual bias towards the

null (e.g., "healthy worker" effect) or residual bias suggesting a spurious effect (e.g., recall bias). Below is the criterion used to determine whether the initial confidence in the body of evidence for each outcome should be upgraded:

- Upgrade one confidence level for evidence that residual confounding or bias would underestimate an apparent association or treatment effect (i.e., bias toward the null) or suggest a spurious effect when results suggest no effect
- Consistency in the body of evidence. Evaluation of consistency across animal models and species, consistency across independent studies of different human populations and exposure scenarios, and consistency across human study types. Below is the criterion used to determine whether the initial confidence in the body of evidence for each outcome should be upgraded:
 - o Upgrade one confidence level if there is a high degree of consistency in the database

C.7 TRANSLATE CONFIDENCE RATING INTO LEVEL OF EVIDENCE OF HEALTH EFFECTS

In the seventh step of the systematic review of the health effects data for chloroethane, the confidence in the body of evidence for specific outcomes was translated to a level of evidence rating. The level of evidence rating reflected the confidence in the body of evidence and the direction of the effect (i.e., toxicity or no toxicity); route-specific differences were noted. The level of evidence for health effects was rated on a five-point scale:

- **High level of evidence:** High confidence in the body of evidence for an association between exposure to the substance and the health outcome
- **Moderate level of evidence:** Moderate confidence in the body of evidence for an association between exposure to the substance and the health outcome
- Low level of evidence: Low confidence in the body of evidence for an association between exposure to the substance and the health outcome
- Evidence of no health effect: High confidence in the body of evidence that exposure to the substance is not associated with the health outcome
- **Inadequate evidence:** Low or moderate confidence in the body of evidence that exposure to the substance is not associated with the health outcome OR very low confidence in the body of evidence for an association between exposure to the substance and the health outcome

A summary of the level of evidence of health effects for chloroethane is presented in Table C-18.

Table C-18. Level of Evidence of Health Effects for Chloroethane				
Outcome	Confidence in body of evidence	Direction of health effect	Level of evidence for health effect	
Human studies				
Neurological	Low	Health effect	Low	
Animal studies				
Neurological	High	Health effect	High	
Reproductive	Low	Health effect	Low	
Developmental	Low	Health effect	Low	

C.8 INTEGRATE EVIDENCE TO DEVELOP HAZARD IDENTIFICATION CONCLUSIONS

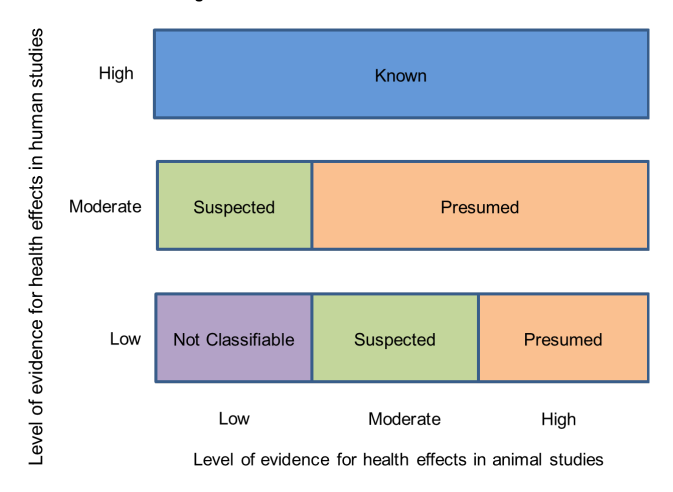
The final step involved the integration of the evidence streams for the human studies and animal studies to allow for a determination of hazard identification conclusions. For health effects, there were four hazard identification conclusion categories:

- **Known** to be a hazard to humans
- **Presumed** to be a hazard to humans
- **Suspected** to be a hazard to humans
- **Not classifiable** as to the hazard to humans

The initial hazard identification was based on the highest level of evidence in the human studies and the level of evidence in the animal studies; if there were no data for one evidence stream (human or animal), then the hazard identification was based on the one data stream (equivalent to treating the missing evidence stream as having low level of evidence). The hazard identification scheme is presented in Figure C-1 and described below:

- **Known:** A health effect in this category would have:
 - o High level of evidence for health effects in human studies **AND** a high, moderate, or low level of evidence in animal studies.
- **Presumed:** A health effect in this category would have:
 - Moderate level of evidence in human studies AND high or moderate level of evidence in animal studies OR
 - o Low level of evidence in human studies AND high level of evidence in animal studies
- **Suspected:** A health effect in this category would have:
 - Moderate level of evidence in human studies AND low level of evidence in animal studies OR
 - Low level of evidence in human studies AND moderate level of evidence in animal studies
- Not classifiable: A health effect in this category would have:
 - o Low level of evidence in human studies AND low level of evidence in animal studies

Figure C-1. Hazard Identification Scheme



Other relevant data such as mechanistic or mode-of-action data were considered to raise or lower the level of the hazard identification conclusion by providing information that supported or opposed biological plausibility.

Two hazard identification conclusion categories were used when the data indicated that there may be no health effect in humans:

- **Not identified** to be a hazard in humans
- **Inadequate** to determine hazard to humans

If the human level of evidence conclusion of no health effect was supported by the animal evidence of no health effect, then the hazard identification conclusion category of "not identified" was used. If the human or animal level of evidence was considered inadequate, then a hazard identification conclusion category of "inadequate" was used. As with the hazard identification for health effects, the impact of other relevant data was also considered for no health effect data.

The hazard identification conclusions for chloroethane are listed below and summarized in Table C-19.

