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IMMEDIATELY DANGEROUS TO LIFE OR HEALTH (IDLH) VALUE PROFILE

FOR

METHYL ISOCYANATE

[CAS No. 624-83-9]

Department of Health and Human Services
Centers for Disease Control and Prevention
National Institute for Occupational Safety and Health

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1 Foreword

2
3 Chemicals are a ubiquitous component of the modern workplace. Occupational exposures to chemicals have the
4 potential to adversely affect the health and lives of workers. Acute or short-term exposures to high concentrations
5 of some airborne chemicals have the ability to quickly overwhelm workers, resulting in a spectrum of undesirable
6 health outcomes that may inhibit the ability to escape from the exposure environment (e.g., irritation of the eyes
7 and respiratory tract or cognitive impairment), cause severe irreversible effects (e.g., damage to the respiratory
8 tract or reproductive toxicity), and in extreme cases, cause death. Airborne concentrations of chemicals capable
9 of causing such adverse health effects or of impeding escape from high-risk conditions may arise from a variety of
10 non-routine workplace situations, including special work procedures (e.g., in confined spaces), industrial
11 accidents (e.g., chemical spills or explosions), and chemical releases into the community (e.g., during
12 transportation incidents or other uncontrolled-release scenarios).

13
14 The “immediately dangerous to life or health air concentration values (IDLH values)” developed by the National
15 Institute for Occupational Safety and Health (NIOSH) characterize these high-risk exposure concentrations and
16 conditions [NIOSH 2013]. IDLH values are based on a 30-minute exposure duration and have traditionally
17 served as a key component of the decision logic for the selection of respiratory protection devices [NIOSH 2004].
18 In addition, occupational health professionals have employed these values beyond their initial purpose as a
19 component of the NIOSH Respirator Selection Logic to assist in developing Risk Management Plans for non-
20 routine work practices governing operations in high-risk environments (e.g., confined spaces) and the
21 development of Emergency Preparedness Plans.

22
23 The approach used to derive IDLH values for high priority chemicals is outlined in the NIOSH Current
24 Intelligence Bulletin (CIB) 66: Derivation of Immediately Dangerous to Life or Health Values [NIOSH 2013].
25 CIB 66 provides 1) an update on the scientific basis and risk assessment methodology used to derive IDLH
26 values, 2) the rationale and derivation process for IDLH values, and 3) a demonstration of the derivation of
27 scientifically credible IDLH values using available data resources.

28
29 The purpose of this technical report is to present the IDLH value for methyl isocyanate (CAS # 624-83-9). The
30 scientific basis, toxicologic data and risk assessment approach used to derive the IDLH value are summarized to
31 ensure transparency and scientific credibility.

32
33 John Howard, M.D.
34 Director
35 National Institute for Occupational Safety and Health

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1 **Abbreviations**

2

3	ACGIH	American Conference of Governmental Industrial Hygienists
4	AEGL	Acute Exposure Guideline Levels
5	AIHA	American Industrial Hygiene Association
6	BMC	benchmark concentration
7	BMCL	benchmark concentration lower confidence limit
8	C	ceiling
9	CAS	chemical abstract service
10	ERPG	Emergency Response Planning Guidelines
11	GD	gestation day
12	IDLH	immediately dangerous to life or health
13	LC ₅₀	median lethal concentration
14	LC _{Lo}	lowest concentration of a chemical that caused death in humans or animals
15	LEL	lower explosive limit
16	LOAEL	lowest observed adverse effect level
17	mg/m ³	milligram(s) per cubic meter
18	NAC	National Advisory Committee
19	NAS	National Academy of Sciences
20	NIOSH	National Institute for Occupational Safety and Health
21	NOAEL	no observed adverse effect level
22	OSHA	Occupational Safety and Health Administration
23	PEL	permissible exposure limit
24	ppm	parts per million
25	RD ₅₀	concentration of a chemical in the air that is estimated to cause a 50% decrease in the respiratory
26		rate
27	REL	recommend exposure limit
28	SCP	Standard Completion Program
29	STEL	short term exposure limit
30	TLV	threshold limit value
31	TWA	time weighted average
32	UEL	upper explosive limit
33	WEEL	workplace environmental exposure level

1 **Glossary**

2
3 **Acute Exposure:** Exposure by the oral, dermal, or inhalation route for 24 hours or less.

