

EPA-450/3-77-008b

October 1976

**PRELIMINARY SCORING
OF SELECTED ORGANIC
AIR POLLUTANTS
APPENDIX I -
CHEMISTRY, PRODUCTION,
AND TOXICITY
OF CHEMICALS
A THROUGH C**



**U.S. ENVIRONMENTAL PROTECTION AGENCY
Office of Air and Waste Management
Office of Air Quality Planning and Standards
Research Triangle Park, North Carolina 27711**

**PRELIMINARY SCORING
OF SELECTED ORGANIC AIR POLLUTANTS
APPENDIX I -
CHEMISTRY, PRODUCTION,
AND TOXICITY
OF CHEMICALS
A THROUGH C**

by

J. Dorigan, B. Fuller and R. Duffy

The Mitre Corporation
Metrek Division
Westgate Research Park
McLean, Virginia 22101

Contract No. 68-02-1495

Project No. 077L

EPA Project Officer: David R. Patrick

Prepared for

ENVIRONMENTAL PROTECTION AGENCY
Office of Air and Waste Management
Office of Air Quality Planning and Standards
Research Triangle Park, North Carolina 27711

October 1976

PREFACE

A program is being developed by the Emission Standards and Engineering Division of the U.S. Environmental Protection Agency to determine the extent and environmental effects of atmospheric industrial emissions of synthetic organic chemicals. As part of the initial effort, EPA requested that the MITRE Corporation, METREK Division, devise a system for identifying those chemicals that may be released to the atmosphere from chemical manufacturing plants and whose health effects have been documented. To address this need, three general categories of information for each compound to be evaluated were deemed directly relevant:

- (1) the quantity of the compound produced;
- (2) the potential for atmospheric release from chemical manufacturing plants, and
- (3) the toxicological effects of the compounds.

The results of this study are presented in the accompanying report "Preliminary Scoring of Organic Air Pollutants."

The following dossiers were prepared for each of the 637 compounds under consideration in an attempt to collate the pertinent data for subsequent evaluation.

TABLE OF CONTENTS

	<u>Page</u>
Acenaphthene	AI-2
Acetal	AI-4
Acetaldehyde	AI-6
Acetaldol	AI-8
Acetamide	AI-10
Acetamidofluorene	AI-12
Acetanilide	AI-14
Acetic acid	AI-16
Acetic anhydride	AI-18
Acetone	AI-20
Acetone cyanohydrin	AI-22
Acetonitrile	AI-24
Acetophenone	AI-26
Acetylene	AI-28
Acetylene tetrabromide	AI-30
Acetylene tetrachloride	AI-32
Acrolein	AI-34
Acrylamide	AI-36
Acrylic acid	AI-38
Acrylonitrile	AI-40
Adipic acid	AI-42
Alachlor	AI-44
Aldrin	AI-46
Allyl alcohol	AI-48
Allyl chloride	AI-50
Allylene	AI-52
Allyl naphthalene	AI-54
Amiben	AI-56
p-Aminobenzoic acid	AI-58
p-Aminobiphenyl	AI-60
Aminoethylethanolamine	AI-62
Amyl acetate	AI-64
Amyl alcohol	AI-66
Amylamines	AI-68
Amyl chloride(s)	AI-70
Amylene	AI-72
Amyl ether	AI-74
Amyl mercaptan	AI-76
Aniline	AI-78
Aniline hydrochloride	AI-80
Anisidine (o and p)	AI-82
Anisole	AI-84
Anthranilic acid	AI-86
Anthraquinone	AI-88
Auramine	AI-90
Azodrin	AI-92

TABLE OF CONTENTS (CONTINUED)

	<u>Page</u>
Banvel D	AI-94
Banvel T	AI-96
Benzaldehyde	AI-98
Benzamide	AI-100
Benzene	AI-102
Benzenedisulfonic acid	AI-104
Benezenesulfonic acid	AI-106
Benzidine	AI-108
Benzil	AI-110
Benzilic acid	AI-112
Benzoic acid	AI-114
Benzoin	AI-116
Benzonitrile	AI-118
Benzophenone	AI-120
Benzoquinone	AI-122
Benzotrichloride	AI-124
Benzoyl chloride	AI-126
Benzyl alcohol	AI-128
Benzylamine	AI-130
Benzylbenzene	AI-132
Benzyl benzoate	AI-134
Benzyl chloride	AI-136
Benzyl dichloride	AI-138
Bis(chloromethyl) ether	AI-140
Bisphenol A	AI-142
Bromacil	AI-144
Bromobenzene	AI-146
Bromonaphthalene	AI-148
1,3-Butadiene	AI-150
Butane	AI-152
1-Butene	AI-154
2-Butene (cis+trans)	AI-156
2-Butoxyethanol	AI-158
n-Butyl acetate	AI-160
sec-Butyl acetate	AI-162
tert-Butyl acetate	AI-164
n-Butylacrylate	AI-166
n-Butyl alcohol	AI-168
sec-Butyl alcohol	AI-170
tert-Butyl alcohol	AI-172
n-Butylaldehyde	AI-174
sec-Butylamine	AI-176
tert-Butylamine	AI-178
Butylbenzoic acid	AI-180
Butylene	AI-182
1,3-Butylene glycol	AI-184

