ATRAZINE

APPENDIX A. ATSDR MINIMAL RISK LEVELS AND WORKSHEETS

The Comprehensive Environmental Response, Compensation, and Liability Act (CERCLA) [42 U.S.C. 9601 et seq.], as amended by the Superfund Amendments and Reauthorization Act (SARA) [Pub. L. 99–499], requires that the Agency for Toxic Substances and Disease Registry (ATSDR) develop jointly with the U.S. Environmental Protection Agency (EPA), in order of priority, a list of hazardous substances most commonly found at facilities on the CERCLA National Priorities List (NPL); prepare toxicological profiles for each substance included on the priority list of hazardous substances; and assure the initiation of a research program to fill identified data needs associated with the substances.

The toxicological profiles include an examination, summary, and interpretation of available toxicological information and epidemiologic evaluations of a hazardous substance. During the development of toxicological profiles, Minimal Risk Levels (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 duration of exposure. MRLs are based on noncancer health effects only and are not based on a consideration of cancer effects. 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 no-observed-adverse-effect level/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 and longer) 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 chemical-induced end point 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.

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APPENDIX A

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 a hundredfold 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 Division of Toxicology, 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 levels. For additional information regarding MRLs, please contact the Division of Toxicology, Agency for Toxic Substances and Disease Registry, 1600 Clifton Road NE, Mailstop E-29, Atlanta, Georgia 30333.

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MINIMAL RISK LEVEL WORKSHEET

2003
-Public
on [X] Oral
] Intermediate [] Chronic

Minimal Risk Level: 0.01 [X] mg/kg/day [] ppm

<u>Reference</u>: Infurna R, Levy B, Meng C, et al. 1988. Teratological evaluations of atrazine technical, atrazine herbicide, in rats and rabbits. J Toxicol Environ Health 24:307-319.

<u>Experimental design</u>: Groups of 19 female New Zealand White rabbits were artificially inseminated (gestation day 0) and administered 0, 1, 5, or 75 mg/kg/day atrazine (Aatrex) in 3% aqueous corn starch containing 0.5% Tween 80 by gavage on gestational days 7–19. Rabbits were observed twice daily for changes in appearance and behavior. Feed consumption and body weight changes were monitored throughout gestation. Dams were necropsied on gestational day 29. Ovaries were examined, corpora lutea were counted, uteri and contents were weighed, live fetuses and resorptions were counted, and liver weights were recorded. Fetuses were weighed, sexed, and examined for external, visceral, skeletal, and soft tissue abnormalities.

<u>Effects noted in study and corresponding doses</u>: Clinical signs related to treatment were increased incidence of stool variations (little, no, or soft stool) and bloody vulva. Absolute, but not relative, liver weight was decreased in the 75 mg/kg/day group. Food consumption and body weight gain were severely reduced during the treatment period in the high dose group, but rebounded after cessation of treatment; however, overall body weight gain corrected for weight of the uterus, placentas, and fetuses was significantly reduced. Slight, but statistically significant, reductions in food consumption and body weight gain were noted in the 5 mg/kg/day group.

Dose and end point used for MRL derivation:

[X] NOAEL [] LOAEL 1 mg/kg/day in pregnant rabbits, decreased body weight gain at ≥ 5 mg/kg/day

Uncertainty Factors used in MRL derivation:

- [] 10 for use of a LOAEL
- [X] 10 for extrapolation from animals to humans
- [X] 10 for human variability

Was a conversion used from ppm in food or water to a mg/body weight dose? If so, explain: None needed.

If an inhalation study in animals, list the conversion factors used in determining human equivalent dose: N/A