4 **Acute Exposure Guideline Levels (AEGLs):** Threshold exposure limits for the general public applicable to
5 emergency exposure periods ranging from 10 minutes to 8 hours. AEGL-1, AEGL 2, and AEGL-3 are
6 developed for five exposure periods (10 and 30 minutes, 1 hour, 4 hours, and 8 hours) and are distinguished
7 by varying degrees of severity of toxic effects ranging from transient, reversible effects to life-threatening
8 effects [NAS 2001]. AEGLs are intended to be guideline levels used during rare events or single once-in-a-
9 lifetime exposures to airborne concentrations of acutely toxic, high-priority chemicals [NAS 2001]. The
10 threshold exposure limits are designed to protect the general population, including the elderly, children or
11 other potentially sensitive groups that are generally not considered in the development of workplace exposure
12 recommendations (additional information available at <http://www.epa.gov/oppt/aegl/>).

13 **Acute Reference Concentration (RfC):** An estimate (with uncertainty spanning perhaps an order of magnitude)
14 of a continuous inhalation exposure for an acute duration (24 hours or less) of the human population
15 (including sensitive subgroups) that is likely to be without an appreciable risk of deleterious effects during a
16 lifetime. It can be derived from a NOAEL, LOAEL, or benchmark concentration, with uncertainty factors
17 (UFs) generally applied to reflect limitations of the data used. Generally used in USEPA noncancer health
18 assessments [USEPA 2014].

19 **Acute Toxicity:** Any poisonous effect produced within a short period of time following an exposure, usually 24
20 to 96 hours.

21 **Adverse Effect:** A substance-related biochemical change, functional impairment, or pathologic lesion that affects
22 the performance of an organ or system or alters the ability to respond to additional environmental challenges.

23 **Benchmark Dose/Concentration (BMD/BMC):** A dose or concentration that produces a predetermined change
24 in response rate of an effect (called the benchmark response, or BMR) compared to background [USEPA
25 2014] (additional information available at <http://www.epa.gov/ncea/bmds/>).

26 **Benchmark Response (BMR):** A predetermined change in response rate of an effect. Common defaults for the
27 BMR are 10% or 5%, reflecting study design, data variability, and sensitivity limits used.

28 **BMCL:** A statistical lower confidence limit on the concentration at the BMC [USEPA 2014].

29 **Bolus Exposure:** A single, relatively large dose.

30 **Ceiling Value (“C”):** U.S. term in occupational exposure indicating the airborne concentration of a potentially
31 toxic substance that should never be exceeded in a worker’s breathing zone.

32 **Chronic Exposure:** Repeated exposure for an extended period of time. Typically exposures are more than
33 approximately 10% of life span for humans and >90 days to 2 years for laboratory species.

34 **Critical Study:** The study that contributes most significantly to the qualitative and quantitative assessment of risk
35 [USEPA 2014].
36

37 **Dose:** The amount of a substance available for interactions with metabolic processes or biologically significant
38 receptors after crossing the outer boundary of an organism [USEPA 2014].

39 **EC₅₀:** A combination of the effective concentration of a substance in the air and the exposure duration that is
40 predicted to cause an effect in 50% (one half) of the experimental test subjects.

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1 **Emergency Response Planning Guidelines (ERPGs):** Maximum airborne concentrations below which nearly all
2 individuals can be exposed without experiencing health effects for 1-hour exposure. ERPGs are presented in a
3 tiered fashion with health effects ranging from mild or transient to serious, irreversible, or life threatening
4 (depending on the tier). ERPGs are developed by the American Industrial Hygiene Association [AIHA 2006].

5 **Endpoint:** An observable or measurable biological event or sign of toxicity ranging from biomarkers of initial
6 response to gross manifestations of clinical toxicity.