Presumed Health Effects

- Neurological
 - Low level of evidence from human studies: several case reports described neurological symptoms after inhaling chloroethane; however, exposure levels are not known (Al-Ajmi et al. 2018; Demarest et al. 2011; Finch and Lobo 2005; Hager et al. 2021; Hes et al. 1979; Kuthiah and Er 2019; Nordin et al. 1988; Senussi and Chalise 2015). Volunteers who inhaled chloroethane reported feeling dizzy and slightly intoxicated and had increased reaction times (Davidson 1925; USBM 1929); however, these studies were poor quality with high risk of bias.
 - High level of evidence from animal studies: neurological effects have been reported in several species following inhalation (Bush et al. 1952; Dow 1985, 1992, 1995; Landry et al. 1982; Lazarew 1929; Morris et al. 1953; NTP 1989; USBM 1929;) and gavage (Dow 1992) exposure.

Not Classifiable Health Effects

- Reproductive
 - Low level of evidence from animal studies: inhalation studies have reported effects on the length of the estrous cycle (Bucher et al. 1995), uterine weight (Fedtke et al. 1994a), and uterine glutathione levels (Fedtke et al. 1994b), although no histopathological and/or hormonal changes have been reported after exposure (Bucher et al. 1995; Landry et al. 1987, 1989; NTP 1989). Breslin et al. (1988) reported that inhalation did not affect the estrous cycle of mice.
- Developmental
 - Low level of confidence from animal studies: one inhalation study reported increased incidence of delayed fetal foramina closure (DFFC) of the skull bones in pups exposed *in utero* on GDs 6–15 (Scortichini et al. 1986), whereas another study of pups exposed *in utero* on GDs 6–15 did not report any fetal abnormalities (Dow 1985).

Table C-19. Hazard Identification Conclusions for Chloroethane			
Outcome	Hazard identification		
Neurological	Presumed		
Reproductive	Not classifiable		
Developmental	Not classifiable		

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APPENDIX D. USER'S GUIDE

Chapter 1. Relevance to Public Health

This chapter provides an overview of U.S. exposures, a summary of health effects based on evaluations of existing toxicologic, epidemiologic, and toxicokinetic information, and an overview of the minimal risk levels. This is designed to present interpretive, weight-of-evidence discussions for human health endpoints by addressing the following questions:

- 1. What effects are known to occur in humans?
- 2. What effects observed in animals are likely to be of concern to humans?
- 3. What exposure conditions are likely to be of concern to humans, especially around hazardous waste sites?

Minimal Risk Levels (MRLs)

Where sufficient toxicologic information is available, ATSDR derives MRLs for inhalation and oral routes of entry at each duration of exposure (acute, intermediate, and chronic). These MRLs are not meant to support regulatory action, but to acquaint health professionals with exposure levels at which adverse health effects are not expected to occur in humans.

MRLs should help physicians and public health officials determine the safety of a community living near a hazardous substance emission, given the concentration of a contaminant in air or the estimated daily dose in water. MRLs are based largely on toxicological studies in animals and on reports of human occupational exposure.

MRL users should be familiar with the toxicologic information on which the number is based. Section 1.2, Summary of Health Effects, contains basic information known about the substance. Other sections, such as Section 3.2 Children and Other Populations that are Unusually Susceptible and Section 3.4 Interactions with Other Substances, provide important supplemental information.

MRL users should also understand the MRL derivation methodology. MRLs are derived using a modified version of the risk assessment methodology that the Environmental Protection Agency (EPA) provides (Barnes and Dourson 1988) to determine reference doses (RfDs) for lifetime exposure.

To derive an MRL, ATSDR generally selects the most sensitive endpoint which, in its best judgement, represents the most sensitive human health effect for a given exposure route and duration. ATSDR cannot make this judgement or derive an MRL unless information (quantitative or qualitative) is available for all potential systemic, neurological, and developmental effects. If this information and reliable quantitative data on the chosen endpoint are available, ATSDR derives an MRL using the most sensitive species (when information from multiple species is available) with the highest no-observed-adverse-effect level (NOAEL) that does not exceed any adverse effect levels. When a NOAEL is not available, a lowest-observed-adverse-effect level (LOAEL) can be used to derive an MRL, and an uncertainty factor of 10 must be employed. Additional uncertainty factors of 10 must be used both for human variability to protect sensitive subpopulations (people who are most susceptible to the health effects caused by the substance) and for interspecies variability (extrapolation from animals to humans). In deriving an MRL, these individual uncertainty factors are multiplied together. The product is then divided into the inhalation concentration or oral dosage selected from the study. Uncertainty factors used in developing a

substance-specific MRL are provided in the footnotes of the levels of significant exposure (LSE) tables that are provided in Chapter 2. Detailed discussions of the MRLs are presented in Appendix A.

Chapter 2. Health Effects

Tables and Figures for Levels of Significant Exposure (LSE)

Tables and figures are used to summarize health effects and illustrate graphically levels of exposure associated with those effects. These levels cover health effects observed at increasing dose concentrations and durations, differences in response by species and MRLs to humans for noncancer endpoints. The LSE tables and figures can be used for a quick review of the health effects and to locate data for a specific exposure scenario. The LSE tables and figures should always be used in conjunction with the text. All entries in these tables and figures represent studies that provide reliable, quantitative estimates of NOAELs, LOAELs, or Cancer Effect Levels (CELs).

The legends presented below demonstrate the application of these tables and figures. Representative examples of LSE tables and figures follow. The numbers in the left column of the legends correspond to the numbers in the example table and figure.

TABLE LEGEND

See Sample LSE Table (page D-5)

- (1) Route of exposure. One of the first considerations when reviewing the toxicity of a substance using these tables and figures should be the relevant and appropriate route of exposure.