Other additional studies or pertinent information which lend support to this MRL: The MRL is supported by a number of studies showing decreased body weight gain in rats (Cantemir et al. 1997; Cooper et al. 1996b, 2000; Cummings et al. 2000b; Dési 1983; Eldridge et al. 1994a, 1999a; EPA 1984f, 1987d, 1987e; Infurna et al. 1988; Kniewald et al. 2000; Pinter et al. 1990; Santa Maria et al. 1987; Šimić et al. 1994; Suschetet et al. 1974; Tennant et al. 1994b; Ugazio et al. 1991a; Vos et al. 1983; Wetzel et al. 1994) and dogs (EPA 1987f). Other effects noted in rabbits in the Infurna et al. (1988) study occurred only in the high-dose group (75 mg/kg/day) and included increased incidence of stool variations (little, no, or soft stool), bloody vulva, absolute, but not relative, liver weight decrease, and severely reduced food consumption and body weight gain. Slight, but statistically significant, decreases in body weight gain occurred in the 5 mg/kg/day group. Other NOAELs and LOAELs for acute-duration exposures include: a NOAEL of 12.5 mg/kg/day for increased inflammation of the lateral prostate in adult male offspring of atrazine-treated rat dams (Stoker et al. 1999); a NOAEL of 5 mg/kg/day for increased resorptions/litter and decreased live fetuses/litter in rabbits exposed on gestational days 7-19 (Infurna et al. 1988); and a NOAEL of 1 mg/kg/day for developmental effects (decreased fetal body weight; increased incidence of nonossification of foot bones and patellae) in offspring of treated rabbit dams (Infurna et al. 1988); and a NOAEL of 10 mg/kg/day for developmental effects (incomplete ossification of skull, hyoid bone, teeth, forepaw metacarpals, and hindpaw distal phalanges) in rat offspring of dams exposed to 70 mg/kg/day (Infurna et al. 1988). The developmental effects were attributed to severe maternal toxicity related to severe decreases in food intake and body weight. Changes in serum and pituitary hormone levels have been seen at exposures of \geq 50 mg/kg/day (Cooper et al. 2000; Stoker et al. 1999).

Agency Contact (Chemical Manager): Alfred S. Dorsey Jr., DVM

MINIMAL RISK LEVEL WORKSHEET

Chemical Name:	Atrazine
CAS Number:	1912-24-9
Date:	September 2003
Profile Status:	Draft 3 Post-Public
Route:	[] Inhalation [X] Oral
Duration:	[] Acute [X] Intermediate [] Chronic
Graph Key:	74
Species:	Pig

Minimal Risk Level: 0.003 [X] mg/kg/day [] ppm

<u>Reference</u>: Gojmerac T, Uremovic M, Uremovic S, et al. 1999. Reproduction disturbance caused by an s-triazine herbicide in pigs. Acta Veterinaria Hungarica 47(1):129-135.

Experimental design: Groups of nine female Swedish Landrace/Large Yorkshire cross pigs (6–7-monthold gilts) were administered 0 or 1 mg/kg/day atrazine in the feed for 19 days beginning with the onset of estrus (day 0). Blood samples were drawn 3 times daily at 3-hour intervals on the 5 days immediately following the final day of atrazine administration (this corresponded to the expected day of the next estrus [day 0] and 2 days before [days -1 and -2] and 2 days [days 1 and 2] after the expected estrus). Serum 17β -estradiol (E₂) concentrations in the blood samples were determined. Histopathological examination of the uterus was performed.

Effects noted in study and corresponding doses: E_2 concentrations were statistically significantly different (p<0.001) from controls on all 5 days measured. In controls, E_2 concentrations were high on days -2 and -1, then dropped on day 0 (beginning of estrus) and remained low on days 1 and 2. In treated animals, E_2 concentrations were lower than controls on days -2 and -1, and higher than controls on days 0 through 2. Treated pigs failed to exhibit overt signs of estrus onset and uterine histopathology indicated a state of uterine rest (diestrus) at the end of the observation period. A slight, but steady increase of E_2 hormone level was seen in the treated animals on day 24 of the estrus cycle (day 2). The study authors suggested that the balance of the E_2 hormone level was being gradually restored, which is the pattern that would be anticipated if the animals were about to go into estrus.

Dose and end point used for MRL derivation:

[] NOAEL [X] LOAEL 1 mg/kg/day in pigs, delayed onset of estrus

Uncertainty Factors used in MRL derivation:

- [X] 10 for use of a LOAEL
- [X] 10 for extrapolation from animals to humans
- [X] 3 for human variability

An uncertainty factor of 3 for human variability was used instead of 10 because the critical effect was identified in a sensitive population (young, developing female pigs).