7 **Exposure:** Contact made between a chemical, physical, or biological agent and the outer boundary of an
8 organism. Exposure is quantified as the amount of an agent available at the exchange boundaries of the
9 organism (e.g., skin, lungs, gut).

10 **Extrapolation:** An estimate of the response at a point outside the range of the experimental data, generally
11 through the use of a mathematical model, although qualitative extrapolation may also be conducted. The
12 model may then be used to extrapolate to response levels that cannot be directly observed.

13 **Hazard:** A potential source of harm. Hazard is distinguished from risk, which is the probability of harm under
14 specific exposure conditions.

15 **Immediately Dangerous to Life or Health (IDLH) condition:** A situation that poses a threat of exposure to
16 airborne contaminants when that exposure is likely to cause death or immediate or delayed permanent adverse
17 health effects or prevent escape from such an environment [NIOSH 2004, 2013].

18 **IDLH value:** A maximum (airborne concentration) level above which only a highly reliable breathing apparatus
19 providing maximum worker protection is permitted [NIOSH 2004, 2013]. IDLH values are based on a 30-
20 minute exposure duration.

21 **LC₀₁:** The statistically determined concentration of a substance in the air that is estimated to cause death in 1% of
22 the test animals.

23 **LC₅₀:** The statistically determined concentration of a substance in the air that is estimated to cause death in 50%
24 (one half) of the test animals; median lethal concentration.

25 **LC₁₀:** The lowest lethal concentration of a substance in the air reported to cause death, usually for a small
26 percentage of the test animals.

27
28 **LD₅₀:** The statistically determined lethal dose of a substance that is estimated to cause death in 50% (one half) of
29 the test animals; median lethal concentration.

30 **LD₁₀:** The lowest dose of a substance that causes death, usually for a small percentage of the test animals.

31 **LEL:** The minimum concentration of a gas or vapor in air, below which propagation of a flame does not occur in
32 the presence of an ignition source.

33 **Lethality:** Pertaining to or causing death; fatal; referring to the deaths resulting from acute toxicity studies. May
34 also be used in lethality threshold to describe the point of sufficient substance concentration to begin to cause
35 death.

36 **Lowest Observed Adverse Effect Level (LOAEL):** The lowest tested dose or concentration of a substance that
37 has been reported to cause harmful (adverse) health effects in people or animals.

38 **Mode of Action:** The sequence of significant events and processes that describes how a substance causes a toxic
39 outcome. Mode of action is distinguished from the more detailed mechanism of action, which implies a more
40 detailed understanding on a molecular level.