 Typically, when sufficient data exist, three LSE tables and two LSE figures are presented in the document. The three LSE tables present data on the three principal routes of exposure (i.e., inhalation, oral, and dermal). LSE figures are limited to the inhalation and oral routes. Not all substances will have data on each route of exposure and will not, therefore, have all five of the tables and figures. Profiles with more than one chemical may have more LSE tables and figures.
- (2) Exposure period. Three exposure periods—acute (<15 days), intermediate (15–364 days), and chronic (≥365 days)—are presented within each relevant route of exposure. In this example, two oral studies of chronic-duration exposure are reported. For quick reference to health effects occurring from a known length of exposure, locate the applicable exposure period within the LSE table and figure.
- (3) <u>Figure key</u>. Each key number in the LSE table links study information to one or more data points using the same key number in the corresponding LSE figure. In this example, the study represented by key number 51 identified NOAELs and less serious LOAELs (also see the three "51R" data points in sample LSE Figure 2-X).
- (4) Species (strain) No./group. The test species (and strain), whether animal or human, are identified in this column. The column also contains information on the number of subjects and sex per group. Chapter 1, Relevance to Public Health, covers the relevance of animal data to human toxicity and Section 3.1, Toxicokinetics, contains any available information on comparative toxicokinetics. Although NOAELs and LOAELs are species specific, the levels are extrapolated to equivalent human doses to derive an MRL.
- (5) <u>Exposure parameters/doses</u>. The duration of the study and exposure regimens are provided in these columns. This permits comparison of NOAELs and LOAELs from different studies. In this case (key number 51), rats were orally exposed to "Chemical X" via feed for 2 years. For a

- more complete review of the dosing regimen, refer to the appropriate sections of the text or the original reference paper (i.e., Aida et al. 1992).
- (6) Parameters monitored. This column lists the parameters used to assess health effects. Parameters monitored could include serum (blood) chemistry (BC), biochemical changes (BI), body weight (BW), clinical signs (CS), developmental toxicity (DX), food intake (FI), gross necropsy (GN), hematology (HE), histopathology (HP), immune function (IX), lethality (LE), neurological function (NX), organ function (OF), ophthalmology (OP), organ weight (OW), reproductive function (RX), urinalysis (UR), and water intake (WI).
- (7) Endpoint. This column lists the endpoint examined. The major categories of health endpoints included in LSE tables and figures are death, body weight, respiratory, cardiovascular, gastrointestinal, hematological, musculoskeletal, hepatic, renal, dermal, ocular, endocrine, immunological, neurological, reproductive, developmental, other noncancer, and cancer. "Other noncancer" refers to any effect (e.g., alterations in blood glucose levels) not covered in these systems. In the example of key number 51, three endpoints (body weight, hematological, and hepatic) were investigated.
- (8) <u>NOAEL</u>. A NOAEL is the highest exposure level at which no adverse effects were seen in the organ system studied. The body weight effect reported in key number 51 is a NOAEL at 25.5 mg/kg/day. NOAELs are not reported for cancer and death; with the exception of these two endpoints, this field is left blank if no NOAEL was identified in the study.
- (9) LOAEL. A LOAEL is the lowest dose used in the study that caused an adverse health effect. LOAELs have been classified into "Less Serious" and "Serious" effects. These distinctions help readers identify the levels of exposure at which adverse health effects first appear and the gradation of effects with increasing dose. A brief description of the specific endpoint used to quantify the adverse effect accompanies the LOAEL. Key number 51 reports a less serious LOAEL of 6.1 mg/kg/day for the hepatic system, which was used to derive a chronic exposure, oral MRL of 0.008 mg/kg/day (see footnote "c"). MRLs are not derived from serious LOAELs. A cancer effect level (CEL) is the lowest exposure level associated with the onset of carcinogenesis in experimental or epidemiologic studies. CELs are always considered serious effects. The LSE tables and figures do not contain NOAELs for cancer, but the text may report doses not causing measurable cancer increases. If no LOAEL/CEL values were identified in the study, this field is left blank.
- (10) <u>Reference</u>. The complete reference citation is provided in Chapter 8 of the profile.
- (11) <u>Footnotes</u>. Explanations of abbreviations or reference notes for data in the LSE tables are found in the footnotes. For example, footnote "c" indicates that the LOAEL of 6.1 mg/kg/day in key number 51 was used to derive an oral MRL of 0.008 mg/kg/day.

FIGURE LEGEND

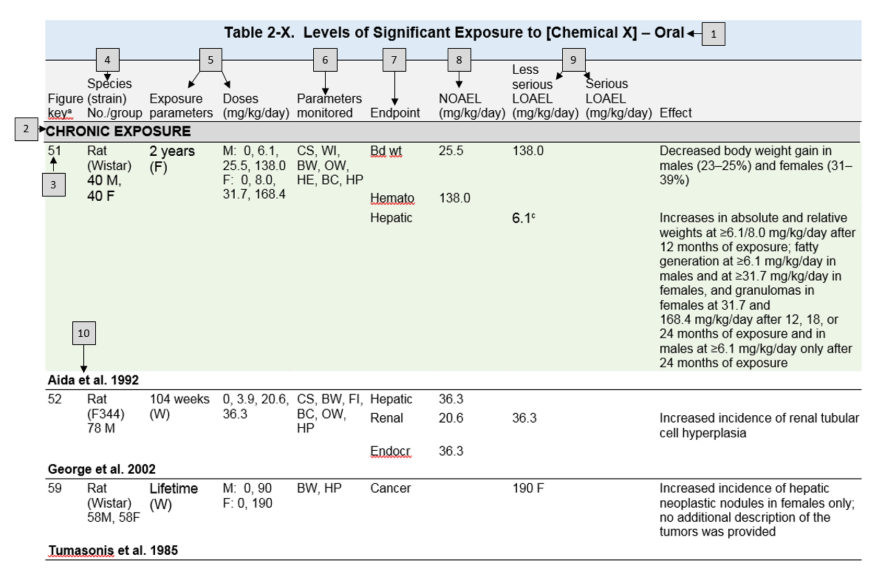
See Sample LSE Figure (page D-6)

LSE figures graphically illustrate the data presented in the corresponding LSE tables. Figures help the reader quickly compare health effects according to exposure concentrations for particular exposure periods.

(12) <u>Exposure period</u>. The same exposure periods appear as in the LSE table. In this example, health effects observed within the chronic exposure period are illustrated.