TABLE OF CONTENTS (CONTINUED)

	<u>Page</u>
n-Butyl glycidyl ether	AI-186
Butylmercaptan	AI-188
tert-Butylphenol	AI-190
tert-Butyltoluene	AI-192
n-Butyric acid	AI-194
n-Butyric anhydride	AI-196
Butyronitrile	AI-198
Buxten	AI-200
Cacodylic acid	AI-202
Calcium propionate	AI-204
Calcium stearate	AI-206
Camphor	AI-208
Caprolactam	AI-210
Captan	AI-212
Carbaryl	AI-214
Carbofuran	AI-216
Carbon disulfide	AI-218
Carbon tetrabromide	AI-220
Carbon tetrachloride	AI-222
Castor oil	AI-224
Cellulose acetate	AI-226
Chloranil	AI-228
Chlordane	AI-230
Chloroacetaldehyde	AI-232
Chloroacetic acid	AI-234
Chloroacetophenone	AI-236
m-Chloroaniline	AI-238
o-Chloroaniline	AI-240
p-Chloroaniline	AI-242
o-Chlorobenzaldehyde	AI-244
p-Chlorobenzaldehyde	AI-246
Chlorobenzene	AI-248
Chlorobenzilate	AI-250
Chlorobenzoic acid	AI-252
o-Chlorobenzoyl chloride	AI-254
o-Chlorobenzylidene malonitrile	AI-256
Chlorodifluoroethane	AI-258
Chlorodifluoromethane	AI-260
2-Chloro-4-ethylamino-6-isopropylamino-s-triazine	AI-262
Chloroform	AI-264
Chloronaphthalene	AI-266
m-Chloronitrobenzene	AI-268
o-Chloronitrobenzene	AI-270
p-Chloronitrobenzene	AI-272
1-Chloro-1-nitropropane	AI-274
m-Chlorophenol	AI-276

TABLE OF CONTENTS (CONCLUDED)

	<u>Page</u>
o-Chlorophenol	AI-278
p-Chlorophenol	AI-280
Chloroprene	AI-282
Chloropropham	AI-284
m-Chlorotoluene	AI-286
o-Chlorotoluene	AI-288
p-Chlorotoluene	AI-290
Citric acid	AI-292
Crag herbicide 1	AI-294
m-Cresol	AI-296
o-Cresol	AI-298
p-Cresol	AI-300
Crotonaldehyde	AI-302
Crotonic acid	AI-304
Cumene	AI-306
Cumene hydroperoxide	AI-308
Cyanoacetic acid	AI-310
Cyanogen chloride	AI-312
Cyanuric acid	AI-314
Cyanuric chloride	AI-316
Cyclohexane	AI-318
Cyclohexanol	AI-320
Cyclohexanone	AI-322
Cyclohexene	AI-324
Cyclohexylamine	AI-326
Cyclopentadiene	AI-328
Cyprex	AI-330

APPENDIX I

CHEMISTRY, PRODUCTION AND TOXICITY OF SELECTED
SYNTHETIC ORGANIC CHEMICALS

Acenaphthene

CHEMICAL NAME

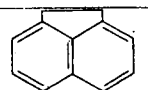
CAS Number: 208-96-8

Chemical Name: Acenaphthylene

Synonyms: 1,2-Dihydro-acenaphthylene
1,8-Ethylene naphthalene
Naphthyleneethylene

**Molecular
Formula:** $C_{12}H_8$

Structure:



Chemical Properties

Molecular Weight: 154.21

Physical State: Solid

Vapor Pressure: 10 mm at 131.2°C

Vapor Density: 5.32

Boiling Point: 277.5°C

Melting Point: 96.2°C

Density: 1.0242 at 99°C/4°C

Solubility: Insoluble (H_2O)

**Octanol/Water Parti-
tion Coefficient:**

Environmental Persistence

BOD:

Atmospheric Reactivity:

Safety Hazard: Fire-slight

Acenapthene

PRODUCTION DATA

Annual U.S.
Production:

Consumption:

Fraction of
Dispersion:

Fraction of Pro-
duction Lost:

Release Rate (million lbs/yr):

TOXICITY DATA

Acute Toxicity

Dosage

Animal

Route

Non-lethal Acute Effects:

Irritant to skin and mucous membranes

Chronic Toxicity

U.S. Occupational Standard:

Carcinogenicity: Neoplastic effects

Mutagenicity:

Teratogenicity:

TLV:

Other Chronic Effects:

Acetal

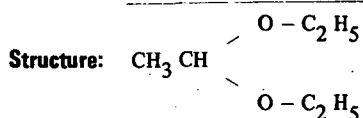
CHEMICAL NAME

CAS Number: 105-57-7

Chemical Name: Diethyl acetal acetaldehyde

Synonyms: 1,1-Diethoxyethane
Ethylidene diethyl ether
Diethylacetal

**Molecular
Formula:** $C_6H_{14}O_2$



Chemical Properties

Molecular Weight: 118.18

Vapor Pressure: 28.8 at 25°C

Boiling Point: 103.2°C at 760 mm

Density: 0.8314 at 20°C/4°C

**Octanol/Water Parti-
tion Coefficient:** 1.17

Physical State: Liquid

Vapor Density: 4.08 mp at 100°C

Melting Point:

Solubility: Soluble (H₂O)

Environmental Persistence

BOD:

Atmospheric Reactivity:

Safety Hazard: Fire—dangerous
Explosion—moderate

PRODUCTION DATA

Annual U.S.
Production: 56.8×10^6 lbs. (1975)

Consumption: 60.4×10^6 lbs.

Fraction of
Dispersion: 1.0

Fraction of Pro-
duction Lost: 0.03

Release Rate (million lbs/yr): 63.3

TOXICITY DATA

Acute Toxicity	Dosage	Animal	Route
LD ₅₀	4570 mg/kg	rat	oral
LC _{Lo}	4000 ppm (4 hours)	rat	inhalation

Non-lethal Acute Effects:

Narcotic

Chronic Toxicity

U.S. Occupational Standard:

Carcinogenicity:

Mutagenicity:

Teratogenicity:

TLV:

Other Chronic Effects:

Acetaldehyde

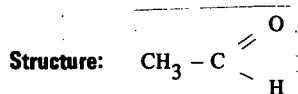
CHEMICAL NAME

CAS Number: 75-07-0

Chemical Name: Acetaldehyde

Synonyms: Acetic aldehyde
Ethyl aldehyde

**Molecular
Formula:** C_2H_4O



Chemical Properties

Molecular Weight: 44.05

Physical State: Liquid

Vapor Pressure: 923 Torrs at 25°C

Vapor Density: 1.52

Boiling Point: 20.8°C at 760 mm

Melting Point: -121°C

Density: 0.7834 at 18°C/4°C

Solubility: Infinite (hot H₂O)

**Octanol/Water Parti-
tion Coefficient:** 0.43

Environmental Persistence

BOD: 126 at 20°C for 5 days

Atmospheric Reactivity:

Safety Hazard: Fire—dangerous—oxidizes
Explosion—severe

Acetaldehyde

PRODUCTION DATA

Annual U.S.
Production: 1500×10^6 lbs. (1976) Consumption: 1300×10^6 lbs. (1976)

Fraction of
Dispersion: 0.01 Fraction of Pro-
duction Lost: 0.015

Release Rate (million lbs/yr): 36.2

TOXICITY DATA

Acute Toxicity	Dosage	Animal	Route
LD ₅₀	1930 mg/kg	rat	oral
	640 mg/kg	rat	subcutaneous
	1232 mg/kg	mouse	oral
	560 mg/kg	mouse	subcutaneous
LD _{Lo}	500 mg/kg	rat	intraperitoneal
LC _{Lo}	4000 ppm (4 hours)	rat	inhalation

Non-lethal Acute Effects:

TC _{Lo}	134 ppm	human	inhalation
------------------	---------	-------	------------

Chronic Toxicity

U.S. Occupational Standard: TWA-200 ppm

Carcinogenicity:

TD _{Lo}	60 mg/kg (79 weeks)	rat	subcutaneous
------------------	---------------------	-----	--------------

Mutagenicity:

Teratogenicity: TLV: 200 ppm

Other Chronic Effects:

Eye irritation	50 ppm
----------------	--------

Acetaldol

CHEMICAL NAME

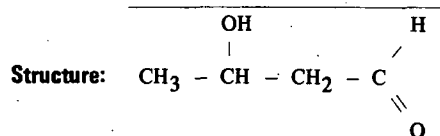
CAS Number: 107-89-1

Chemical Name: 3-Hydroxybutyraldehyde

Synonyms: Acetaldol
Aldol
3-Butanolal

Oxybutyric aldehyde

**Molecular
Formula:** C₄H₈O₂



Chemical Properties

Molecular Weight: 88.12

Physical State: Liquid

Vapor Pressure: 10 mm at 25°C

Vapor Density: 3.04

Boiling Point: 83°C at 20 mm
Decomposes at 85°C

Melting Point:

Density: 1.103 at 20°C/4°C

Solubility: Infinite (H₂O)

**Octanol/Water Parti-
tion Coefficient:**

Environmental Persistence

BOD:

Atmospheric Reactivity:

Safety Hazard:
Fire—moderate

PRODUCTION DATA**Annual U.S.
Production:****Consumption:****Fraction of
Dispersion:****Fraction of Pro-
duction Lost:****Release Rate (million lbs/yr):****TOXICITY DATA****Acute Toxicity
LD₅₀****Dosage**
2,180 mg/kg
140 mg/kg**Animal**
rat
rat**Route**
oral
skin**Chronic Toxicity****U.S. Occupational Standard:****Carcinogenicity:** Not tested**Mutagenicity:****Teratogenicity:****TLV:****Other Chronic Effects:**

Acetamide

CHEMICAL NAME

CAS Number: 60-35-5

Chemical Name: Acetamide

Synonyms: Acetic acid amide
Ethanamide

**Molecular
Formula:** C₂H₅NO

Structure:
$$\begin{array}{c} \text{O} \\ || \\ \text{CH}_3 - \text{C} - \text{NH}_2 \end{array}$$

Chemical Properties

Molecular Weight: 59.07

Physical State: Solid

Vapor Pressure: 1 mm at 65°C

Vapor Density:

Boiling Point: 221.2°C at 760 mm

Melting Point: 82.3°C

Density: 1.1590 at 20°C/4°C

Solubility: Soluble (H₂O)

**Octanol/Water Parti-
tion Coefficient:** -1.46

Environmental Persistence

BOD:

Atmospheric Reactivity:

Safety Hazard:

Disaster hazard--dangerous--cyanide

Acetamide

PRODUCTION DATA

Annual U.S.
Production:

Consumption:

Fraction of
Dispersion:

Fraction of Pro-
duction Lost:

Release Rate (million lbs/yr):

TOXICITY DATA

Acute Toxicity
LD₅₀

Dosage
10 gm/kg
8300 mg/kg

Animal
rat
mouse

Route
subcutaneous
subcutaneous

Non-lethal Acute Effects:
Mild irritant

Chronic Toxicity

U.S. Occupational Standard:

Carcinogenicity:
TD_{Lo}

456 gm/kg (52 weeks)

rat

oral

Mutagenicity:

Teratogenicity:

TLV:

Other Chronic Effects:

Acetamidofluorene

CHEMICAL NAME

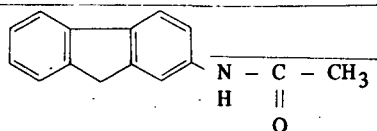
CAS Number: 53-96-3

Chemical Name: N-fluoren-2-yl-acetamide

Synonyms: 2-Acetamidofluorene
N-(2-fluorenyl)acetamide
2-Acetylaminofluorene

Molecular Formula: $C_{15}H_{13}NO$

Structure:



Chemical Properties

Molecular Weight: 223.28

Physical State: Solid

Vapor Pressure:

Vapor Density:

Boiling Point:

Melting Point: 194°C

Density:

Solubility: Insoluble (H₂O)

Octanol/Water Partition Coefficient:

Environmental Persistence

BOD:

Atmospheric Reactivity:

Safety Hazard:

Acetamidofluorene

PRODUCTION DATA

Annual U.S.
Production:

Consumption:

Fraction of
Dispersion:

Fraction of Pro-
duction Lost:

Release Rate (million lbs/yr):

TOXICITY DATA

Acute Toxicity
LD₅₀

Dosage
1020 mg/kg

Animal
mouse

Route
oral

Chronic Toxicity

U.S. Occupational Standard: (air) Carcinogen

Carcinogenicity:

TD_{Lo}

15 gm/kg (4 weeks)
475 mg/kg (11 weeks)
260 mg/kg (1 week)
192 mg/kg (8 weeks)
560 mg/kg (14 days)
4400 mg/kg (65 weeks)
7980 mg/kg (38 weeks)

hamster
rat
rat
rat
mouse
rabbit
hamster

intravenous
oral
skin
intraperitoneal
oral
oral
oral

Mutagenicity:

Teratogenicity:

TD_{Lo}
Other Chronic Effects:

100 mg/kg

TLV:
mouse

parenteral

Acetanilide

CHEMICAL NAME

CAS Number: 103-84-4

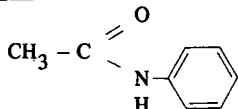
Chemical Name: Acetanilide

Synonyms: N-phenylacetamide
Antifibrin

Molecular

Formula: C_8H_9NO

Structure:



Chemical Properties

Molecular Weight: 135.17

Vapor Pressure: 1 mm at 114°C

Boiling Point: 303.8°C at 760 mm

Density: 1.2190 at 15°C/4°C

Octanol/Water Partition Coefficient: 1.59

Physical State: Solid

Vapor Density: 4.65

Melting Point: 114.3°C

Solubility: Slightly soluble (H₂O)

Environmental Persistence

BOD:

Atmospheric Reactivity:

Safety Hazard: Fire—slight

Acetanilide

PRODUCTION DATA

Annual U.S.**Production:** 5.5×10^6 lbs. (1967)**Consumption:****Fraction of****Dispersion:** 0.01**Fraction of Pro-****duction Lost:** 0.015**Release Rate (million lbs/yr):**

TOXICITY DATA

Acute Toxicity**Dosage****Animal****Route**LD₅₀

800 mg/kg

rat

oral

1210 mg/kg

mouse

oral

LD_{Lo}

1000 mg/kg

mouse

intraperitoneal

Chronic Toxicity**U.S. Occupational Standard:****Carcinogenicity:****Mutagenicity:****Teratogenicity:****TLV:****Other Chronic Effects:**

Acetic Acid

CHEMICAL NAME

CAS Number: 64-19-7

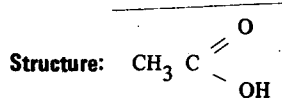
Chemical Name: Acetic acid

Synonyms: Methane carboxylic acid
Ethylic acid
Glacial acetic acid

Ethanoic acid
Vinegar acid

Molecular

Formula: $C_2H_4O_2$



Chemical Properties

Molecular Weight: 60.05

Physical State: Liquid

Vapor Pressure: 11.4 mm at 20°C

Vapor Density: 2.07

Boiling Point: 117.9°C at 760 mm

Melting Point: 16.6°C

Density: 1.0492 at 20°C/4°C

Solubility: Infinite (H₂O)

Octanol/Water Partition Coefficient: -0.31

Environmental Persistence

BOD: 102.7 at 20°C for 20 days

Atmospheric Reactivity:

Safety Hazard: Fire--moderate
Explosion--moderate

Acetic Acid

PRODUCTION DATA

Annual U.S.
Production: 2097.0 x 10⁶ lbs. (1975) Consumption: 2097.6 x 10⁶ lbs.

Fraction of
Dispersion: 0.01 Fraction of Pro-
duction Lost: 0.015

Release Rate (million lbs/yr): 55.9

TOXICITY DATA

Acute Toxicity	Dosage	Animal	Route
LD ₅₀	3310 mg/kg	rat	oral
	4960 mg/kg	mouse	oral
	525 mg/kg	mouse	intravenous
LD _{Lo}	1200 mg/kg	rabbit	oral
	1200 mg/kg	rabbit	subcutaneous
	600 mg/kg	rabbit	rectal
LC ₅₀	5620 ppm (1 hour)	mouse	inhalation

Non-lethal Acute Effects:

Irritant-LD _{Lo}	816 ppm (3 months)	human	inhalation
	50 ppm	human	inhalation
Gastrointestinal tract-LD _{Lo}	1470 mg/kg	human	oral

Chronic Toxicity

U.S. Occupational Standard: TWA-10 ppm

Carcinogenicity:

Mutagenicity:

Teratogenicity: TLV: 10 ppm

Other Chronic Effects:

Dermatitis
Ulcers
Slight irritation of
respiratory tract 100-260 ppm (1 hour)
Conjunctivitis
Bronchitis, pharyngitis
Erosion of exposed teeth

Acetic anhydride

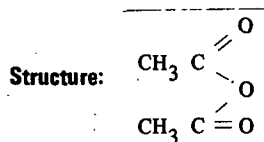
CHEMICAL NAME

CAS Number: 108-24-7

Chemical Name: Acetic anhydride

Synonyms: Acetyl oxide
Acetic oxide
Ethanoic anhydride

**Molecular
Formula:** $C_4H_6O_3$



Chemical Properties

Molecular Weight: 102.09

Physical State: Liquid

Vapor Pressure: 5.09 mm at 25°C

Vapor Density: 3.50

Boiling Point: 139.55°C

Melting Point: -73.1°C

Density: 1.082 at 20°C/4°C

Solubility: Very soluble (H₂O)

Octanol/Water Partition Coefficient:

Environmental Persistence

BOD:

Atmospheric Reactivity:

Safety Hazard: Fire—moderate
Explosion—moderate

Acetic anhydride

PRODUCTION DATA

Annual U.S.

Production: 1633.1×10^6 lbs. (1974)

Consumption: 1637×10^6 lbs.