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- 1 **No Observed Adverse Effect Level (NOAEL):** The highest tested dose or concentration of a substance that has
2 been reported to cause no harmful (adverse) health effects in people or animals.
- 3 **Occupational Exposure Limit (OEL):** Workplace exposure recommendations developed by governmental
4 agencies and non-governmental organizations. OELs are intended to represent the maximum airborne
5 concentrations of a chemical substance below which workplace exposures should not cause adverse health
6 effects. OELs may apply to ceiling, short-term (STELs), or time-weighted average (TWA) limits.
- 7 **Peak Concentration:** Highest concentration of a substance recorded during a certain period of observation.
- 8 **Permissible Exposure Limit (PEL):** Occupational exposure limits developed by OSHA (29 CFR 1910.1000) or
9 MSHA (30 CFR 57.5001) for allowable occupational airborne exposure concentrations. PELs are legally
10 enforceable and may be designated as ceiling, STEL, or TWA limits.
- 11
- 12 **Point of Departure (POD):** The point on the dose–response curve from which dose extrapolation is initiated.
13 This point can be the lower bound on dose for an estimated incidence or a change in response level from a
14 concentration-response model (BMC), or it can be a NOAEL or LOAEL for an observed effect selected from
15 a dose evaluated in a health effects or toxicology study.
- 16 **RD₅₀:** The statistically determined concentration of a substance in the air that is estimated to cause a 50% (one
17 half) decrease in the respiratory rate.
- 18 **Recommended Exposure Limit (REL):** Recommended maximum exposure limit to prevent adverse health
19 effects based on human and animal studies and established for occupational (up to 10-hour shift, 40-hour
20 week) inhalation exposure by NIOSH. RELs may be designated as ceiling, STEL, or TWA limits.
- 21 **Short-Term Exposure Limit (STEL):** A worker’s 15-minute time-weighted average exposure concentration that
22 shall not be exceeded at any time during a work day.
- 23 **Target Organ:** Organ in which the toxic injury manifests in terms of dysfunction or overt disease.
- 24 **Threshold Limit Values (TLVs®):** Recommended guidelines for occupational exposure to airborne
25 contaminants, published by the American Conference of Governmental Industrial Hygienists (ACGIH). TLVs
26 refer to airborne concentrations of chemical substances and represent conditions under which it is believed
27 that nearly all workers may be repeatedly exposed, day after day, over a working lifetime, without adverse
28 effects. TLVs may be designated as ceiling, short-term (STELs), or 8-hr TWA limits.
- 29 **Time-Weighted Average (TWA):** A worker’s 8-hour (or up to 10-hour) time-weighted average exposure
30 concentration that shall not be exceeded during an 8-hour (or up to 10-hour) work shift of a 40-hour week.
31 The average concentration is weighted to take into account the duration of different exposure concentrations.
- 32 **Toxicity:** The degree to which a substance is able to cause an adverse effect on an exposed organism.
- 33
- 34 **Uncertainty Factors (UFs):** Mathematical adjustments applied to the POD when developing IDLH values. The
35 UFs for IDLH value derivation are determined by considering the study and effect used for the POD, with
36 further modification based on the overall database.
- 37 **Workplace Environmental Exposure Levels (WEELs):** Exposure levels developed by the American Industrial
38 Hygiene Association (AIHA) that provide guidance for protecting most workers from adverse health
39 effects related to occupational chemical exposures expressed as a TWA or ceiling limit.

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1 **Acknowledgments**

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8 **Education and Information Division**

9 Devin Baker, M.Ed.
10 Charles L. Geraci, Ph.D.
11 Thomas J. Lentz, Ph.D.
12 Richard Niemeier, Ph.D.
13 Chris Sofge, Ph.D.

14
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17
18 Michael S. Bisesi, Ph.D., R.E.H.S., C.I.H., Senior Associate Dean for Academic Affairs; Director, Center
19 for Public Health Practice; Interim Chair & Associate Professor, Division of Environmental Health
20 Science, College of Public Health, Ohio State University

21
22 Richard B. Schlesinger, Ph.D., Fellow A.T.S., Senior Associate Dean for Academic Affairs and Research
23 Professor of Biology, Dyson College of Arts and Sciences, Pace University

24
25

1.0 Introduction

1.1 IDLH Value for Methyl Isocyanate

IDLH Value: 0.12 ppm (0.28 mg/m³)

Basis for IDLH Value: The IDLH value for methyl isocyanate is based on the LOAEL for severe reproductive and developmental effects, specifically fetal resorption and fetal skeletal malformations. Varma et al. [1987] reported a LOAEL of 2 ppm for these developmental and reproductive effects in dams exposed for 3 hours on gestation day (GD) 8. This represents a one-time acute exposure at a critical developmental time of the dam and fetuses. The LOAEL was adjusted to a 30-minute equivalent duration concentration of 3.6 ppm. A composite uncertainty factor of 30 was applied to account for extrapolation from a concentration that causes severe effects in animals, animal to human differences, human variability, and uncertainty about the threshold for escape-impairing effects, yielding an IDLH value of **0.12 ppm**.