Was a conversion used from ppm in food or water to a mg/body weight dose? If so, explain: N/A

If an inhalation study in animals, list the conversion factors used in determining human equivalent dose: $N\!/\!A$

Other additional studies or pertinent information which lend support to this MRL: Other systemic and reproductive effects have been observed in animals exposed to atrazine for 15–365 days. Decreased body weight gain was seen in rats at LOAELs of 2.7 mg/kg/day and above (Cantemir et al. 1997; Cooper et al. 1996b, 2000; Dési 1983; Eldridge et al. 1994a, 1999a; Kniewald et al. 2000; Laws et al. 2000; Suschetet et al. 1974; Trentacoste et al. 2001; Wetzel et al. 1994). Endocrine gland weights and serum and pituitary gland hormone levels were altered in rats at LOAELs as low as 6.9 mg/kg/day for 1 month (Cooper et al. 1996b, 2000; Eldridge et al. 1994a; Friedmann 2002; Laws et al. 2000; Trentacoste et al. 2001; Wetzel et al. 1994). Disrupted estrus cyclicity was seen in rats at the lowest LOAEL of 6.9 mg/kg/day (Wetzel et al. 1994). Other NOAELs and LOAELs observed included: a LOAEL of 50 mg/kg/day (NOAEL of 5 mg/kg/day) for increased relative liver weights in Sprague-Dawley and Donryu rats (Aso et al. 2000); a LOAEL of 2 mg/kg/day for degeneration of a small number of myocardial fibers, mild degeneration and inflammation and mild chronic interstitial hepatitis and subacute glomerulitis, degeneration and desquamation of proximal tubules, a 350% increase in serum gamma-glutamyl-transferase, and mild liver histological changes in pigs (Ćurić et al. 1999; Gojmerac et al. 1995); a LOAEL of 33 mg/kg/day (NOAEL of 4.6 mg/kg/day) for abnormal estrus cycle in Sprague-Dawley rats (Eldridge et al. 1999a); and a LOAEL of 2 mg/kg/day for ovarian histopathology, disrupted estrogen and progesterone levels, and delaved onset of estrus (Gojmerac et al. 1996), and ovarian cysts and disruption of estrus cyclicity (Ćurić et al. 1999).

Agency Contact (Chemical Manager): Alfred S. Dorsey Jr., DVM

APPENDIX B. USER'S GUIDE

Chapter 1

Public Health Statement

This chapter of the profile is a health effects summary written in non-technical language. Its intended audience is the general public especially people living in the vicinity of a hazardous waste site or chemical release. If the Public Health Statement were removed from the rest of the document, it would still communicate to the lay public essential information about the chemical.

The major headings in the Public Health Statement are useful to find specific topics of concern. The topics are written in a question and answer format. The answer to each question includes a sentence that will direct the reader to chapters in the profile that will provide more information on the given topic.

Chapter 2

Relevance to Public Health

This chapter provides a health effects summary based on evaluations of existing toxicologic, epidemiologic, and toxicokinetic information. This summary is designed to present interpretive, weight-of-evidence discussions for human health end points 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?

The chapter covers end points in the same order they appear within the Discussion of Health Effects by Route of Exposure section, by route (inhalation, oral, dermal) and within route by effect. Human data are presented first, then animal data. Both are organized by duration (acute, intermediate, chronic). *In vitro* data and data from parenteral routes (intramuscular, intravenous, subcutaneous, etc.) are also considered in this chapter. If data are located in the scientific literature, a table of genotoxicity information is included.

The carcinogenic potential of the profiled substance is qualitatively evaluated, when appropriate, using existing toxicokinetic, genotoxic, and carcinogenic data. ATSDR does not currently assess cancer potency or perform cancer risk assessments. Minimal risk levels (MRLs) for noncancer end points (if derived) and the end points from which they were derived are indicated and discussed.

Limitations to existing scientific literature that prevent a satisfactory evaluation of the relevance to public health are identified in the Chapter 3 Data Needs section.

Interpretation of Minimal Risk Levels

Where sufficient toxicologic information is available, we have derived minimal risk levels (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.

They should help physicians and public health officials determine the safety of a community living near a chemical 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. Chapter 2, "Relevance to Public Health," contains basic information known about the substance. Other sections such as Chapter 3 Section 3.9, "Interactions with Other Substances," and Section 3.10, "Populations that are Unusually Susceptible" 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 the Environmental Protection Agency (EPA) provides (Barnes and Dourson 1988) to determine reference doses for lifetime exposure (RfDs).