- (13) <u>Endpoint</u>. These are the categories of health effects for which reliable quantitative data exist. The same health effect endpoints appear in the LSE table.
- (14) <u>Levels of exposure</u>. Concentrations or doses for each health effect in the LSE tables are graphically displayed in the LSE figures. Exposure concentration or dose is measured on the log scale "y" axis. Inhalation exposure is reported in mg/m³ or ppm and oral exposure is reported in mg/kg/day.
- (15) <u>LOAEL</u>. In this example, the half-shaded circle that is designated 51R identifies a LOAEL critical endpoint in the rat upon which a chronic oral exposure MRL is based. The key number 51 corresponds to the entry in the LSE table. The dashed descending arrow indicates the extrapolation from the exposure level of 6.1 mg/kg/day (see entry 51 in the sample LSE table) to the MRL of 0.008 mg/kg/day (see footnote "c" in the sample LSE table).
- (16) <u>CEL</u>. Key number 59R is one of studies for which CELs were derived. The diamond symbol refers to a CEL for the test species (rat). The number 59 corresponds to the entry in the LSE table.
- (17) <u>Key to LSE figure</u>. The key provides the abbreviations and symbols used in the figure.

APPENDIX D



The number corresponds to entries in Figure 2-x.

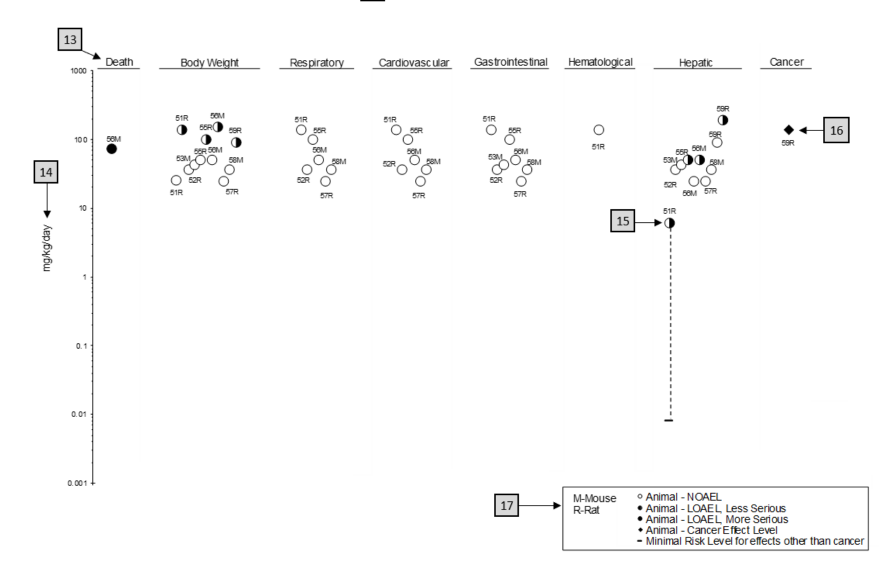
¹¹ bused to derive an acute-duration oral minimal risk level (MRL) of 0.1 mg/kg/day based on the BMDLos of 10 mg/kg/day and an uncertainty factor of 100 (10 for extrapolation from animals to humans and 10 for human variability).

^{*}Used to derive a chronic-duration oral MRL of 0.008 mg/kg/day based on the BMDL10 of 0.78 mg/kg/day and an uncertainty factor of 100 (10 for extrapolation from animals to humans and 10 for human variability).

APPENDIX D

Figure 2-X. Levels of Significant Exposure to [Chemical X] - Oral

12 → Chronic (≥365 days)



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APPENDIX E. QUICK REFERENCE FOR HEALTH CARE PROVIDERS

Toxicological Profiles are a unique compilation of toxicological information on a given hazardous substance. Each profile reflects a comprehensive and extensive evaluation, summary, and interpretation of available toxicologic and epidemiologic information on a substance. Health care providers treating patients potentially exposed to hazardous substances may find the following information helpful for fast answers to often-asked questions.

Primary Chapters/Sections of Interest

- **Chapter 1: Relevance to Public Health:** The Relevance to Public Health Section provides an overview of exposure and health effects and evaluates, interprets, and assesses the significance of toxicity data to human health. A table listing minimal risk levels (MRLs) is also included in this chapter.
- **Chapter 2: Health Effects**: Specific health effects identified in both human and animal studies are reported by type of health effect (e.g., death, hepatic, renal, immune, reproductive), route of exposure (e.g., inhalation, oral, dermal), and length of exposure (e.g., acute, intermediate, and chronic).

NOTE: Not all health effects reported in this section are necessarily observed in the clinical setting.

Pediatrics:

Section 3.2 Children and Other Populations that are Unusually Susceptible

Section 3.3 Biomarkers of Exposure and Effect

ATSDR Information Center

Phone: 1-800-CDC-INFO (800-232-4636) or 1-888-232-6348 (TTY)

Internet: http://www.atsdr.cdc.gov

ATSDR develops educational and informational materials for health care providers categorized by hazardous substance, clinical condition, and/or by susceptible population. The following additional materials are available online:

- Clinician Briefs and Overviews discuss health effects and approaches to patient management in a brief/factsheet style. They are narrated PowerPoint presentations with Continuing Education credit available (see https://www.atsdr.cdc.gov/environmental-medicine/hcp/emhsis/index.html).
- Managing Hazardous Materials Incidents is a set of recommendations for on-scene (prehospital) and hospital medical management of patients exposed during a hazardous materials incident (see https://www.atsdr.cdc.gov/MHMI/index.html).
- Fact Sheets (ToxFAQsTM) provide answers to frequently asked questions about toxic substances (see https://wwwn.cdc.gov/TSP/ToxFAQs/ToxFAQsLanding.aspx).

