2-HEXANONE

### 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. Serious health effects (such as irreparable damage to the liver or kidneys, or 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.

A-1

#### APPENDIX A

A-2

Proposed MRLs undergo a rigorous review process: Health Effects/MRL Workgroup reviews within the Division of Toxicology and Human Health Sciences, 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 Division of Toxicology and Human Health Sciences, Agency for Toxic Substances and Disease Registry, 1600 Clifton Road NE, Mailstop S102-1, Atlanta, Georgia 30329-4027.

Chemical Name:	2-Hexanone
CAS Numbers:	591-78-6
Date:	February 2020
<b>Profile Status:</b>	Final
Route:	Inhalation
Duration:	Acute

*MRL Summary:* There are insufficient data for derivation of an acute-duration inhalation MRL.

**Rationale for Not Deriving an MRL:** An acute-duration inhalation MRL for 2-hexanone was not derived due to lack of appropriate studies in humans or in animals. The only information located regarding effects in humans is that men (unknown number) exposed to  $\geq 2,300$  ppm vapors of a commercial-grade 2-hexanone for brief periods of time (25–60 seconds) found the atmospheres extremely disagreeable due to a strong odor and had irritation of the eyes and nasal passages (Schrenk et al. 1936). The same group of investigators reported that an unspecified number of guinea pigs exposed to 2,300 ppm 2-hexanone showed signs of eye and nose irritation after 1 minute of exposure and lacrimation after 10 minutes of exposure; no such signs were reported in guinea pigs exposed to 1,000 ppm 2-hexanone. Exposure of guinea pigs to 6,500 ppm for 540 minutes caused lethality.

Chemical Name:	2-Hexanone
CAS Numbers:	591-78-6
Date:	February 2020
Profile Status:	Final
Route:	Inhalation
Duration:	Intermediate

*MRL Summary:* There are insufficient data for derivation of an intermediate-duration inhalation MRL.

Rationale for Not Deriving an MRL: An intermediate-duration inhalation MRL for 2-hexanone was not derived because the lowest dose tested in an animal study induced serious neurological effects; ATSDR does not derive MRLs based on serious LOAELs. The lowest LOAEL reported was 50 ppm for sciatic nerve demyelination (a serious effect) in rats exposed for 8 hours/day, 5 days/week for 6 months (Duckett et al. 1979). Other intermediate-duration inhalation studies reported neurological effects (decreased nerve conduction velocity, neuropathy, and histopathological changes to central and peripheral nerves) occurring at higher exposure levels (≥100–1,000 ppm) (Duckett et al. 1979; Egan et al. 1980; Johnson et al. 1977; Katz et al. 1980; Mendell et al. 1974; Saida et al. 1976; Spencer et al. 1975). Other adverse effects included decreased body weight and decreased white blood cell count (Johnson et al. 1977; Katz et al. 1980); however, these effects were observed at exposure levels of 700–1,000 ppm.

Chemical Name:	2-Hexanone
CAS Numbers:	591-78-6
Date:	February 2020
<b>Profile Status:</b>	Final
Route:	Inhalation
Duration:	Chronic

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

**Rationale for Not Deriving an MRL:** A chronic-duration inhalation MRL for 2-hexanone was not derived due to lack of adequate data. Peripheral neuropathy, rated as moderate to severe, was observed in workers exposed to 2-hexanone (Allen et al. 1975; Billmaier et al. 1974). Workers were also exposed to methyl ethyl ketone, which has been shown to potentiate the effects induced by 2,5-hexanedione, the toxic metabolite of 2-hexanone (Saida et al. 1976; Yu et al. 2002). Therefore, these data are not suitable for derivation of the chronic-duration inhalation MRL. Two chronic-duration inhalation studies have been conducted in animals, one in rats exposed intermittently to 2-hexanone for 72 weeks (Krasavage and O'Donoghue 1977) and one in cats similarly exposed for 2 years (O'Donoghue and Krasavage 1979). In both studies, the lowest LOAEL reported was 330 ppm for serious neurological effects (peripheral neuropathy and axonal degeneration of central and peripheral nerves), with a NOAEL of 100 ppm. However, because a serious LOAEL of 50 ppm for sciatic nerve demyelination was observed in an intermediate-duration study in rats (Duckett et al. 1979), a chronic-duration inhalation MRL was not derived.

Chemical Name:	2-Hexanone
CAS Numbers:	591-78-6
Date:	February 2020
<b>Profile Status:</b>	Final
Route:	Oral
Duration:	Acute

*MRL Summary:* There are insufficient data for derivation of an acute-duration oral MRL.