To derive an MRL, ATSDR generally selects the most sensitive end point 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 end point are available, ATSDR derives an MRL using the most sensitive species (when information from multiple species is available) with the highest 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 (UF) 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 LSE Tables.

Chapter 3

Health Effects

Tables and Figures for Levels of Significant Exposure (LSE)

Tables (3-1, 3-2, and 3-3) and figures (3-1 and 3-2) 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, minimal risk levels (MRLs) to humans for noncancer end points, and EPA's estimated range associated with an upper- bound individual lifetime cancer risk of 1 in 10,000 to 1 in 10,000,000. Use the LSE tables and figures for a quick review of the health effects and to locate data for a specific exposure scenario. The LSE tables and figures represent studies that provide reliable, quantitative estimates of No-Observed-Adverse-Effect Levels (NOAELs), Lowest-Observed-Adverse-Effect Levels (LOAELs), or Cancer Effect Levels (CELs).

The legends presented below demonstrate the application of these tables and figures. Representative examples of LSE Table 3-1 and Figure 3-1 are shown. The numbers in the left column of the legends correspond to the numbers in the example table and figure.

LEGEND

See LSE Table 3-1

- (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. When sufficient data exists, 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 Table 3-1, 3-2, and 3-3, respectively). LSE figures are limited to the inhalation (LSE Figure 3-1) and oral (LSE Figure 3-2) routes. Not all substances will have data on each route of exposure and will not therefore have all five of the tables and figures.
- (2) Exposure Period Three exposure periods acute (less than 15 days), intermediate (15–364 days), and chronic (365 days or more) are presented within each relevant route of exposure. In this example, an inhalation study of intermediate exposure duration is 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>Health Effect</u> The major categories of health effects included in LSE tables and figures are death, systemic, immunological, neurological, developmental, reproductive, and cancer. NOAELs and LOAELs can be reported in the tables and figures for all effects but cancer. Systemic effects are further defined in the "System" column of the LSE table (see key number 18).
- (4) <u>Key to Figure</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 18 has been used to derive a NOAEL and a Less Serious LOAEL (also see the 2 "18r" data points in Figure 3-1).
- (5) <u>Species</u> The test species, whether animal or human, are identified in this column. Chapter 2, "Relevance to Public Health," covers the relevance of animal data to human toxicity and Section 3.4, "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.
- (6) Exposure Frequency/Duration The duration of the study and the weekly and daily exposure regimen are provided in this column. This permits comparison of NOAELs and LOAELs from different studies. In this case (key number 18), rats were exposed to 1,1,2,2-tetrachloroethane via inhalation for 6 hours per day, 5 days per week, for 3 weeks. For a more complete review of the dosing regimen refer to the appropriate sections of the text or the original reference paper, i.e., Nitschke et al. 1981.
- (7) <u>System</u> This column further defines the systemic effects. These systems include: respiratory, cardiovascular, gastrointestinal, hematological, musculoskeletal, hepatic, renal, and dermal/ocular. "Other" refers to any systemic effect (e.g., a decrease in body weight) not covered in these systems. In the example of key number 18, 1 systemic effect (respiratory) was investigated.

- (8) <u>NOAEL</u> A No-Observed-Adverse-Effect Level (NOAEL) is the highest exposure level at which no harmful effects were seen in the organ system studied. Key number 18 reports a NOAEL of 3 ppm for the respiratory system which was used to derive an intermediate exposure, inhalation MRL of 0.005 ppm (see footnote "b").
- (9) <u>LOAEL</u> A Lowest-Observed-Adverse-Effect Level (LOAEL) is the lowest dose used in the study that caused a harmful 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 end point used to quantify the adverse effect accompanies the LOAEL. The respiratory effect reported in key number 18 (hyperplasia) is a Less serious LOAEL of 10 ppm. MRLs are not derived from Serious LOAELs.
- (10) <u>Reference</u> The complete reference citation is given in Chapter 9 of the profile.
- (11) <u>CEL</u> 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.
- (12) <u>Footnotes</u> Explanations of abbreviations or reference notes for data in the LSE tables are found in the footnotes. Footnote "b" indicates the NOAEL of 3 ppm in key number 18 was used to derive an MRL of 0.005 ppm.