Fraction of

Dispersion: 0.01

Fraction of Pro-

duction Lost: 0.015

Release Rate (million lbs/yr): 39.4

TOXICITY DATA

Acute Toxicity

LD₅₀

LC_{Lo}

Dosage

1780 mg/kg

1000 ppm (4 hours)

Animal

rat

rat

Route

oral

inhalation

Non-lethal Acute Effects:

Irritant to eyes and upper respiratory tract

Ingestion—burning pain in stomach followed by nausea and vomiting

Chronic Toxicity

U.S. Occupational Standard: TWA: 5 ppm (air)

Carcinogenicity:

Mutagenicity:

Teratogenicity:

TLV: 5 ppm

Other Chronic Effects:

Acetone

CHEMICAL NAME

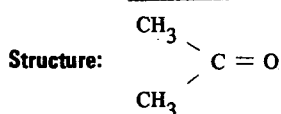
CAS Number: 67-64-1

Chemical Name: Acetone

Synonyms: 2-Propanone
Dimethyl ketone
 β -Ketone propane

Methylketone
Pyroacetic ether

Molecular
Formula: C_3H_6O



Chemical Properties

Molecular Weight: 58.08

Physical State: Liquid

Vapor Pressure: 400 mm at 39.5°C

Vapor Density: 2.0

Boiling Point: 56.2°C at 760 mm

Melting Point: -95.35°C

Density: 0.7972 at 15°C/4°C

Solubility: Infinite (H_2O)

Octanol/Water Partition Coefficient: 0.55

Environmental Persistence

BOD: 1.84 for 20 days at 20°C
72% of theoretical for 10 days at 20°C

Atmospheric Reactivity: Oxygen uptake: 50 mg/l during 12 hours

Safety Hazard: Fire—dangerous
Explosion—moderate

Acetone

PRODUCTION DATA

Annual U.S.**Production:** 1980.3×10^6 lbs. (1974)**Consumption:** 1946.0×10^6 lbs.**Fraction of****Dispersion:** 0.31**Fraction of Pro-****duction Lost:** 0.015**Release Rate (million lbs/yr):** 533.9

TOXICITY DATA

Acute ToxicityLD₅₀**Dosage**

1295 mg/kg

5300 mg/kg

8000 mg/kg

Animal

mouse

rabbit

dog

Route

intraperitoneal

oral

oral

Non-lethal Acute Effects:Eye effects—TC_{Lo} 500 ppm

Central nervous system

TC_{Lo} 12,000 ppm (4 hours)

Paralysis

human

air exposure

Chronic Toxicity**U.S. Occupational Standard:** TWA: 1000 ppm**Carcinogenicity:****Mutagenicity:****Teratogenicity:**

TLV: 1000 ppm

Other Chronic Effects:

Skin irritation

Chronic respiratory
tract irritation,
dizzinessMinor irritation of
eyes and nose**Odor perception:** 1.6 ppm

1000 ppm (3 hours/day)

3000 ppm

Acetone cyanohydrin

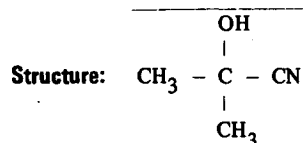
CHEMICAL NAME

CAS Number: 75-86-5

Chemical Name: 2-Methyl lactonitrile

Synonyms: Hydroxyisobutyronitrile

**Molecular
Formula:** C_4H_7NO



Chemical Properties

Molecular Weight: 85.11

Physical State: Liquid

Vapor Pressure:

Vapor Density: 2.93

Boiling Point: 82°C at 23 mm

Melting Point: -20°C

Density: 0.932 at 20°C/4°C

Solubility: Very soluble, decomposes in H_2O

**Octanol/Water Parti-
tion Coefficient:**

Environmental Persistence

BOD:

Atmospheric Reactivity:

Safety Hazard: Fire—slight

Acetone cyanohydrin

PRODUCTION DATA

Annual U.S.**Production:** 537.9 x 10⁶ lbs (1969)**Consumption:** 100 x 10⁶ lbs (1975)**Fraction of****Dispersion:** 0.01**Fraction of Pro-****duction Lost:** 0.015**Release Rate (million lbs/yr):** 6.0

TOXICITY DATA

Acute Toxicity**LD₅₀****Dosage**

17 mg/kg

3 mg/kg

14 mg/kg

17 mg/kg

9 mg/kg

Animal

rat

mouse

rabbit

rabbit

guinea pig

mouse

rat

Route

oral

oral

oral

skin

oral

inhalation

inhalation

LC₅₀

575 ppm (2 hours)

LC_{Lo}

63 ppm (4 hours)

Chronic Toxicity**U.S. Occupational Standard:****Carcinogenicity:****Mutagenicity:****Teratogenicity:****TLV:** Unconfirmed**Other Chronic Effects:**

Acetonitrile

CHEMICAL NAME

CAS Number: 75-05-8

Chemical Name: Acetonitrile

Synonyms: Methyl cyanide
Cyanomethane
Ethanenitrile

Ethyl nitrile
Methane carbonitrile

**Molecular
Formula:** C_2H_3N

Structure: $CH_3 - C \equiv N$

Chemical Properties

Molecular Weight: 41.05

Physical State: Liquid

Vapor Pressure: 92.8 mm at 25°C

Vapor Density: 1.42

Boiling Point: 81.6°C

Melting Point: -45.72°C

Density: 0.786 at 20°C/20°C

Solubility: Infinite (H₂O)