**Rationale for Not Deriving an MRL:** An acute-duration oral MRL was not derived for 2-hexanone because of insufficient database. There are no human data, and the database in animals consists of a report of an oral  $LD_{50}$  in rats (Smyth et al. 1954) and a study of the potentiation action of 2-hexanone on liver and kidney toxicity caused by chloroform (Brown and Hewitt 1984). In that study, a single high dose of 1,500 mg/kg alone (only dose tested) did not induce morphological alterations in the liver, but produced epithelial degeneration in proximal tubules of the kidneys. However, the Brown and Hewitt (1984) study did not assess neurological effects, the most sensitive effect of 2-hexanone for both inhalation and oral exposure. Therefore, an acute-duration oral MRL was not derived.

Chemical Name:	2-Hexanone
CAS Numbers:	591-78-6
Date:	February 2020
<b>Profile Status:</b>	Final
Route:	Oral
Duration:	Intermediate

*MRL Summary:* There are insufficient data for derivation of an intermediate-duration oral MRL.

**Rationale for Not Deriving an MRL:** An intermediate-duration oral MRL for 2-hexanone was not derived because the lowest dose tested in an animal study induced serious neurological effects; a serious LOAEL is not considered a suitable basis for an MRL. The lowest LOAEL reported was 310 mg/kg/day for an approximately 40% reduction in locomotor activity in guinea pigs exposed to 2-hexanone in drinking water for 24 weeks; this effect was classified as a serious LOAEL. Pupillary responses to light stimuli were also significantly reduced at this dose level. Other intermediate-duration oral studies reported adverse neurological effects at higher daily doses. These effects include peripheral neuropathy in rats administered  $\geq$ 480 mg/kg/day (Union Carbide 1977); hindlimb weakness in rats administered 400 mg/kg/day (Eben et al. (1979), and paralysis in rats exposed to 660 mg/kg/day (Krasavage et al. 1980).

Chemical Name:	2-Hexanone
CAS Number:	591-78-6
Date:	February 2020
Profile Status:	Final
Route:	Oral
Duration:	Chronic
MRL:	0.05 mg/kg/day
Critical Effect:	Axonal swelling in peripheral nerve and spinal cord
Reference:	O'Donoghue et al. 1978
Point of Departure:	LOAEL of 143 mg/kg/day
Uncertainty Factor:	1,000
Modifying factor:	3
LSE Graph Key:	7
Species:	Rat

*MRL Summary:* A chronic-duration oral MRL of 0.05 mg/kg/day was derived for 2-hexanone based on axonal swelling in peripheral nerve in rats administered via drinking water for 13 months (O'Donoghue et al. 1978). The MRL is based on a LOAEL of 143 mg/kg/day and a total uncertainty factor of 1,000 (10 for use of a LOAEL, 10 for extrapolation from animals to humans, 10 for human variability) and a modifying factor of 3 for a high response at the lowest dose tested.

*Selection of the Critical Effect:* One chronic-duration oral study evaluated the toxicity of 2-hexanone (O'Donoghue et al. 1978). The lowest LOAEL reported was 143 mg/kg/day for neurotoxicity. The study also reported reduced body weight at 266 mg/kg/day and skeletal muscle myofiber atrophy at 560 mg/kg/day. Neurotoxicity also has been observed in intermediate-duration oral studies and in animals and humans exposed to 2-hexanone via inhalation; see discussion below (*Other Additional Studies or Pertinent Information that Lend Support to this MRL*).

*Selection of the Principal Study:* One chronic-duration oral exposure study was identified (O'Donoghue et al. 1978). This is a well-conducted study that conducted neurological examinations and microscopic examinations of numerous tissues.

#### Summary of the Principal Study:

O'Donoghue JL, Krasavage WJ, Terhaar CJ. 1978. A comparative chronic toxicity study of methyl npropyl ketone, methyl n-butyl ketone, and hexane by ingestion. Eastman Kodak Co. Submitted to the U.S. Environmental Protection Agency under TSCA Section 8E. EPA Document No. 88920008233. OTS0555051.

Groups of male Sprague-Dawley rats (10/group) were exposed to drinking water containing 0, 0.25, 0.5, or 1.0% 2-hexanone (96.1% purity) for 13 months. Based on water consumption and body weight data, the investigators calculated daily doses of 2-hexanone of 0, 143, 266, or 560 mg/kg/day. There were two control groups, each with 10 rats. Rats were observed daily for clinical signs; body weight measurements and neurological examinations were performed weekly. At termination of exposure, the hindlimb sciatic-plantar nerve, multiple levels of the spinal cord, medulla, and cerebellum from five rats per group were embedded in plastic for microscopic examination. Most tissues and organs from the highest dose group and target organs from lower dose groups were embedded in paraffin for light microscopy examination.