LEGEND

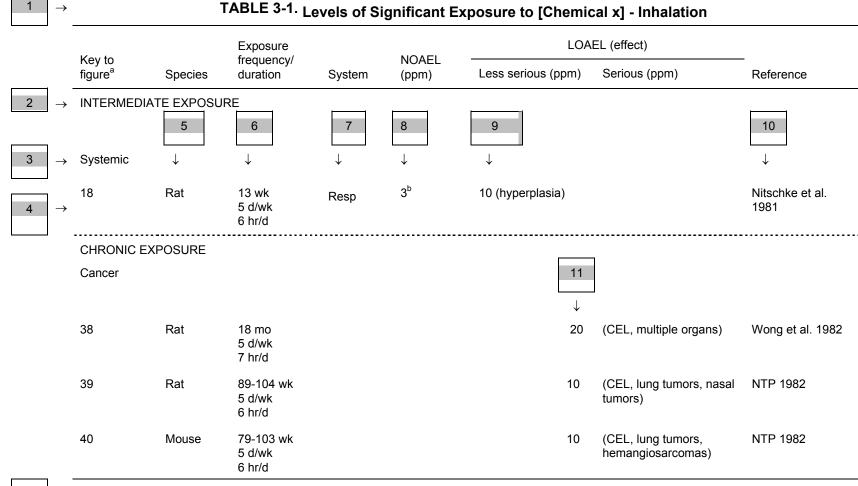
See Figure 3-1

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 intermediate and chronic exposure periods are illustrated.
- (14) <u>Health Effect</u> These are the categories of health effects for which reliable quantitative data exists. The same health effects 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/m³ or ppm and oral exposure is reported in mg/kg/day.
- (16) <u>NOAEL</u> In this example, the open circle designated 18r identifies a NOAEL critical end point in the rat upon which an intermediate inhalation exposure MRL is based. The key number 18 corresponds to the entry in the LSE table. The dashed descending arrow indicates the extrapolation from the exposure level of 3 ppm (see entry 18 in the Table) to the MRL of 0.005 ppm (see footnote "b" in the LSE table).

- (17) <u>CEL</u> Key number 38r is 1 of 3 studies for which Cancer Effect Levels were derived. The diamond symbol refers to a Cancer Effect Level for the test species-mouse. The number 38 corresponds to the entry in the LSE table.
- (18) Estimated Upper-Bound Human Cancer Risk Levels This is the range associated with the upperbound for lifetime cancer risk of 1 in 10,000 to 1 in 10,000,000. These risk levels are derived from the EPA's Human Health Assessment Group's upper-bound estimates of the slope of the cancer dose response curve at low dose levels (q_1^*) .
- (19) <u>Key to LSE Figure</u> The Key explains the abbreviations and symbols used in the figure.

SAMPLE

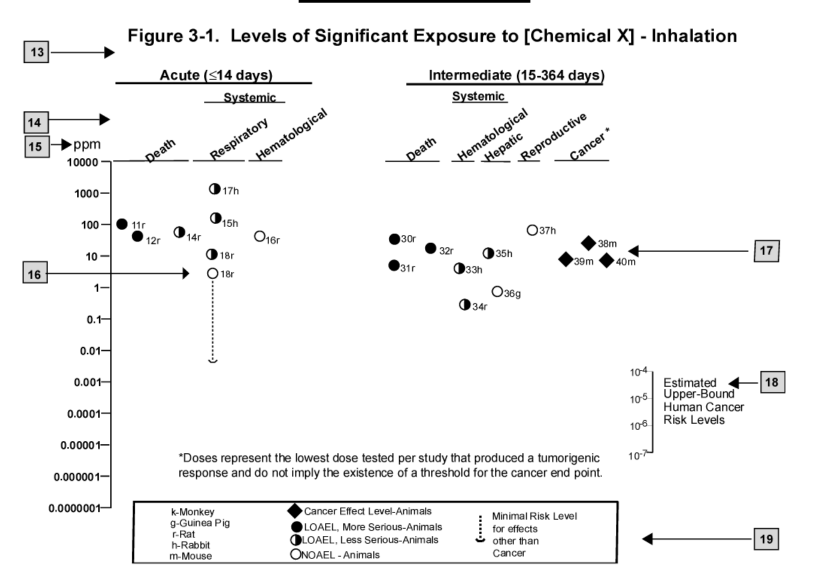


 \rightarrow ^a The number corresponds to entries in Figure 3-1.