Other Agencies and Organizations

- The National Center for Environmental Health (NCEH) focuses on preventing or controlling disease, injury, and disability related to the interactions between people and their environment outside the workplace. Contact: NCEH, Mailstop F-29, 4770 Buford Highway, NE, Atlanta, GA 30341-3724 Phone: 770-488-7000 FAX: 770-488-7015 Web Page: https://www.cdc.gov/nceh/.
- The National Institute for Occupational Safety and Health (NIOSH) conducts research on occupational diseases and injuries, responds to requests for assistance by investigating problems of health and safety in the workplace, recommends standards to the Occupational Safety and Health Administration (OSHA) and the Mine Safety and Health Administration (MSHA), and trains professionals in occupational safety and health. Contact: NIOSH, 400 7th Street, S.W., Suite 5W, Washington, DC 20024 Phone: 202-245-0625 or 1-800-CDC-INFO (800-232-4636) Web Page: https://www.cdc.gov/niosh/.
- The National Institute of Environmental Health Sciences (NIEHS) is the principal federal agency for biomedical research on the effects of chemical, physical, and biologic environmental agents on human health and well-being. Contact: NIEHS, PO Box 12233, 104 T.W. Alexander Drive, Research Triangle Park, NC 27709 Phone: 919-541-3212 Web Page: https://www.niehs.nih.gov/.

Clinical Resources (Publicly Available Information)

- The Association of Occupational and Environmental Clinics (AOEC) has developed a network of clinics in the United States to provide expertise in occupational and environmental issues. Contact:

 AOEC, 1010 Vermont Avenue, NW, #513, Washington, DC 20005 Phone: 202-347-4976
 FAX: 202-347-4950 e-mail: AOEC@AOEC.ORG Web Page: http://www.aoec.org/.
- The American College of Occupational and Environmental Medicine (ACOEM) is an association of physicians and other health care providers specializing in the field of occupational and environmental medicine. Contact: ACOEM, 25 Northwest Point Boulevard, Suite 700, Elk Grove Village, IL 60007-1030 Phone: 847-818-1800 FAX: 847-818-9266 Web Page: http://www.acoem.org/.
- The American College of Medical Toxicology (ACMT) is a nonprofit association of physicians with recognized expertise in medical toxicology. Contact: ACMT, 10645 North Tatum Boulevard, Suite 200-111, Phoenix AZ 85028 Phone: 844-226-8333 FAX: 844-226-8333 Web Page: http://www.acmt.net.
- The Pediatric Environmental Health Specialty Units (PEHSUs) is an interconnected system of specialists who respond to questions from public health professionals, clinicians, policy makers, and the public about the impact of environmental factors on the health of children and reproductive-aged adults. Contact information for regional centers can be found at https://www.pehsu.net/.
- The American Association of Poison Control Centers (AAPCC) provide support on the prevention and treatment of poison exposures. Contact: AAPCC, 515 King Street, Suite 510, Alexandria VA 22314 Phone: 701-894-1858 Poison Help Line: 1-800-222-1222 Web Page: http://www.aapcc.org/.

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APPENDIX F. GLOSSARY

Absorption—The process by which a substance crosses biological membranes and enters systemic circulation. Absorption can also refer to the taking up of liquids by solids, or of gases by solids or liquids.

Acute Exposure—Exposure to a chemical for a duration of \leq 14 days, as specified in the Toxicological Profiles.

Adsorption—The adhesion in an extremely thin layer of molecules (as of gases, solutes, or liquids) to the surfaces of solid bodies or liquids with which they are in contact.

Adsorption Coefficient (K_{oc}) —The ratio of the amount of a chemical adsorbed per unit weight of organic carbon in the soil or sediment to the concentration of the chemical in solution at equilibrium.

Adsorption Ratio (Kd)—The amount of a chemical adsorbed by sediment or soil (i.e., the solid phase) divided by the amount of chemical in the solution phase, which is in equilibrium with the solid phase, at a fixed solid/solution ratio. It is generally expressed in micrograms of chemical sorbed per gram of soil or sediment.

Benchmark Dose (BMD) or Benchmark Concentration (BMC)—is the dose/concentration corresponding to a specific response level estimate using a statistical dose-response model applied to either experimental toxicology or epidemiology data. For example, a BMD₁₀ would be the dose corresponding to a 10% benchmark response (BMR). The BMD is determined by modeling the dose-response curve in the region of the dose-response relationship where biologically observable data are feasible. The BMDL or BMCL is the 95% lower confidence limit on the BMD or BMC.

Bioconcentration Factor (BCF)—The quotient of the concentration of a chemical in aquatic organisms at a specific time or during a discrete time period of exposure divided by the concentration in the surrounding water at the same time or during the same period.

Biomarkers—Indicators signaling events in biologic systems or samples, typically classified as markers of exposure, effect, and susceptibility.

Cancer Effect Level (CEL)—The lowest dose of a chemical in a study, or group of studies, that produces significant increases in the incidence of cancer (or malignant tumors) between the exposed population and its appropriate control.

Carcinogen—A chemical capable of inducing cancer.

Case-Control Study—A type of epidemiological study that examines the relationship between a particular outcome (disease or condition) and a variety of potential causative agents (such as toxic chemicals). In a case-control study, a group of people with a specified and well-defined outcome is identified and compared to a similar group of people without the outcome.

Case Report—A report that describes a single individual with a particular disease or exposure. These reports may suggest some potential topics for scientific research, but are not actual research studies.

Case Series—Reports that describe the experience of a small number of individuals with the same disease or exposure. These reports may suggest potential topics for scientific research, but are not actual research studies.

Ceiling Value—A concentration that must not be exceeded.

Chronic Exposure—Exposure to a chemical for \geq 365 days, as specified in the Toxicological Profiles.

Clastogen—A substance that causes breaks in chromosomes resulting in addition, deletion, or rearrangement of parts of the chromosome.

Cohort Study—A type of epidemiological study of a specific group or groups of people who have had a common insult (e.g., exposure to an agent suspected of causing disease or a common disease) and are followed forward from exposure to outcome, and who are disease-free at start of follow-up. Often, at least one exposed group is compared to one unexposed group, while in other cohorts, exposure is a continuous variable and analyses are directed towards analyzing an exposure-response coefficient.