Clinical signs were restricted to neurological effects and reduced body weight. Final body weights were reduced approximately 6, 14, and 30% in the 143, 266, and 560 mg/kg/day dose groups, respectively. No information was provided regarding food consumption. Clinical neurological signs were seen in the 266 and 560 mg/kg/day groups. Signs first appeared on day 42 in the 560 mg/kg/day group and on day 77 in the 266 mg/kg/day group. All rats in the 560 mg/kg/day group showed severe deficits. Signs included decreased extension of hindlimbs, hindlimb weakness, waddling gait, dragging of hind paws, and loss of tone in hindlimb musculature with grossly observable atrophy of hindlimb musculature and axial muscles of the lumbar area. Weakness of the forelimbs was seen in three out of nine rats in the 560 mg/kg/day group by the end of the study. No clinical progression was apparent in the 266 mg/kg/day group. Histological examination showed that rats from all treated groups had "giant" axonal neuropathy. Axonal swelling and giant axonopathy were common in peripheral nerves and spinal cord from 560 mg/kg/day rats, less common in dorsal root ganglia, and rare in the brain. Sections embedded in plastic showed clumping of axonal organelles. Myelin alterations were also seen in peripheral nerves. Neurogenic skeletal muscle atrophy occurred in proximal and distal hindlimb musculature. Alterations in the 266 mg/kg/day group were similar but less severe. Less severe changes were seen in peripheral nerves from 143 mg/kg/day rats; fewer giant axons were evident, but myelin changes were more common. Spinal lesions and neurogenic muscle atrophy were minimal. Relevant incidence data are shown in Table A-1. No treatment-related gross or microscopic alterations were reported in tissues other than the nervous system and skeletal muscle.

	2-Hexanone for 13 Months					•
	Axonal swelling Myofibrillar atrophy				lar atrophy	
Dose Dorsal root Peripheral Quadriceps (mg/kg/day) Brain Spinal cord ganglia nerve muscle					Quadriceps muscle	Calf muscle
0	0/10	0/5	0/5	0/10	0/10	0/10
143	2/10	7/10 <sup>a</sup>	0/7	8/10 <sup>a</sup>	1/10	2/10
266	4/10 <sup>a</sup>	5/5 <sup>a</sup>	0/5	10/10 <sup>a</sup>	5/10 <sup>a</sup>	6/10 <sup>a</sup>
560	8/10 <sup>a</sup>	5/5ª	3/5	10/10 <sup>a</sup>	10/10 <sup>a</sup>	10/10 <sup>a</sup>

## Table A-1. Incidence Data for Neuropathological Lesions in Rats exposed to

<sup>a</sup>p<0.05 per Fisher Exact Test conducted by SRC, Inc.

Source: O'Donoghue et al. 1978

Selection of the Point of Departure for the MRL: Benchmark dose (BMD) modelling of the incidence data for axonal swelling in peripheral nerve of rats in the O'Donoghue et al. (1978) study was considered and rejected because a nearly maximum response level (80%) was reached with the lowest dose tested. In such cases, there is great uncertainty because the BMD may be just below the first dose or orders of magnitude lower (EPA 2012b). Therefore, the NOAEL/LOAEL approach was used to derive a chronicduration oral MRL for 2-hexanone.

*Calculations:* Conversion of drinking water concentrations to daily doses (mg/kg/day) was done by the investigators.

#### Uncertainty Factor (UF) and Modifying Factor (MF):

- 10 UF for use of a LOAEL
- 10 UF for extrapolation from animals to humans
- 10 UF for human variability

• 3 MF to account for an 80% response rate at the lowest dose

 $MRL = LOAEL \div (UFs \ x \ MF)$ 0.05 mg/kg/day = 143 mg/kg/day ÷ (1,000 x 3)