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Used to derive an intermediate inhalation Minimal Risk Level (MRL) of 5 x 10⁻³ ppm; dose adjusted for intermittent exposure and divided by an uncertainty factor of 100 (10 for extrapolation from animal to humans, 10 for human variability).

SAMPLE



APPENDIX C. ACRONYMS, ABBREVIATIONS, AND SYMBOLS

ACOEM	American Callege of Occurational and Environmental Medicine		
ACOEM	American College of Occupational and Environmental Medicine		
ACGIH	American Conference of Governmental Industrial Hygienists		
ADI	acceptable daily intake		
ADME	absorption, distribution, metabolism, and excretion		
AED	atomic emission detection		
AOEC	Association of Occupational and Environmental Clinics		
AFID	alkali flame ionization detector		
AFOSH	Air Force Office of Safety and Health		
ALT	alanine aminotransferase		
AML	acute myeloid leukemia		
AOAC	Association of Official Analytical Chemists		
AP	alkaline phosphatase		
APHA	American Public Health Association		
AST	aspartate aminotranferase		
atm	atmosphere		
ATSDR	Agency for Toxic Substances and Disease Registry		
AWQC	Ambient Water Quality Criteria		
BAT	best available technology		
BCF	bioconcentration factor		
BEI	Biological Exposure Index		
BSC	Board of Scientific Counselors		
С	centigrade		
CAA	Clean Air Act		
CAG	Cancer Assessment Group of the U.S. Environmental Protection Agency		
CAS	Chemical Abstract Services		
CDC	Centers for Disease Control and Prevention		
CEL	cancer effect level		
CELDS	Computer-Environmental Legislative Data System		
CERCLA	Comprehensive Environmental Response, Compensation, and Liability Act		
CFR	Code of Federal Regulations		
Ci	curie		
CI	confidence interval		
CL	ceiling limit value		
CLP	Contract Laboratory Program		
cm	centimeter		
CML	chronic myeloid leukemia		
CPSC	Consumer Products Safety Commission		
CWA	Clean Water Act		
DHEW	Department of Health, Education, and Welfare		
DHHS	Department of Health and Human Services		
DNA	deoxyribonucleic acid		
DOD	Department of Defense		
DOE	Department of Energy		
DOL	Department of Labor		
DOT	Department of Transportation		
DOT/UN/	Department of Transportation/United Nations/		
NA/IMCO	North America/International Maritime Dangerous Goods Code		
DWEL	drinking water exposure level		

ECD	alastron contura dataction
	electron capture detection
ECG/EKG	electrocardiogram
EEG	electroencephalogram
EEGL	Emergency Exposure Guidance Level
EPA	Environmental Protection Agency
F	Fahrenheit
F_1	first-filial generation
FAO	Food and Agricultural Organization of the United Nations
FDA	Food and Drug Administration
FEMA	Federal Emergency Management Agency
FIFRA	Federal Insecticide, Fungicide, and Rodenticide Act
FPD	flame photometric detection
fpm	feet per minute
FR	Federal Register
FSH	follicle stimulating hormone
g	gram
GC	gas chromatography
gd	gestational day
GLC	gas liquid chromatography
GPC	gel permeation chromatography
HPLC	high-performance liquid chromatography
HRGC	high resolution gas chromatography
HSDB	Hazardous Substance Data Bank
IARC	International Agency for Research on Cancer
IDLH	immediately dangerous to life and health
ILO	International Labor Organization
IRIS	Integrated Risk Information System
Kd	adsorption ratio
kg	kilogram
K _{oc}	organic carbon partition coefficient
K _{ow}	octanol-water partition coefficient
L	liter
LC	liquid chromatography
LC_{Lo}	lethal concentration, low
LC_{50}	lethal concentration, 50% kill
LD_{Lo}	lethal dose, low
LD_{50}	lethal dose, 50% kill
LDH	lactic dehydrogenase
LH	luteinizing hormone
LT_{50}	lethal time, 50% kill
LOAEL	lowest-observed-adverse-effect level
LSE	Levels of Significant Exposure
m	meter
MA	trans, trans-muconic acid
MAL	maximum allowable level
mCi	millicurie
MCL	maximum contaminant level
MCLG	maximum contaminant level goal
MFO	mixed function oxidase
mg	milligram
mL	milliliter