Cross-sectional Study—A type of epidemiological study of a group or groups of people that examines the relationship between exposure and outcome to a chemical or to chemicals at a specific point in time.

Data Needs—Substance-specific informational needs that, if met, would reduce the uncertainties of human health risk assessment.

Developmental Toxicity—The occurrence of adverse effects on the developing organism that may result from exposure to a chemical prior to conception (either parent), during prenatal development, or postnatally to the time of sexual maturation. Adverse developmental effects may be detected at any point in the life span of the organism.

Dose-Response Relationship—The quantitative relationship between the amount of exposure to a toxicant and the incidence of the response or amount of the response.

Embryotoxicity and Fetotoxicity—Any toxic effect on the conceptus as a result of prenatal exposure to a chemical; the distinguishing feature between the two terms is the stage of development during which the effect occurs. Effects include malformations and variations, altered growth, and *in utero* death.

Epidemiology—The investigation of factors that determine the frequency and distribution of disease or other health-related conditions within a defined human population during a specified period.

Excretion—The process by which metabolic waste products are removed from the body.

Genotoxicity—A specific adverse effect on the genome of living cells that, upon the duplication of affected cells, can be expressed as a mutagenic, clastogenic, or carcinogenic event because of specific alteration of the molecular structure of the genome.

Half-life—A measure of rate for the time required to eliminate one-half of a quantity of a chemical from the body or environmental media.

Health Advisory—An estimate of acceptable drinking water levels for a chemical substance derived by EPA and based on health effects information. A health advisory is not a legally enforceable federal standard, but serves as technical guidance to assist federal, state, and local officials.

Immediately Dangerous to Life or Health (IDLH)—A condition that poses a threat of life or health, or conditions that pose an immediate threat of severe exposure to contaminants that are likely to have adverse cumulative or delayed effects on health.

Immunotoxicity—Adverse effect on the functioning of the immune system that may result from exposure to chemical substances.

Incidence—The ratio of new cases of individuals in a population who develop a specified condition to the total number of individuals in that population who could have developed that condition in a specified time period.

Intermediate Exposure—Exposure to a chemical for a duration of 15–364 days, as specified in the Toxicological Profiles.

In Vitro—Isolated from the living organism and artificially maintained, as in a test tube.

In Vivo—Occurring within the living organism.

Lethal Concentration_(LO) (LC_{LO)}—The lowest concentration of a chemical in air that has been reported to have caused death in humans or animals.

Lethal Concentration₍₅₀₎ (LC₅₀)—A calculated concentration of a chemical in air to which exposure for a specific length of time is expected to cause death in 50% of a defined experimental animal population.

Lethal Dose_(LO) (LD_{Lo)}—The lowest dose of a chemical introduced by a route other than inhalation that has been reported to have caused death in humans or animals.

Lethal Dose₍₅₀₎ (LD₅₀)—The dose of a chemical that has been calculated to cause death in 50% of a defined experimental animal population.

Lethal Time $_{(50)}$ (LT₅₀)—A calculated period of time within which a specific concentration of a chemical is expected to cause death in 50% of a defined experimental animal population.

Lowest-Observed-Adverse-Effect Level (LOAEL)—The lowest exposure level of chemical in a study, or group of studies, that produces statistically or biologically significant increases in frequency or severity of adverse effects between the exposed population and its appropriate control.

Lymphoreticular Effects—Represent morphological effects involving lymphatic tissues such as the lymph nodes, spleen, and thymus.

Malformations—Permanent structural changes that may adversely affect survival, development, or function.

Metabolism—Process in which chemical substances are biotransformed in the body that could result in less toxic and/or readily excreted compounds or produce a biologically active intermediate.

Minimal LOAEL—Indicates a minimal adverse effect or a reduced capacity of an organ or system to absorb additional toxic stress that does not necessarily lead to the inability of the organ or system to function normally.

Minimal Risk Level (MRL)—An estimate of daily human exposure to a hazardous substance that is likely to be without an appreciable risk of adverse noncancer health effects over a specified route and duration of exposure.

Modifying Factor (MF)—A value (greater than zero) that is applied to the derivation of a Minimal Risk Level (MRL) to reflect additional concerns about the database that are not covered by the uncertainty factors. The default value for a MF is 1.

Morbidity—The state of being diseased; the morbidity rate is the incidence or prevalence of a disease in a specific population.

Mortality—Death; the mortality rate is a measure of the number of deaths in a population during a specified interval of time.

Mutagen—A substance that causes mutations, which are changes in the DNA sequence of a cell's DNA. Mutations can lead to birth defects, miscarriages, or cancer.

Necropsy—The gross examination of the organs and tissues of a dead body to determine the cause of death or pathological conditions.

Neurotoxicity—The occurrence of adverse effects on the nervous system following exposure to a hazardous substance.

No-Observed-Adverse-Effect Level (NOAEL)—The dose of a chemical at which there were no statistically or biologically significant increases in frequency or severity of adverse effects seen between the exposed population and its appropriate control. Although effects may be produced at this dose, they are not considered to be adverse.

Octanol-Water Partition Coefficient (K_{ow})—The equilibrium ratio of the concentrations of a chemical in n-octanol and water, in dilute solution.

Odds Ratio (OR)—A means of measuring the association between an exposure (such as toxic substances and a disease or condition) that represents the best estimate of relative risk (risk as a ratio of the incidence among subjects exposed to a particular risk factor divided by the incidence among subjects who were not exposed to the risk factor). An odds ratio that is greater than 1 is considered to indicate greater risk of disease in the exposed group compared to the unexposed group.

Permissible Exposure Limit (PEL)—An Occupational Safety and Health Administration (OSHA) regulatory limit on the amount or concentration of a substance not to be exceeded in workplace air averaged over any 8-hour work shift of a 40-hour workweek.