*Other Additional Studies or Pertinent Information that Lend Support to this MRL:* 2-Hexanone is well-known as a neurotoxic chemical that has been tested in a variety of animal species. Neurotoxicity, including hind limb weakness, paralysis, decreased locomotor activity, and peripheral neuropathy, was observed in intermediate-duration oral exposure studies reported in rats and guinea pigs (Abdel-Rahman et al. 1978; Eben et al. 1979; Krasavage et al. 1980; Union Carbide 1977). Although no studies examining oral exposure of 2-hexanone in humans were identified, case reports and studies in workers also provide evidence that 2-hexanone produces neurotoxicity in humans following inhalation exposure (Allen et al. 1975; Billmaier et al. 1974; Davenport et al. 1976; Mallov 1976). Inhalation studies in animals also demonstrate that 2-hexanone is neurotoxic, with adverse neurological effects following acute exposure of guinea pigs (Schrenk et al. 1936), intermediate exposure of monkeys, rats, and cats (Duckett et al. 1979; Egan et al. 1980; Johnson et al. 1977; Katz et al. 1980; Mendell et al. 1974; Saida et al. 1976; Spencer et al. 1975), and chronic exposure of rats and cats (Krasavage and O' Donoghue 1977, 1979). Because the toxic chemical form of 2-hexanone is the metabolite, 2,5-hexanedione, and 2,5-hexanedione is also a metabolite of *n*-hexane, additional relevant information can be found in documents on *n*-hexane (i.e., ATSDR 1999).

## APPENDIX B. LITERATURE SEARCH FRAMEWORK FOR 2-HEXANONE

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 2-hexanone.

## **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 2-hexanone. ATSDR primarily focused on peer-reviewed articles without publication date or language restrictions. Non-peer-reviewed studies that were considered relevant to the assessment of the health effects of 2-hexanone 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 2-hexanone 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 **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	

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

#### **B.1.1 Literature Search**

The current literature search was intended to update the draft toxicological profile for 2-hexanone released for public comment in April 2018. The following main databases were searched in March 2019:

- PubMed
- National Library of Medicine's TOXLINE
- 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 2-hexanone. 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 2-hexanone 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	Query string
PubMed	
03/2019	((591-78-6[rn] OR "Methyl n-Butyl Ketone"[mh] OR "2-Hexanone"[tw] OR "2- Oxohexane"[tw] OR "Butyl methyl ketone"[tw] OR "Butylmethyl Ketone"[tw] OR "Hexan-2- one"[tw] OR "Hexanone-2"[tw] OR ("MBK"[tw] AND (ketone OR hexanone)) OR "Methyl butyl ketone"[tw] OR "Methyl n-butyl ketone"[tw] OR "n-Butyl methyl ketone"[tw] OR "Propylacetone"[tw]) AND (2014/12/01 : 3000[dp] OR 2015/12/01 : 3000[mhda] OR 2015/12/01 : 3000[crdat] OR 2015/12/01 : 3000[edat])) OR ("2-HEXANON"[tw] OR "Hexan- 2-on"[tw] OR "hexan-2-ona"[tw] OR "hexane-2-one"[tw] OR "Ketone, butyl methyl"[tw] OR "MnBK"[tw])
Toxline	
03/2019	Year of Publication 2014 through 2019 (591-78-6[rn] OR "2-Hexanone" OR "2-Oxohexane" OR "Butyl methyl ketone" OR "Butylmethyl Ketone" OR "Hexan-2-one" OR "Hexanone-2" OR "Methyl butyl ketone" OR "Methyl n-butyl ketone" OR "n-Butyl methyl ketone" OR "Propylacetone" OR "2- HEXANON" OR "Hexan-2-on" OR "hexan-2-ona" OR "hexane-2-one" OR "Ketone, butyl methyl") AND ( ANEUPL [org] OR BIOSIS [org] OR CIS [org] OR DART [org] OR EMIC [org] OR EPIDEM [org] OR HEEP [org] OR HMTC [org] OR IPA [org] OR RISKLINE [org] OR MTGABS [org] OR NIOSH [org] OR NTIS [org] OR PESTAB [org] OR PPBIB [org] ) AND NOT PubMed [org] AND NOT pubdart [org]
	Year of Publication 2014 through 2019 ("MBK" AND (ketone OR hexanone)) AND (ANEUPL [org] OR BIOSIS [org] OR CIS [org] OR DART [org] OR EMIC [org] OR EPIDEM [org] OR HEEP [org] OR HMTC [org] OR IPA [org] OR RISKLINE [org] OR MTGABS [org] OR NIOSH [org] OR NTIS [org] OR PESTAB [org] OR PPBIB [org] ) AND NOT PubMed [org] AND NOT pubdart [org] ("2-HEXANON" OR "Hexan-2-on" OR "hexan-2-ona" OR "hexane-2-one" OR "Ketone, butyl methyl" OR "MnBK") AND (ANEUPL [org] OR BIOSIS [org] OR CIS [org] OR DART
	[org] OR EMIC [org] OR EPIDEM [org] OR HEEP [org] OR HMTC [org] OR IPA [org] OR RISKLINE [org] OR MTGABS [org] OR NIOSH [org] OR NTIS [org] OR PESTAB [org] OR PPBIB [org] ) AND NOT PubMed [org] AND NOT pubdart [org]
Toxcenter	
03/2019	<ul> <li>L1 1426 SEA FILE=TOXCENTER 591-78-6</li> <li>L2 1286 SEA FILE=TOXCENTER L1 NOT PATENT/DT</li> <li>L3 1253 SEA FILE=TOXCENTER L2 NOT TSCATS/FS</li> <li>L4 82 SEA FILE=TOXCENTER L3 AND ED&gt;=20151201</li> <li>L5 116 SEA FILE=TOXCENTER L3 AND PY&gt;2014</li> <li>L6 116 SEA FILE=TOXCENTER L4 OR L5 ANSWERS '1-115' FROM FILE TOXCENTER</li> </ul>