1.2 Purpose

This *IDLH Value Profile* presents (1) a brief summary of technical data associated with acute inhalation exposures to acrylonitrile and (2) the rationale behind the Immediately Dangerous to Life or Health (IDLH) value for methyl isocyanate. IDLH values are developed based on the scientific rationale and logic outlined in the NIOSH Current Intelligence Bulletin (CIB) 66: Derivation of Immediately Dangerous to Life or Health (IDLH) values [NIOSH 2013]. As described in CIB 66, NIOSH performs in-depth literature searches to ensure that all relevant data from human and animal studies with acute exposures to the substance are identified. Information included in CIB 66 on the literature search includes pertinent databases, key terms, and guides for evaluating data quality and relevance for the establishment of an IDLH value. The information that is identified in the in-depth literature search is evaluated with general considerations that include description of studies (i.e., species, study protocol, exposure concentration and duration), health endpoint evaluated, and critical effect levels (e.g., NOAELs, LOAELs, LC₅₀ values). For methyl isocyanate, the in-depth literature search was conducted through February 2014.

1.3 General Substance Information

Chemical: Methyl Isocyanate

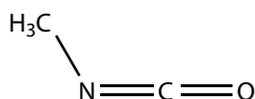
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1 **Synonyms:** Methyl ester isocyanic acid; Isocyanato-methane; MIC [Isocyanate]*

2 **Chemical category:** Organic isocyanates†

3 **Structural formula:**



8 Table 1 highlights selected physiochemical properties of methyl isocyanate relevant to IDLH conditions. Table 2
9 provides alternative exposure guidelines for methyl isocyanate. Table 3 summarizes the Acute Exposure
10 Guidelines Level (AEGL) values for methyl isocyanate.

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Table 1: Physiochemical Properties of Methyl Isocyanate

Property	Value
Molecular weight	57.06‡
Chemical formula	C ₂ H ₃ NO
Description	Colorless liquid
Odor	Sharp, unpleasant
Odor Threshold	2.1 ppm ⁺
UEL	26% [†]
LEL	5.3% [†]
Vapor pressure	348 mmHg at 20°C (68°F) [‡]
Flash point	-7°C (19°F) [‡] - closed cup
Ignition temperature	534°C (994°F) [‡]
Solubility	Hydrolysis [†]

15 **Abbreviation:** °C – Celsius; °F – Fahrenheit; mmHg – millimeter mercury; LEL – lower explosive limit; UEL – upper explosive limit

16 * NLM [2012]

17 † AIHA [1989]

18 ‡ IFA [2012]

19 ‡ HSDB [2012]

20 § AIHA [1989]

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1 **Table 2: Alternative Exposure Guidelines for Methyl Isocyanate**

2

Organization	Value
Revised (1994) IDLH value*	3 ppm
NIOSH REL [2014] [†]	0.02 ppm (0.05 mg/m ³)
OSHA PEL[2014] [^]	0.02 ppm (0.05 mg/m ³) [skin]
ACGIH TLV [2014] [‡]	TWA 0.02 ppm (0.05 mg/m ³) STEL 0.06 ppm (0.14 mg/m ³)
AIHA ERPG [2010] ⁺	ERPG 1: 0.025 ppm ERPG 2: 0.5 ppm ERPG 3: 5 ppm
AIHA WEEL [2010] [§]	Not available

3 **Abbreviation:** ACGIH – American Conference of Governmental Industrial Hygienists; AIHA – American Industrial Hygiene
4 Association; ERPG – Emergency Response Preparedness Guidelines; IDLH – immediately dangerous to life or health; NIOSH – National
5 Institute for Occupational Safety and Health; OSHA – Occupational Safety and Health Administration; PEL – permissible exposure limit;
6 REL – recommended exposure limit; SCP – Standards Completion Program; STEL - short term exposure limit; TWA - time weighted
7 average; WEEL – workplace environmental exposure level

8 **References:** *NIOSH [1994]; ^OSHA [2014]; †NIOSH [2014]; ‡ ACGIH [2014]; AIHA [2010a]⁺; §AIHA [2010b]

