

Provisional Peer-Reviewed Toxicity Values for
1-Chlorooctadecane
(CASRN 3386-33-2)

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COMMONLY USED ABBREVIATIONS AND ACRONYMS

| | | | |
|------------------|---|--------------------|--|
| α 2u-g | alpha 2u-globulin | MN | micronuclei |
| ACGIH | American Conference of Governmental Industrial Hygienists | MNPCE | micronucleated polychromatic erythrocyte |
| AIC | Akaike's information criterion | MOA | mode of action |
| ALD | approximate lethal dosage | MTD | maximum tolerated dose |
| ALT | alanine aminotransferase | NAG | N-acetyl- β -D-glucosaminidase |
| AST | aspartate aminotransferase | NCEA | National Center for Environmental Assessment |
| atm | atmosphere | NCI | National Cancer Institute |
| ATSDR | Agency for Toxic Substances and Disease Registry | NOAEL | no-observed-adverse-effect level |
| BMD | benchmark dose | NTP | National Toxicology Program |
| BMDL | benchmark dose lower confidence limit | NZW | New Zealand White (rabbit breed) |
| BMDS | Benchmark Dose Software | OCT | ornithine carbamoyl transferase |
| BMR | benchmark response | ORD | Office of Research and Development |
| BUN | blood urea nitrogen | PBPK | physiologically based pharmacokinetic |
| BW | body weight | PCNA | proliferating cell nuclear antigen |
| CA | chromosomal aberration | PND | postnatal day |
| CAS | Chemical Abstracts Service | POD | point of departure |
| CASRN | Chemical Abstracts Service Registry Number | POD _{ADJ} | duration-adjusted POD |
| CBI | covalent binding index | QSAR | quantitative structure-activity relationship |
| CHO | Chinese hamster ovary (cell line cells) | RBC | red blood cell |
| CL | confidence limit | RDS | replicative DNA synthesis |
| CNS | central nervous system | RfC | inhalation reference concentration |
| CPN | chronic progressive nephropathy | RfD | oral reference dose |
| CYP450 | cytochrome P450 | RGDR | regional gas dose ratio |
| DAF | dosimetric adjustment factor | RNA | ribonucleic acid |
| DEN | diethylnitrosamine | SAR | structure activity relationship |
| DMSO | dimethylsulfoxide | SCE | sister chromatid exchange |
| DNA | deoxyribonucleic acid | SD | standard deviation |
| EPA | Environmental Protection Agency | SDH | sorbitol dehydrogenase |
| FDA | Food and Drug Administration | SE | standard error |
| FEV1 | forced expiratory volume of 1 second | SGOT | glutamic oxaloacetic transaminase, also known as AST |
| GD | gestation day | SGPT | glutamic pyruvic transaminase, also known as ALT |
| GDH | glutamate dehydrogenase | SSD | systemic scleroderma |
| GGT | γ -glutamyl transferase | TCA | trichloroacetic acid |
| GSH | glutathione | TCE | trichloroethylene |
| GST | glutathione-S-transferase | TWA | time-weighted average |
| Hb/g-A | animal blood-gas partition coefficient | UF | uncertainty factor |
| Hb/g-H | human blood-gas partition coefficient | UF _A | interspecies uncertainty factor |
| HEC | human equivalent concentration | UF _H | intraspecies uncertainty factor |
| HED | human equivalent dose | UF _S | subchronic-to-chronic uncertainty factor |
| i.p. | intraperitoneal | UF _D | database uncertainty factor |
| IRIS | Integrated Risk Information System | U.S. | United States of America |
| IVF | in vitro fertilization | WBC | white blood cell |
| LC ₅₀ | median lethal concentration | | |
| LD ₅₀ | median lethal dose | | |
| LOAEL | lowest-observed-adverse-effect level | | |

PROVISIONAL PEER-REVIEWED TOXICITY VALUES FOR 1-CHLOROCTADECANE (CASRN 3386-33-2)

BACKGROUND

A Provisional Peer-Reviewed Toxicity Value (PPRTV) is defined as a toxicity value derived for use in the Superfund Program. PPRTVs are derived after a review of the relevant scientific literature using established Agency guidance on human health toxicity value derivations. All PPRTV assessments receive internal review by a standing panel of National Center for Environment Assessment (NCEA) scientists and an independent external peer review by three scientific experts.

The purpose of this document is to provide support for the hazard and dose-response assessment pertaining to chronic and subchronic exposures to substances of concern, to present the major conclusions reached in the hazard identification and derivation of the PPRTVs, and to characterize the overall confidence in these conclusions and toxicity values. It is not intended to be a comprehensive treatise on the chemical or toxicological nature of this substance.