Table B-2. Database Query Strings		
Database search date Qu	uery string	
	ACT TOXQUERY/Q	
L1	BIOMARKER? OR NEUROLOG?)	
L1 EF	2 QUE (PHARMACOKIN? OR SUBCHRONIC OR PBPK OR PIDEMIOLOGY/ST,CT, IT)	
L1		
L1	<ul> <li>4 QUÉ (TOXICITY OR ADVERSE OR POISONING)/ST,CT,IT</li> <li>5 QUE (INHAL? OR PULMON? OR NASAL? OR LUNG? OR RESPIR?)</li> <li>6 QUE ((OCCUPATION? OR WORKPLACE? OR WORKER?) AND EXPOS?)</li> <li>7 QUE (ORAL OR ORALLY OR INGEST? OR GAVAGE? OR DIET OR DIETS</li> </ul>	
L1 PE	DIETARY OR DRINKING(W)WATER?) 8 QUE (MAXIMUM AND CONCENTRATION? AND (ALLOWABLE OR ERMISSIBLE))	
L1 L2 OF	QUE (FOETUS? OR FETAL? OR FOETAL? OR FERTIL? OR MALFORM?	
L2 L2		
L2 SF	23 QUE (SPERM OR SPERMAC? OR SPERMAG? OR SPERMATI? OR PERMAS? OR	
L2 SF	PERMATOX? OR	
	SPERMATOZ? OR SPERMATU? OR SPERMI? OR SPERMO?) 25 QUE (NEONAT? OR NEWBORN? OR DEVELOPMENT OR EVELOPMENTAL?)	
L2 L2	QUE (ENDOCRIN? AND DISRUPT?)	
L2 L2 L3 OF	<ul> <li>QUE (WEAN? OR OFFSPRING OR AGE(W)FACTOR?)</li> <li>QUE (DERMAL? OR DERMIS OR SKIN OR EPIDERM? OR CUTANEOUS?)</li> <li>QUE (CARCINOG? OR COCARCINOG? OR CANCER? OR PRECANCER?</li> </ul>	
L3		
L3	ARCINOM?) 22 QUE (GENETOX? OR GENOTOX? OR MUTAGEN? OR ENETIC(W)TOXIC?)	
L3 L3 L3 L3	QUE (NEPHROTOX? OR HEPATOTOX?)QUE (ENDOCRIN? OR ESTROGEN? OR ANDROGEN? OR HORMON?)QUE (OCCUPATION? OR WORKER? OR WORKPLACE? OR EPIDEM?)	

Table B-2.	Database	Query	/ Strings
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Database	
search date Que	ery string
	L28 OR L29 OR L30 OR L31 OR L32 OR L33 OR L34 OR L35
L37	QUE (RAT OR RATS OR MOUSE OR MICE OR GUINEA(W)PIG? OR
MUI	RIDAE
	OR DOG OR DOGS OR RABBIT? OR HAMSTER? OR PIG OR PIGS OR
SWI	NE
	OR PORCINE OR MONKEY? OR MACAQUE?)
L38	QUE (MARMOSET? OR FERRET? OR GERBIL? OR RODENT? OR
LAG	OMORPHA
	OR BABOON? OR CANINE OR CAT OR CATS OR FELINE OR MURINE)
L39	
L40	QUE (HUMAN OR HUMANS OR HOMINIDAE OR MAMMALS OR MAMMAL?
OR	
	PRIMATES OR PRIMATE?)
L41	QUE L39 OR L40
L42	
L43	
	ANSWERS '1-46' FROM FILE TOXCENTER
	D SCAN L43