Pesticide—General classification of chemicals specifically developed and produced for use in the control of agricultural and public health pests (insects or other organisms harmful to cultivated plants or animals).

Pharmacokinetics—The dynamic behavior of a material in the body, used to predict the fate (disposition) of an exogenous substance in an organism. Utilizing computational techniques, it provides the means of studying the absorption, distribution, metabolism, and excretion of chemicals by the body.

Pharmacokinetic Model—A set of equations that can be used to describe the time course of a parent chemical or metabolite in an animal system. There are two types of pharmacokinetic models: data-based and physiologically-based. A data-based model divides the animal system into a series of compartments, which, in general, do not represent real, identifiable anatomic regions of the body, whereas the physiologically-based model compartments represent real anatomic regions of the body.

Physiologically Based Pharmacodynamic (PBPD) Model—A type of physiologically based dose-response model that quantitatively describes the relationship between target tissue dose and toxic endpoints. These models advance the importance of physiologically based models in that they clearly describe the biological effect (response) produced by the system following exposure to an exogenous substance.

Physiologically Based Pharmacokinetic (PBPK) Model—A type of physiologically based dose-response model that is comprised of a series of compartments representing organs or tissue groups with realistic weights and blood flows. These models require a variety of physiological information, including tissue volumes, blood flow rates to tissues, cardiac output, alveolar ventilation rates, and possibly membrane permeabilities. The models also utilize biochemical information, such as blood:air partition coefficients, and metabolic parameters. PBPK models are also called biologically based tissue dosimetry models.

Prevalence—The number of cases of a disease or condition in a population at one point in time.

Prospective Study—A type of cohort study in which a group is followed over time and the pertinent observations are made on events occurring after the start of the study.

Recommended Exposure Limit (REL)—A National Institute for Occupational Safety and Health (NIOSH) time-weighted average (TWA) concentration for up to a 10-hour workday during a 40-hour workweek.

Reference Concentration (RfC)—An estimate (with uncertainty spanning perhaps an order of magnitude) of a continuous inhalation exposure to the human population (including sensitive subgroups) that is likely to be without an appreciable risk of deleterious noncancer health effects during a lifetime. The inhalation RfC is expressed in units of mg/m³ or ppm.

Reference Dose (RfD)—An estimate (with uncertainty spanning perhaps an order of magnitude) of the daily oral exposure of the human population to a potential hazard that is likely to be without risk of deleterious noncancer health effects during a lifetime. The oral RfD is expressed in units of mg/kg/day.

Reportable Quantity (RQ)—The quantity of a hazardous substance that is considered reportable under the Comprehensive Environmental Response, Compensation, and Liability Act (CERCLA). RQs are (1) ≥1 pound or (2) for selected substances, an amount established by regulation either under CERCLA or under Section 311 of the Clean Water Act. Quantities are measured over a 24-hour period.

Reproductive Toxicity—The occurrence of adverse effects on the reproductive system that may result from exposure to a hazardous substance. The toxicity may be directed to the reproductive organs and/or the related endocrine system. The manifestation of such toxicity may be noted as alterations in sexual behavior, fertility, pregnancy outcomes, or modifications in other functions that are dependent on the integrity of this system.

Retrospective Study—A type of cohort study based on a group of persons known to have been exposed at some time in the past. Data are collected from routinely recorded events, up to the time the study is undertaken. Retrospective studies are limited to causal factors that can be ascertained from existing records and/or examining survivors of the cohort.

Risk—The possibility or chance that some adverse effect will result from a given exposure to a hazardous substance.

Risk Factor—An aspect of personal behavior or lifestyle, an environmental exposure, existing health condition, or an inborn or inherited characteristic that is associated with an increased occurrence of disease or other health-related event or condition.

Risk Ratio/Relative Risk—The ratio of the risk among persons with specific risk factors compared to the risk among persons without risk factors. A risk ratio that is greater than 1 indicates greater risk of disease in the exposed group compared to the unexposed group.

Serious LOAEL—A dose that evokes failure in a biological system and can lead to morbidity or mortality.

Short-Term Exposure Limit (STEL)—A STEL is a 15-minute TWA exposure that should not be exceeded at any time during a workday.

Standardized Mortality Ratio (SMR)—A ratio of the observed number of deaths and the expected number of deaths in a specific standard population.

Target Organ Toxicity—This term covers a broad range of adverse effects on target organs or physiological systems (e.g., renal, cardiovascular) extending from those arising through a single limited exposure to those assumed over a lifetime of exposure to a chemical.

Teratogen—A chemical that causes structural defects that affect the development of an organism.

Threshold Limit Value (TLV)—An American Conference of Governmental Industrial Hygienists (ACGIH) concentration of a substance to which it is believed that nearly all workers may be repeatedly exposed, day after day, for a working lifetime without adverse effect. The TLV may be expressed as a Time-Weighted Average (TLV-TWA), as a Short-Term Exposure Limit (TLV-STEL), or as a ceiling limit (TLV-C).

Time-Weighted Average (TWA)—An average exposure within a given time period.

Toxicokinetic—The absorption, distribution, metabolism, and elimination of toxic compounds in the living organism.

Toxics Release Inventory (TRI)—The TRI is an EPA program that tracks toxic chemical releases and pollution prevention activities reported by industrial and federal facilities.

Uncertainty Factor (UF)—A factor used in operationally deriving the Minimal Risk Level (MRL), Reference Dose (RfD), or Reference Concentration (RfC) from experimental data. UFs are intended to account for (1) the variation in sensitivity among the members of the human population, (2) the uncertainty in extrapolating animal data to the case of human, (3) the uncertainty in extrapolating from data obtained in a study that is of less than lifetime exposure, and (4) the uncertainty in using lowest-observed-adverse-effect level (LOAEL) data rather than no-observed-adverse-effect level (NOAEL) data. A default for each individual UF is 10; if complete certainty in data exists, a value of 1 can be used; however, a reduced UF of 3 may be used on a case-by-case basis (3 being the approximate logarithmic average of 10 and 1).