1 **Table 3: AEGL Values for Methyl Isocyanate**

2

Classification	10-min	30-min	1-hour	4-hour	8-hour	End Point [reference]
AEGL-1	NR	NR	NR	NR	NR	Insufficient data
AEGL-2	0.40 ppm 0.94 mg/m ³	0.13 ppm 0.32 mg/m ³	0.067 ppm 0.16 mg/m ³	0.017 ppm 0.034 mg/m ³	0.008 ppm 0.02 mg/m ³	Decreased fetal body weights [Varma 1987]; Cardiac arrhythmias [Tepper et al. 1987]
AEGL-3	1.2 ppm 2.8 mg/m ³	0.40 ppm 0.95 mg/m ³	0.20 ppm 0.47 mg/m ³	0.05 ppm 0.12 mg/m ³	0.025 ppm 0.06 mg/m ³	Decreased pup survival during lactation [Schwetz et al. 1987]

3 **Abbreviation:** AEGL – acute exposure guideline levels; mg/m³ – milligrams per cubic meter; min – minute; NR – not recommended due to insufficient data; ppm – parts per million

4 * **References:** NAS [2003]

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2.0 Animal Toxicity Data

Methyl isocyanate is one of the most reactive isocyanates and rapidly degrades in water [Varma and Guest 1993; NAS 2003]. The reactive nature of methyl isocyanate contributes to its toxicological potential and ability to cause irritation and cytotoxicity in the respiratory tract. The database for acute lethality in animals consists of multiple studies in rabbits [Pant et al. 1987; Dow Chemical 1990], guinea pigs [Mellon Institute 1966, 1970; Dodd et al. 1985, 1986, 1987; Kolb et al. 1987; Troup et al. 1987; Ferguson and Alarie 1991], rats [Mellon Institute 1963b, 1970; IRDC 1964; Kimmerle and Eben 1964; Eastman Kodak 1966; Fait and Dodd 1981; Dodd et al. 1985, 1986, 1987; Nemery et al. 1985a; Salmon et al. 1985; Bucher et al. 1987a,b; Dinsdale et al. 1987; Pant et al. 1987; Vijayaraghavan and Kaushik 1987; Dutta et al. 1988; Sethi et al. 1989; Dow Chemical 1990; Jeevaratnam et al. 1990; Man Tech Environmental 1992; Jeevaratnam and Sriramachari 1994; Sriramachari and Jeevaratnam 1994], and mice [Dodd et al. 1985, 1986; Boorman et al. 1987a,b; Bucher et al. 1987a; Vijayaraghavan and Kaushik 1987; Varma et al. 1988]. Based on histopathology and necropsy reports, lethality appears to be primarily caused by damage to the respiratory system.

Multiple studies have reported varied non-lethal effects following acute exposure to methyl isocyanate. Studies in guinea pigs indicate that exposure to methyl isocyanate for 15 minutes causes hypoxia and metabolic acidosis at concentrations as low as 240 ppm [Fedde et al. 1987; Maginniss et al. 1987]. Rats acutely exposed to methyl isocyanate developed severe inflammation and erosion of the respiratory tract [Mitsumori et al. 1987] and hypoxia [Troup et al. 1987]. Studies that observed acutely exposed rats for extended periods of time reported evidence of pulmonary obstruction [Stevens et al. 1987; Bucher and Uriah 1989]. Tepper et al. [1987] reported an increase in cardiac arrhythmia in rats at 4-6 months post-exposure to a 2 hour exposure to 3 ppm methyl isocyanate. In mice, an RD_{50} value of 1.3 – 2.9 ppm and an RD_{50TC} of 1.9 ppm have been estimated [Ferguson et al. 1986; James et al. 1987].

Developmental and reproductive studies have determined that acute exposure to methyl isocyanate significantly decreases maternal body, fetal, and placental weight and significantly increases the total number of resorptions and skeletal malformations in both rats and mice [Varma 1987; Varma et al. 1990; Singh et al. 1994]. Varma et al. [1987] reported a LOAEL of 2 ppm for developmental and reproductive effects, including increased fetal resorptions and increased skeletal malformations, in dams exposed for 3 hours on GD 8. This represents a one-time acute exposure at a critical developmental time of the dam and fetuses.