## Table B-3. Strategies to Augment the Literature Search

Source	Query and number screened when available			
TSCATS via ChemView				
03/2019	Compounds searched: 591-78-6			
NTP				
03/2019	"591-78-6" "2-Hexanone" "Butylmethyl Ketone" "Methyl butyl ketone" "Methyl n-butyl ketone" "n-Butyl methyl ketone" "Hexan-2-one" "Hexanone-2" "Butyl methyl ketone" "2-Oxohexane" "Propylacetone" "Ketone, butyl methyl" "MBK" "mnbk" "2-HEXANON" "Hexan-2-on" "hexan-2-ona" "hexane-2-one"			
DTIC				
05/2019	<ul> <li>Synonyms in all fields search box</li> <li>"591-78-6" OR "2-Hexanone" OR "2-Oxohexane" OR "Butyl methyl ketone" OR</li> <li>"Hexanone-2" OR "Methyl butyl ketone" OR "Methyl n-butyl ketone" OR</li> <li>"Propylacetone" OR "n-Butyl methyl ketone" OR "Butylmethyl Ketone" OR "Hexan-2-one" OR "2-Hexanone Methyl n-butyl ketone" OR "2-hexanon" OR "hexan-2-on" OR</li> <li>"hexan-2-ona" OR "hexane-2-one" OR "ketone, butyl methyl" OR "MnBK" OR ("MBK" AND ("ketone" OR "hexanone"))</li> <li>Keywords in citation terms box</li> <li>"toxicity" OR "toxicology" OR "poisoning" OR "cancer" OR "carcinogens" OR</li> <li>"carcinogen" OR "neoplasms" OR "neoplasm" OR "oncogenesis" OR "teratogenic compounds" OR "lethality" OR "death" OR "body weight" OR "immunology" OR</li> <li>"genotoxicity" OR "inhalation" OR "dermal" OR "metabolism" OR "pharmacokinetics" OR</li> </ul>			

	Table B-3. Strategies to Augment the Literature Search
Source	Query and number screened when available
	"pharmacokinetic" OR "PBPK" OR "pharmacology" OR "organs" OR "skin" OR "tissues" OR "body fluids" OR "toxic agents" OR "rats" OR "mice" OR "mouse" OR "rat"
NIH RePORTER	
09/2019	"2-Hexanone" OR "2-Oxohexane" OR "Butyl methyl ketone" OR "Butylmethyl Ketone" OR "Hexan-2-one" OR "Hexanone-2" OR ("MBK" AND (ketone OR hexanone)) OR "Methyl butyl ketone" OR "Methyl n-butyl ketone" OR "n-Butyl methyl ketone" OR "Propylacetone" OR "2-HEXANON" OR "Hexan-2-on" OR "hexan-2-ona" OR "hexane- 2-one" OR "Ketone, butyl methyl" OR "MnBK" (Advanced), Search in: Projects AdminIC: All, Fiscal Year: Active Projects
Other	Identified throughout the assessment process

The 2019 results were:

- Number of records identified from PubMed, TOXLINE, and TOXCENTER (after duplicate removal): 86
- Number of records identified from other strategies: 23
- Total number of records to undergo literature screening: 109

## **B.1.2 Literature Screening**

A two-step process was used to screen the literature search to identify relevant studies on 2-hexanone:

- Title and abstract screen
- Full text screen

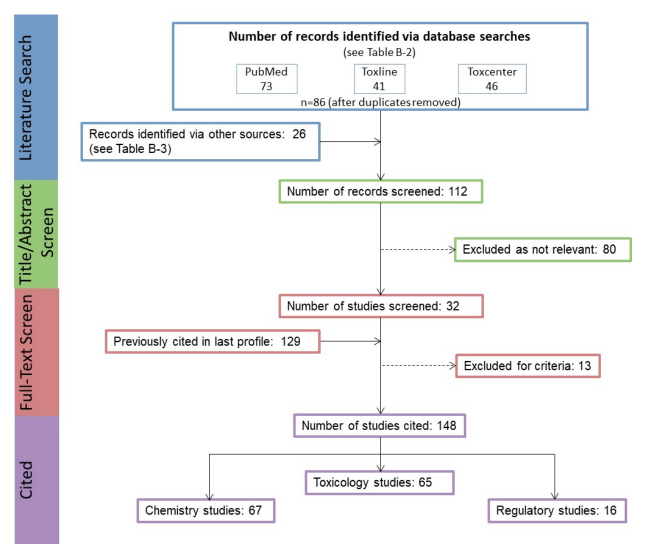
*Title and Abstract Screen.* Within the reference library, titles and abstracts were screened manually for relevance. Studies that were considered relevant (see Table B-1 for inclusion criteria) were moved to the second step of the literature screening process. Studies were excluded when the title and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