The PPRTV review process provides needed toxicity values in a quick turnaround timeframe while maintaining scientific quality. PPRTV assessments are updated approximately on a 5-year cycle for new data or methodologies that might impact the toxicity values or characterization of potential for adverse human health effects and are revised as appropriate. It is important to utilize the PPRTV database (<http://hhpprtv.ornl.gov>) to obtain the current information available. When a final Integrated Risk Information System (IRIS) assessment is made publicly available on the Internet (<http://www.epa.gov/iris>), the respective PPRTVs are removed from the database.

DISCLAIMERS

The PPRTV document provides toxicity values and information about the adverse effects of the chemical and the evidence on which the value is based, including the strengths and limitations of the data. All users are advised to review the information provided in this document to ensure that the PPRTV used is appropriate for the types of exposures and circumstances at the site in question and the risk management decision that would be supported by the risk assessment.

Other U.S. Environmental Protection Agency (EPA) programs or external parties who may choose to use PPRTVs are advised that Superfund resources will not generally be used to respond to challenges, if any, of PPRTVs used in a context outside of the Superfund program.

QUESTIONS REGARDING PPRTVs

Questions regarding the contents and appropriate use of this PPRTV assessment should be directed to the EPA Office of Research and Development's National Center for Environmental Assessment, Superfund Health Risk Technical Support Center (513-569-7300).

INTRODUCTION

1-Chlorooctadecane (CASRN 3386-33-2) is a high production volume chemical used as a solvent and an intermediate in the manufacture of surfactant, pharmaceuticals, and other organic compounds. The molecular formula of 1-chlorooctadecane is $\text{CH}_3(\text{CH}_2)_{17}\text{Cl}$ (see Figure 1). A list of physicochemical properties is provided in Table 1.

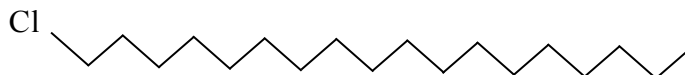


Figure 1. 1-Chlorooctadecane (CASRN 3386-33-2) Structure

| Table 1. Physicochemical Properties of 1-Chlorooctadecane (CASRN 3386-33-2)^a | |
|--|-----------------------|
| Property (Unit) | Value |
| Boiling point (°C) | 348 |
| Melting point (°C) | 28.6 |
| Density (g/cm ³ at 20°C) | ND |
| Vapor pressure (mm Hg at 25°C) | 1.71×10^{-5} |
| pH (unitless) | ND |
| Solubility in water (mg/L at 25°C) | 1.25×10^{-4} |
| Relative vapor density (air = 1) | ND |
| Molecular weight (g/mol) | 288.943 |

^a[ChemIDplus \(2013\)](#).

ND = no data.

Table 2 provides a summary of available toxicity values for 1-chlorooctadecane from EPA and other regulatory agencies or organizations.

**Table 2. Summary of Available Toxicity Values for
1-Chlorooctadecane (CASRN 3386-33-2)**

| Source/Parameter ^{a,b} | Value (Applicability) | Reference |
|---------------------------------|-----------------------|--|
| Noncancer | | |
| ACGIH | NA | ACGIH (2013) |
| ATSDR | NA | ATSDR (2013) |
| Cal/EPA | NA | (Cal/EPA); Cal/EPA (2014)^c |
| NIOSH | NA | NIOSH (2010) |
| OSHA | NA | OSHA (2011); OSHA (2006) |
| IRIS | NA | (U.S. EPA) |
| DWSHA | NA | U.S. EPA (2012a) |
| HEAST | NA | U.S. EPA (2011) |
| CARA HEEP | NA | U.S. EPA (1994) |
| WHO | NA | (WHO) |
| Cancer | | |
| IRIS | NA | (U.S. EPA) |
| HEAST/WOE | NA | U.S. EPA (2011) |
| IARC | NA | (IARC) |
| NTP | NA | NTP (2014) |
| Cal/EPA | NA | (Cal/EPA); Cal/EPA (2015a); Cal/EPA (2011) |

^aSources: ACGIH = American Conference of Governmental Industrial Hygienists; ATSDR = Agency for Toxic Substances and Disease Registry; Cal/EPA = California Environmental Protection Agency; CARA = Chemical Assessments and Related Activities; DWSHA = Drinking Water Standards and Health Advisories; HEAST = Health Effects Assessment Summary Tables; HEEP = Health and Environmental Effects Profile; IARC = International Agency for Research on Cancer; IRIS = Integrated Risk Information System; NIOSH = National Institute for Occupational Safety and Health; NTP = National Toxicology Program; OSHA = Occupational Safety and Health Administration; WHO = World Health Organization.

^bParameters: Cancer weight of evidence (WOE) [\(U.S. EPA, 1986\)](#).