	millimeter	
mm		
mmHg	millimeters of mercury	
mmol	millimole	
mppcf	millions of particles per cubic foot	
MRL	Minimal Risk Level	
MS	mass spectrometry	
NAAQS	National Ambient Air Quality Standard	
NAS	National Academy of Science	
NATICH	National Air Toxics Information Clearinghouse	
NATO	North Atlantic Treaty Organization	
NCE	normochromatic erythrocytes	
NCEH	National Center for Environmental Health	
NCI	National Cancer Institute	
ND	not detected	
NFPA	National Fire Protection Association	
ng	nanogram	
NIEHS	National Institute of Environmental Health Sciences	
NIOSH	National Institute for Occupational Safety and Health	
NIOSHTIC	NIOSH's Computerized Information Retrieval System	
NLM	National Library of Medicine	
nm	nanometer	
NHANES	National Health and Nutrition Examination Survey	
nmol	nanomole	
NOAEL	no-observed-adverse-effect level	
NOES	National Occupational Exposure Survey	
NOHS	National Occupational Hazard Survey	
NPD	nitrogen phosphorus detection	
NPDES	National Pollutant Discharge Elimination System	
NPL	National Priorities List	
NR	not reported	
NRC	National Research Council	
NS	not specified	
NSPS	New Source Performance Standards	
NTIS	National Technical Information Service	
NTP	National Toxicology Program	
ODW	Office of Drinking Water, EPA	
OERR	Office of Emergency and Remedial Response, EPA	
OHM/TADS	Oil and Hazardous Materials/Technical Assistance Data System	
OPP	Office of Pesticide Programs, EPA	
OPPTS	Office of Prevention, Pesticides and Toxic Substances, EPA	
OPPT	Office of Pollution Prevention and Toxics, EPA	
OR	odds ratio	
OSHA	Occupational Safety and Health Administration	
OSW	Office of Solid Waste, EPA	
OW	Office of Water	
OWRS	Office of Water Regulations and Standards, EPA	
РАН	polycyclic aromatic hydrocarbon	
PBPD	physiologically based pharmacodynamic	
PBPK	physiologically based pharmacokinetic	
PCE	polychromatic erythrocytes	
PEL	permissible exposure limit	

pg	picogram	
PB PHS	Public Health Service	
PID	photo ionization detector	
pmol	picomole	
PMR	proportionate mortality ratio	
ppb	parts per billion	
ppm	parts per million	
ppt	parts per trillion	
PSNS	pretreatment standards for new sources	
RBC	red blood cell	
REL	recommended exposure level/limit	
RfC	reference concentration	
RfD	reference dose	
RNA	ribonucleic acid	
RTECS	Registry of Toxic Effects of Chemical Substances	
RQ	reportable quantity	
SARA	Superfund Amendments and Reauthorization Act	
SCE	sister chromatid exchange	
SGOT	serum glutamic oxaloacetic transaminase	
SGPT	serum glutamic pyruvic transaminase	
SIC	standard industrial classification	
SIM	selected ion monitoring	
SMCL	secondary maximum contaminant level	
SMR	standardized mortality ratio	
SNARL	suggested no adverse response level	
SPEGL	Short-Term Public Emergency Guidance Level	
STEL	short term exposure limit	
STORET	Storage and Retrieval	
TD_{50}	toxic dose, 50% specific toxic effect	
TLV	threshold limit value	
TOC	total organic carbon	
TPQ	threshold planning quantity	
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	
VOC	volatile organic compound	
WBC	white blood cell	
WHO	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
q_1^*	cancer slope factor
_	negative
+	positive
(+)	weakly positive result
(-)	weakly negative result