- Number of titles and abstracts screened: 112
- Number of studies considered relevant and moved to the next step: 32

*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: 32
- Number of studies cited in the pre-public draft of the toxicological profile: 129
- Total number of studies cited in the profile: 148

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



## Figure B-1. March 2019 Literature Search Results and Screen for 2-Hexanone

## APPENDIX C. 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 C-5)

- (1) <u>Route of exposure</u>. 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) <u>Exposure period</u>. 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.</p>
- (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) <u>Species (strain) No./group</u>. 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) <u>Parameters monitored.</u> This column lists the parameters used to assess health effects. Parameters monitored could include serum (blood) chemistry (BC), behavioral (BH), biochemical changes (BI), body weight (BW), clinical signs (CS), developmental toxicity (DX), enzyme activity (EA), food intake (FI), fetal toxicity (FX), gross necropsy (GN), hematology (HE), histopathology (HP), lethality (LE), maternal toxicity (MX), organ function (OF), ophthalmology (OP), organ weight (OW), teratogenicity (TG), urinalysis (UR), and water intake (WI).
- (7) <u>Endpoint</u>. 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 C-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.

(13) <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.

- (14) <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.
- (15) <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/m<sup>3</sup> or ppm and oral exposure is reported in mg/kg/day.
- (16) <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).
- (17) <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.
- (18) <u>Key to LSE figure</u>. The key provides the abbreviations and symbols used in the figure.

APPENDIX C

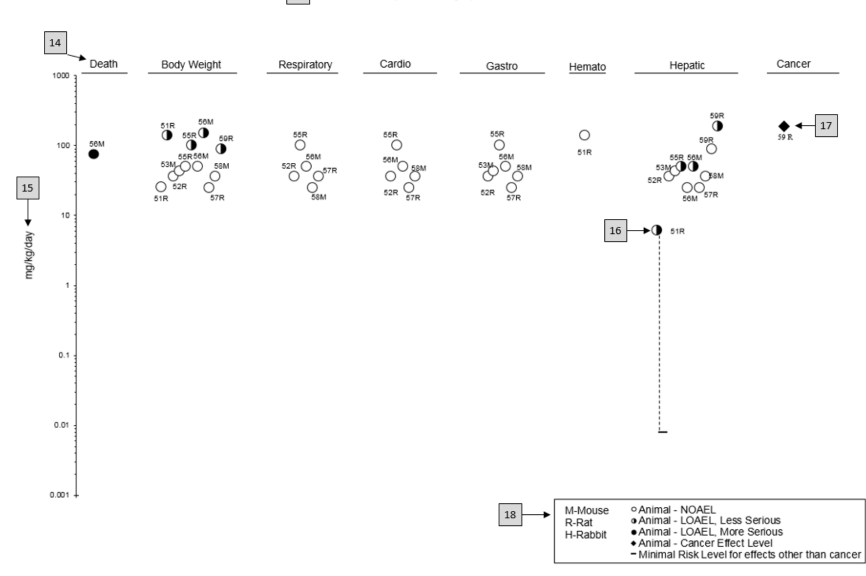
		-	1		_			
	4	5		6	7	8	Less 9	
	Species	₩	4	Ļ		¥	serious Serious	
<u> </u>	(strain)	Exposure	Doses	Parameters	_ +	NOAEL	LOAEL LOAEL	
<u>key</u> ª	<u> </u>	parameters	(mg/kg/day)	monitored	Endpoint	(mg/kg/day)	(mg/kg/day) (mg/kg/day)	Effect
CHRC	NIC EXP	DSURE						
51 ↑ 3	Rat (Wistar) 40 M, 40 F	2 years (F)	M: 0, 6.1, 25.5, 138.0 F: 0, 8.0, 31.7, 168.4	CS, WI, BW, OW, HE, BC, HP	Bd wt	25.5	138.0	Decreased body weight gain in males (23–25%) and females (31–39%)
	40 F		51.7, 100.4		Hemato	138.0		
,	0				Hepatic		6.1°	Increases in absolute and relative weights at $\geq 6.1/8.0$ mg/kg/day after 12 months of exposure; fatty generation at $\geq 6.1$ mg/kg/day in males and at $\geq 31.7$ mg/kg/day in females, and granulomas in females at 31.7 and 168.4 mg/kg/day after 12, 18, or 24 months of exposure and in males at $\geq 6.1$ mg/kg/day only after 24 months of exposure
	et al. 1992							
52	Rat	104 weeks		CS, BW, FI,	Hepatic	36.3		
	(F344) 78 M	(W)	36.3	BC, OW, HP	Renal	20.6	36.3	Increased incidence of renal tubula cell hyperplasia
Georg	e et al. 200	2			Endocr	36.3		
59	Rat (Wistar) 58M, 58F	Lifetime (W)	M: 0, 90 F: 0, 190	BW, HP	Cancer		190 F	Increased incidence of hepatic neoplastic nodules in females only; no additional description of the tumors was provided

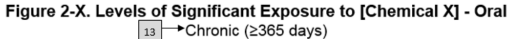
The number corresponds to entries in Figure 2-x.