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1 Table 4 summarizes the LC data identified in animal studies and provides 30-minute equivalent derived values for
2 methyl isocyanate. Table 5 provides non-lethal data reported in animal studies with 30-minute equivalent derived
3 values. Information in these tables includes species of test animals, toxicological metrics (i.e., LC, NOAEL,
4 LOAEL), adjusted 30-minute concentration, and the justification for the composite uncertainty factors applied to
5 calculate the derived values.
6

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1 **Table 4: Lethal Concentration Data for Methyl Isocyanate**

2

Reference	Species	LC ₅₀ (ppm)	LC _{L0} (ppm)	Time (min)	Adjusted 30-min Concentration* (ppm)	Composite Uncertainty Factor	Derived Value (ppm)†
Dodd et al. [1985, 1986]	Guinea Pig	5.4	--	360	12.4	10 [‡]	0.12
Ferguson and Alarie [1991]	Guinea Pig	26.5	--	180	48.2	10 [‡]	0.48
Vijayaraghavan and Kaushik [1987]	Mouse	112.4	--	30	112.4	10 [‡]	1.1
Kimmerle and Eben [1964]	Rat	5.0	--	240	10.0	10 [‡]	0.01
ManTech Environmental [1992]	Rat	45.0	--	60	56.7	10 [‡]	0.57

3 **Abbreviation:** LC – lethal concentration; LC₅₀ – median lethal concentration; LC_{L0} – lowest concentration of a chemical that caused death in humans or animals; min – minute; ppm – parts
4 per million

5 * For exposures other than 30 minutes the ten Berge et al. [1986] relationship is used for duration adjustment ($C^n \times t = k$); no empirically estimated n values were
6 available, therefore the default values were used, n = 3 for exposures greater than 30 minutes and n = 1 for exposures less than 30 minutes.

7 †The derived value is the result of the adjusted 30-minute concentration divided by the composite uncertainty factor.

8 ‡Composite uncertainty factor to account for adjustment of LC₅₀ values to LC₀₁ values in animals, interspecies differences and human variability.

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1 **Table 5: Non-lethal Concentration Data for Methyl Isocyanate**

2

Reference	Species (reference)	NOAEL (ppm)	LOAEL (ppm)	Time (min)	Adjusted 30-min Concentration* (ppm)	Composite Uncertainty Factor	Derived Value (ppm)†
Mellon Institute [1970]	Human	0.5	--	10	0.2	3‡	0.07
Mellon Institute [1963a]	Human	--	1	10	0.3	10±	0.03
Kimmerle and Eben [1964]	Human	--	4	5	0.7	10±	0.07
Tepper et al. [1987]	Rats	--	3	120	4.8	30^	0.16
Fedde et al. [1987]	Guinea Pig	--	240	15	120	30^	4.00
Varma [1987]⁺	Mouse	--	2	180	3.6	30^	0.12
Varma et al. [1990]	Rat	--	9	180	16.4	30^	0.55

3 **Abbreviation:** NOAEL – no observed adverse effect level; min – minute; LOAEL – lowest observed adverse effect level; ppm – parts per million

4 * For exposures other than 30 minutes the ten Berge et al. [1986] relationship is used for duration adjustment ($C^n \times t = k$), with n=3 for exposures greater than 30 minutes and n = 1 for

5 exposures less than 30 minutes.

6 † The derived value is the result of the adjusted 0.5-hour value divided by the composite uncertainty factor. The composite uncertainty factor used varies for each study based on the nature

7 and severity of the endpoint observed.

8 ‡ Composite uncertainty factor assigned to account for interspecies differences and human variability.

9 ± Composite uncertainty factor assigned to account for adjusting from a LOAEL to NOAEL, interspecies differences and human variability.

10 ^ Composite uncertainty factor assigned to account for adjusting from a LOAEL to NOAEL, severe effects, interspecies differences, human variability, and uncertainty about the threshold for

11 escape-impairing effects

12 † Identified study is the primary basis of the IDLH value for methyl isocyanate.