^cThe Cal/EPA Office of Environmental Health Hazard Assessment (OEHHA) Toxicity Criteria Database (<http://oehha.ca.gov/tcdb/index.asp>) was also reviewed and found to contain no information on 1-chlorooctadecane.

NA = not available.

Literature searches were conducted on sources published from 1900 through February 2015 for studies relevant to the derivation of provisional toxicity values for 1-chlorooctadecane (CASRN 3386-33-2). The following databases were searched by chemical name, synonyms, or CASRN: ACGIH, ANEUPL, ATSDR, BIOSIS, Cal/EPA, CCRIS, CDAT, ChemIDplus, CIS, CRISP, DART, EMIC, EPIDEM, ETICBACK, FEDRIP, GENE-TOX, HAPAB, HERO, HMTc, HSDB, IARC, INCHEM IPCS, IPA, ITER, IUCLID, LactMed, NIOSH, NTIS, NTP, OSHA, OPP/RED, PESTAB, PPBIB, PPRTV, PubMed (toxicology subset), RISKLINE, RTECS, TOXLINE, TRI, U.S. EPA IRIS, U.S. EPA HEAST, U.S. EPA HEEP, U.S. EPA OW, and U.S. EPA TSCATS/TSCATS2. The following databases were searched for toxicity values or exposure limits: ACGIH, ATSDR, Cal EPA, U.S. EPA IRIS, U.S. EPA HEAST, U.S. EPA HEEP, U.S. EPA OW, U.S. EPA TSCATS/TSCATS2, NIOSH, NTP, OSHA, and RTECS.

REVIEW OF POTENTIALLY RELEVANT DATA (NONCANCER AND CANCER)

The available data on 1-chlorooctadecane primarily focuses on its biodegradation, biotransformation by marine creatures, incorporation into the fatty acids of microorganisms, and usage in the development of analytical methods. No usable information is available regarding repeated-dose oral or inhalation exposure of humans or animals to 1-chlorooctadecane.

DERIVATION OF PROVISIONAL VALUES

DERIVATION OF ORAL REFERENCE DOSES

Feasibility of Deriving Subchronic and Chronic p-RfDs

No subchronic-duration, chronic-duration, developmental toxicity, reproductive toxicity, or carcinogenicity studies on 1-chlorooctadecane via the oral route were identified. Thus, no oral reference doses could be derived. However, as noted below, a computational toxicological surrogate approach was attempted.

DERIVATION OF INHALATION REFERENCE CONCENTRATIONS

Feasibility of Deriving Subchronic and Chronic Provisional Reference Concentrations (p-RfCs)

No subchronic-duration, chronic-duration, developmental toxicity, reproductive toxicity, or carcinogenicity studies on 1-chlorooctadecane via the inhalation route were identified. Thus, no inhalation reference doses could be derived. However, as noted below, a computational toxicological surrogate approach was attempted.

CANCER WEIGHT-OF-EVIDENCE (WOE) DESCRIPTOR

Limitations in the available data preclude development of a weight-of-evidence (WOE) descriptor.

MODE-OF-ACTION (MOA) DISCUSSION

Limitations in the available data preclude determination of a mode-of-action (MOA) discussion.

ALTERNATIVE METHODS

The surrogate approach allows for the use of data from related compounds to calculate screening values when data for the compound of interest are limited or unavailable. Details regarding searches and methods for surrogate analysis are presented in [Wang et al. \(2012\)](#). Three types of potential surrogates (structural, metabolic, and toxicity) are identified to facilitate the final surrogate chemical selection. The surrogate approach may or may not be route-specific or applicable to multiple routes of exposure. All information was considered together as part of the final weight-of-evidence (WOE) approach to select the most suitable surrogate both toxicologically and chemically.

An initial surrogate search focused on the identification of structurally similar chemicals with toxicity values from the Integrated Risk Information System (IRIS), PPRTV, and Health Effects Assessment Summary Tables (HEAST) databases to take advantage of the well-characterized chemical-class information. This was accomplished by searching the US EPA's DSSTox database ([DSSTox, 2012](#)) at similarity levels >60%, and the National Library of Medicine's ChemIDplus database ([ChemIDplus, 2013](#)) at similarity levels >80%. There were 20 compounds identified in DSSTox and 76 compounds identified in ChemIDplus, for a total of 69 unique compounds. The larger number of compounds identified in ChemIDplus largely reflects its inclusion of other halogen substituted (fluorinated or brominated) compounds. However, there was no in vivo repeated-dose information on any of these related compounds. Due to a lack of repeat-dose toxicity information for any of the potential structural surrogates, derivation of risk values (e.g., RfD, RfC, and oral cancer slope factor) based on the computational toxicological surrogate approach ([Wang et al., 2012](#)) is not feasible for 1-chlorooctadecane.

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