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 C





## APPENDIX D. 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 3.2Children and Other Populations that are Unusually SusceptibleSection 3.3Biomarkers 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

The following additional materials are available online:

- *Case Studies in Environmental Medicine* are self-instructional publications designed to increase primary health care providers' knowledge of a hazardous substance in the environment and to aid in the evaluation of potentially exposed patients (see https://www.atsdr.cdc.gov/csem/csem.html).
- Managing Hazardous Materials Incidents is a three-volume 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.asp). Volumes I and II are planning guides to assist first responders and hospital emergency department personnel in planning for incidents that involve hazardous materials. Volume III—*Medical Management Guidelines for Acute Chemical Exposures*—is a guide for health care professionals treating patients exposed to hazardous materials.

*Fact Sheets (ToxFAQs*<sup>TM</sup>) provide answers to frequently asked questions about toxic substances (see https://www.atsdr.cdc.gov/toxfaqs/Index.asp).

#### **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, 395 E Street, S.W., Suite 9200, Patriots Plaza Building, Washington, DC 20201 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 http://pehsu.net/findhelp.html.
- *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/.

## APPENDIX E. 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 BMD<sub>10</sub> 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 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**<sub>(LO)</sub> ( $LC_{LO}$ )—The lowest concentration of a chemical in air that has been reported to have caused death in humans or animals.

**Lethal Concentration**<sub>(50)</sub> ( $LC_{50}$ )—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  $Dose_{(50)}$  (LD<sub>50</sub>)—The dose of a chemical that has been calculated to cause death in 50% of a defined experimental animal population.

**Lethal Time**<sub>(50)</sub> ( $LT_{50}$ )—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 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 doseresponse 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.

E-5

**Physiologically Based Pharmacokinetic (PBPK) Model**—A type of physiologically based doseresponse 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/m<sup>3</sup> 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  $(1) \ge 1$  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.

**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.

## APPENDIX F. 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 <sub>x</sub>	dose that produces a X% change in response rate of an adverse effect			
BMDL <sub>X</sub>	95% lower confidence limit on the $BMD_X$			
BMDS	Benchmark Dose Software			
BMR	benchmark response			
BUN	blood urea nitrogen			
С	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			
	centimeter			
cm				
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			

FSH	follicle stimulating hormone			
g	gram			
ĞC	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 Hazardous Substance Data Bank			
HSDB				
IARC	International Agency for Research on Cancer			
IDLH	immediately dangerous to life and health			
IRIS	Integrated Risk Information System			
Kd	adsorption ratio			
kg	kilogram			
kkg	kilokilogram; 1 kilokilogram is equivalent to 1,000 kilograms and 1 metric ton			
Koc	organic carbon partition coefficient			
K <sub>ow</sub>	octanol-water partition coefficient			
L	liter			
LC	liquid chromatography			
$LC_{50}$	lethal concentration, 50% kill			
LC <sub>Lo</sub>	lethal concentration, low			
$LD_{50}$	lethal dose, 50% kill			
$DL_{Lo}$	lethal dose, low			
LDH	lactic dehydrogenase			
LH	luteinizing hormone			
LOAEL	lowest-observed-adverse-effect level			
LSE	Level of Significant Exposure			
$LT_{50}$	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			

NIOGU	
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
PS	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 level/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 deviation
SGOT	
SGPT	serum glutamic oxaloacetic transaminase (same as aspartate aminotransferase or AST)
SIC	serum glutamic pyruvic transaminase (same as alanine aminotransferase or ALT) standard industrial classification
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
USNRC	U.S. Nuclear Regulatory Commission

VOC WBC WHO	volatile organic compound white blood cell World Health Organization
>	greater than
> > = < < < %	greater than or equal to
=	equal to
<	less than
$\leq$	less than or equal to
%	percent
α	alpha
β	beta
γ	gamma
δ	delta
μm	micrometer
μg	microgram
$\mathbf{q}_1^*$	cancer slope factor
_	negative
+	positive
(+)	weakly positive result
(-)	weakly negative result