**Octanol/Water Parti-
tion Coefficient:** -0.34

Environmental Persistence

BOD: Total theoretical oxygen demand 3.32

Atmospheric Reactivity: Reacts to oxidizing materials

Safety Hazard: Fire—dangerous
Disaster hazard—dangerous, toxic fumes (HCN)

Acetonitrile

PRODUCTION DATA

Annual U.S.**Production:** 135×10^6 lbs.**Consumption:** 135×10^6 lbs.**Fraction of****Dispersion:** 0.50**Fraction of Pro-****duction Lost:** 0.015**Release Rate (million lbs/yr):** 69.5

TOXICITY DATA

Acute ToxicityLD₅₀**Dosage**

3800 mg/kg

1920 mg/kg

LD_{Lo}

700 mg/kg

LC_{Lo}

8000 ppm (4 hours)

Animal

rat

mouse

mouse

rat

Route

oral

intraperitoneal

subcutaneous

inhalation

Chronic Toxicity**U.S. Occupational Standard:** TWA: 40 ppm**Carcinogenicity:** Equivocal results from 2-year rat study.**Mutagenicity:****Teratogenicity:** Fetal malformations in rats, 0.2
to 1.0 ml/kg intraperitoneal**TLV:** 40 ppm**Other Chronic Effects:**

Growth retardation

Metabolic disturbances

Liver enlargement

Acetophenone

CHEMICAL NAME

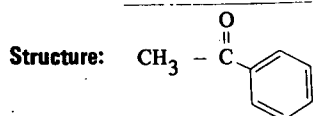
CAS Number: 98-86-2

Chemical Name: Acetophenone

Synonyms: 1-Phenyl-ethanone
Methyl phenyl ketone
Phenyl methyl ketone

Acetylbenzene
Hypnone

Molecular
Formula: C_8H_8O



Chemical Properties

Molecular Weight: 120.16

Physical State: Liquid or solid

Vapor Pressure: 1 mm at 37.1°C

Vapor Density: 4.14

Boiling Point: 202.0°C

Melting Point: 20.5°C

Density: 1.0281 at 20°C/4°C

Solubility: Insoluble (H_2O)

Octanol/Water Partition Coefficient:

Environmental Persistence

BOD:

Atmospheric Reactivity: Reacts with oxidizing materials

Safety Hazard: Fire—slight

Acetophenone

PRODUCTION DATA

Annual U.S.
Production: 2.68×10^6 lbs.

Consumption:

Fraction of
Dispersion: 0.01

Fraction of Pro-
duction Lost: 0.015

Release Rate (million lbs/yr):

TOXICITY DATA

Acute Toxicity

LD₅₀
LD_{Lo}

Dosage
900 mg/kg
200 mg/kg

Animal
rat
mouse

Route
oral
intraperitoneal

Non-lethal Acute Effects:

Dermatitis
Narcosis
CNS depressant

Chronic Toxicity

U.S. Occupational Standard:

Carcinogenicity:

Mutagenicity:

Teratogenicity:

TLV:

Other Chronic Effects:

Acetylene

CHEMICAL NAME

CAS Number: 74-86-2

Chemical Name: Acetylene

Synonyms: Ethyne
Ethine

Molecular
Formula: C_2H_2

Structure: $HC \equiv CH$

Chemical Properties

Molecular Weight: 26.04

Physical State: Gas

Vapor Pressure: 5600 mm at 16.8°C

Vapor Density: 0.91

Boiling Point: -84.0°C at 760 mm (sublimes)

Melting Point: -80.8°C

Density: 0.6181 at -32°C/4°C

Solubility: Slight (H_2O)

Octanol/Water Parti-
tion Coefficient:

Environmental Persistence

BOD:

Atmospheric Reactivity:

Safety Hazard: Fire—dangerous
Explosion—moderate

Acetylene

PRODUCTION DATA

Annual U.S.**Production:** 538 x 10⁶ lbs. (1974)**Consumption:** 617 x 10⁶ lbs.**Fraction of****Dispersion:** 0.01**Fraction of Pro-****duction Lost:** 0.015**Release Rate (million lbs/yr):** 9.1

TOXICITY DATA

Acute ToxicityLC_{Lo}**Dosage**

90,000 ppm

Animal

rat

Route

inhalation

Non-lethal Acute Effects:

Nausea

Cyanosis

CNS depression

Chronic Toxicity**U.S. Occupational Standard:****Carcinogenicity:****Mutagenicity:****Teratogenicity:****TLV:****Other Chronic Effects:**

Intoxication:	100,000 ppm	man (slight)
	200,000 ppm	man (marked)
Incoordination:	300,000 ppm	man
Unconsciousness:	350,000 ppm (5 minutes)	man