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3.0 Human Data

An accidental release in Bhopal, India is the source of most information on human exposures to methyl isocyanate. Although actual exposure concentrations are unknown, Karlsson et al. [1985] estimated a concentration range of 10 to 3,000 ppm, with higher concentrations occurring closer to the site of release, based on dispersion calculations. Although multiple deaths were reported, most occurred 8-72 hours post-release [Varma 1989; Varma and Guest 1993]. Symptoms reported with the most frequency were ocular irritation, coughing, respiratory distress, pulmonary congestion, nausea, vomiting, muscle weakness, and hypoxia resulting in CNS involvement [Kamat et al. 1985; Misra et al. 1987; Lorin and Kulling 1986; Andersson et al. 1988; Weill 1988; Kamat et al. 1992]. Most of these symptoms resolved within 2 weeks post-release, but many of those exposed, including adults and children, reported continued restrictive respiratory, ophthalmological, neuromuscular, and gastrointestinal symptoms for years after the accident [Andersson et al. 1984, 1985; Kamat et al. 1985; Irani and Mahashur 1986; Maskati 1986; Naik et al. 1986; Khurram and Ahmad 1987; Andersson et al. 1990; Kamat et al. 1992; Cullinan et al. 1997; Misra and Kalita 1997].

In addition to information from the Bhopal release, a case report and multiple experimental studies on human volunteers exist. Skin irritation and respiratory irritation were reported in workers exposed to unknown concentrations of methyl isocyanate for unknown durations [Union Carbide 1973]. In one experimental study, four volunteers were exposed to 0.4 to 21 ppm methyl isocyanate for 1-5 minutes [Kimmerle and Eben 1964]; minor irritation of the mucous membranes was reported at 2 ppm with more pronounced ocular irritation at 4 ppm. Volunteers reported that exposure to 21 ppm was instantaneously intolerable. Another experimental study exposed 8 volunteers to 1.75 ppm for 1 minute and then re-exposed 6 of the 8 to 0.5 ppm for 10 minutes [Mellon Institute 1970]. All volunteers reported ocular irritation and multiple volunteers reported nose and throat irritation in both exposures. A third study exposed 7 male volunteers to 0.3 to 5.0 ppm for 1 minute or to 1 ppm for 10 minutes [Mellon Institute 1963a]. Exposure to 2.5 and 5.0 ppm for 1 minute resulted in eye and nose irritation. One volunteer exposed to 5.0 ppm also reported throat irritation. Volunteers exposed to 1 ppm for 10 minutes reported eye, nose, and throat irritation.

4.0 Summary

Methyl isocyanate is one of the most reactive isocyanates and rapidly degrades in water [Varma and Guest 1993; NAS 2003]. The reactive nature of methyl isocyanate contributes to its toxicological potential and ability to cause

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1 irritation and cytotoxicity in the respiratory tract. The IDLH value for methyl isocyanate is based on a LOAEL,
2 which corresponds to severe reproductive and development effects in animals. Tepper et al. [1987] reported
3 cardiac arrhythmia in rats at 4-6 months post-exposure to a 2-hour exposure to 3 ppm methyl isocyanate. Varma
4 et al. [1987] reported a LOAEL of 2 ppm for developmental and reproductive effects, including increased fetal
5 resorptions and increased skeletal malformations, in dams exposed for 3 hours on GD 8. This represents a one-
6 time acute exposure at a critical developmental time of the dam and fetuses. The reproductive and developmental
7 effects occurred at lower concentrations than the cardiac effects and are identified as the most sensitive health
8 endpoint. The LOAEL of 2 ppm when adjusted to a 30-minute equivalent duration concentration is 3.6 ppm. A
9 composite uncertainty factor of 30 was applied to account for extrapolation from a concentration that causes
10 severe effects in animals, animal to human differences, human variability, and uncertainty about the threshold for
11 escape-impairing effects, yielding an IDLH value of 0.12 ppm for methyl isocyanate. This IDLH value should be
12 protective against the reproductive, developmental and cardiac effects associated with methyl isocyanate
13 exposure. Additionally, it is supported by duration-adjusted estimates of the irritation threshold in humans
14 [Mellon Institute 1963a; Kimmerle and Eben 1964].

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