Xenobiotic—Any substance that is foreign to the biological system.

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APPENDIX G. ACRONYMS, ABBREVIATIONS, AND SYMBOLS

AAPCC American Association of Poison Control Centers

ACGIH American Conference of Governmental Industrial Hygienists
ACOEM American College of Occupational and Environmental Medicine

ACMT American College of Medical Toxicology

ADI acceptable daily intake

ADME absorption, distribution, metabolism, and excretion

AEGL Acute Exposure Guideline Level AIC Akaike's information criterion

AIHA American Industrial Hygiene Association

ALT alanine aminotransferase

AOEC Association of Occupational and Environmental Clinics

AP alkaline phosphatase AST aspartate aminotransferase

atm atmosphere

ATSDR Agency for Toxic Substances and Disease Registry

AWQC Ambient Water Quality Criteria

BCF bioconcentration factor

BMD/C benchmark dose or benchmark concentration

BMD_X dose that produces a X% change in response rate of an adverse effect

BMDL_X 95% lower confidence limit on the BMD_X

BMDS Benchmark Dose Software BMR benchmark response BUN blood urea nitrogen

C centigrade CAA Clean Air Act

CAS Chemical Abstract Services

CDC Centers for Disease Control and Prevention

CEL cancer effect level

CERCLA Comprehensive Environmental Response, Compensation, and Liability Act

CFR Code of Federal Regulations

Ci curie

CI confidence interval

cm centimeter

CPSC Consumer Products Safety Commission

CWA Clean Water Act
DNA deoxyribonucleic acid
DOD Department of Defense
DOE Department of Energy
DWEL drinking water exposure level

EAFUS Everything Added to Food in the United States

ECG/EKG electrocardiogram
EEG electroencephalogram

EPA Environmental Protection Agency
ERPG emergency response planning guidelines

F Fahrenheit

F1 first-filial generation

FDA Food and Drug Administration

FIFRA Federal Insecticide, Fungicide, and Rodenticide Act

FR Federal Register

CHLOROETHANE G-2 APPENDIX G

FSH follicle stimulating hormone

g gram

GC gas chromatography
gd gestational day
GGT γ-glutamyl transferase
GRAS generally recognized as safe
HEC human equivalent concentration

HED human equivalent dose

HHS Department of Health and Human Services HPLC high-performance liquid chromatography

HSDB Hazardous Substances Data Bank

IARC International Agency for Research on Cancer IDLH immediately dangerous to life and health IRIS Integrated Risk Information System

Kd adsorption ratio kg kilogram

kilokilogram; 1 kilokilogram is equivalent to 1,000 kilograms and 1 metric ton

 K_{oc} organic carbon partition coefficient K_{ow} octanol-water partition coefficient

L liter

 $\begin{array}{lll} LC & liquid chromatography \\ LC_{50} & lethal concentration, 50\% \ kill \\ LC_{Lo} & lethal concentration, low \\ LD_{50} & lethal dose, 50\% \ kill \\ LD_{Lo} & lethal dose, low \\ LDH & lactate dehydrogenase \\ LH & luteinizing hormone \\ \end{array}$

LOAEL lowest-observed-adverse-effect level LSE Level of Significant Exposure

LESE Level of Significant Expo

LT₅₀ lethal time, 50% kill

m meter mCi millicurie

MCL maximum contaminant level MCLG maximum contaminant level goal

MF modifying factor mg milligram mL milliliter mm millimeter

mmHg millimeters of mercury

mmol millimole

MRL Minimal Risk Level MS mass spectrometry

MSHA Mine Safety and Health Administration

Mt metric ton

NAAQS National Ambient Air Quality Standard

NAS National Academy of Science

NCEH National Center for Environmental Health

ND not detected ng nanogram

NHANES National Health and Nutrition Examination Survey
NIEHS National Institute of Environmental Health Sciences

CHLOROETHANE G-3 APPENDIX G

NIOSH National Institute for Occupational Safety and Health

NLM National Library of Medicine

nm nanometer nmol nanomole

NOAEL no-observed-adverse-effect level

NPL National Priorities List

NR not reported

NRC National Research Council

NS not specified

NTP National Toxicology Program

OR odds ratio

OSHA Occupational Safety and Health Administration

PAC Protective Action Criteria

PAH polycyclic aromatic hydrocarbon

PBPD physiologically based pharmacodynamic PBPK physiologically based pharmacokinetic

PEHSU Pediatric Environmental Health Specialty Unit

PEL permissible exposure limit

PEL-C permissible exposure limit-ceiling value

pg picogram
PND postnatal day
POD point of departure
ppb parts per billion

ppbv parts per billion by volume

ppm parts per million ppt parts per trillion

REL recommended exposure limit

REL-C recommended exposure level-ceiling value

RfC reference concentration

RfD reference dose RNA ribonucleic acid

SARA Superfund Amendments and Reauthorization Act

SCE sister chromatid exchange

SD standard deviation SE standard error

SGOT serum glutamic oxaloacetic transaminase (same as aspartate aminotransferase or AST)
SGPT serum glutamic pyruvic transaminase (same as alanine aminotransferase or ALT)

SIC standard industrial classification

SLOAEL serious lowest-observed-adverse-effect level

SMR standardized mortality ratio sRBC sheep red blood cell STEL short term exposure limit TLV threshold limit value

TLV-C threshold limit value-ceiling value

TRI Toxics Release Inventory
TSCA Toxic Substances Control Act

TWA time-weighted average UF uncertainty factor U.S. United States

USDA United States Department of Agriculture

USGS United States Geological Survey

CHLOROETHANE G-4 APPENDIX G

USNRC U.S. Nuclear Regulatory Commission

VOC volatile organic compound

WBC white blood cell

WHO World Health Organization

> greater than

greater than or equal toequal to

= equal to < less than

 \leq less than or equal to

 q_1^* cancer slope factor

negativepositive

(+) weakly positive result(-) weakly negative result