Acetylene tetrabromide

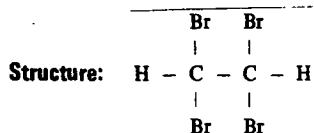
CHEMICAL NAME

CAS Number: 79-27-6

Chemical Name: 1,1,2,2-Tetrabromoethane

Synonyms: Acetylenetetrabromide
Muthmann's liquid

**Molecular
Formula:** $C_2H_2Br_4$



Chemical Properties

Molecular Weight: 345.67

Physical State: Liquid

Vapor Pressure: 1 mm at 65°C

Vapor Density:

Boiling Point: 243.5°C at 760 mm

Melting Point: 0°C

Density: 2.9656 at 20°C/4°C

Solubility: Insoluble (H_2O)

Octanol/Water Partition Coefficient:

Environmental Persistence

BOD:

Atmospheric Reactivity:

Safety Hazard:

Disaster hazard—dangerous—toxic fumes of carbonyl bromide when heated

Acetylene tetrabromide

PRODUCTION DATA

Annual U.S.
Production:

Consumption:

Fraction of
Dispersion:

Fraction of Pro-
duction Lost:

Release Rate (million lbs/yr):

TOXICITY DATA

Acute Toxicity
LD₅₀

Dosage
400 mg/kg
400 mg/kg

Animal
rabbit
guinea pig

Route
oral
oral

Non-lethal Acute Effects:

Irritant
Narcotic

Chronic Toxicity

U.S. Occupational Standard: (air) TWA 1 ppm

Carcinogenicity:

Mutagenicity:

Teratogenicity:

TLV:

Other Chronic Effects:

Acetylene tetrachloride

CHEMICAL NAME

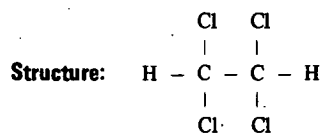
CAS Number: 79-34-5

Chemical Name: 1,1,2,2-Tetrachloroethane

Synonyms: Acetylene tetrachloride

Molecular

Formula: $C_2H_2Cl_4$



Chemical Properties

Molecular Weight: 167.85

Physical State: Liquid

Vapor Pressure: 6.75 mm at 25°C

Vapor Density:

Boiling Point: 146.2°C at 760 mm

Melting Point: -43.8°C

Density: 1.5953 at 20°C/4°C

Solubility: Slightly (H_2O)

Octanol/Water Partition Coefficient:

Environmental Persistence

BOD:

Atmospheric Reactivity:

Safety Hazard:

Explosion—moderate if contact with Na or K

Acetylene tetrachloride

PRODUCTION DATA

Annual U.S.
Production:

Consumption:

Fraction of
Dispersion:

Fraction of Pro-
duction Lost:

Release Rate (million lbs/yr):

TOXICITY DATA

Acute Toxicity	Dosage	Animal	Route
LD _{Lo}	30 mg/kg	mouse	intraperitoneal
	700 mg/kg	dog	oral
	50 mg/kg	dog	intravenous
	500 mg/kg	rabbit	subcutaneous
LC _{Lo}	1000 ppm (4 hours)	rat	inhalation

Non-lethal Acute Effects:

Central nervous system			
TD _{Lo}	30 mg/kg	human	oral

Chronic Toxicity

U.S. Occupational Standard: (air) TWA 5 ppm

Carcinogenicity:

Mutagenicity:

Teratogenicity:

TLV:

Other Chronic Effects:

Acrolein

CHEMICAL NAME

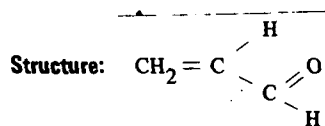
CAS Number: 107-02-8

Chemical Name: Acrolein

Synonyms: 2-Propenal Acraldehyde
Acrylic aldehyde
Allyl aldehyde

Molecular

Formula: C_3H_4O



Chemical Properties

Molecular Weight: 56.06

Physical State: Liquid

Vapor Pressure: 288.2 mm at 25°C

Vapor Density: 1.94

Boiling Point: 52.5°C

Melting Point: -86.95°C

Density: 0.8410 at 20°C/4°C

Solubility: Very soluble, (400,000 mg/l of H₂O)

Octanol/Water Partition Coefficient:

Environmental Persistence

BOD: 33% of theoretical after 10 days

Atmospheric Reactivity: Reactive with oxidizing materials

Safety Hazard: Fire-dangerous

PRODUCTION DATA

Annual U.S. Production: 61.73×10^6 lbs.

Consumption:

Fraction of Dispersion: 0.01

Fraction of Production Lost: 0.015

Release Rate (million lbs/yr):

TOXICITY DATA

Acute Toxicity	Dosage	Animal	Route
LD₅₀	46 mg/kg	rat	oral
	50 mg/kg	rat	subcutaneous
	30 mg/kg	mouse	subcutaneous
	7 mg/kg	rabbit	oral
LD_{Lo}	562 mg/kg	rabbit	skin
	2 mg/kg	mouse	intraperitoneal
	150 mg/kg	rabbit	subcutaneous
	150 mg/kg	guinea pig	subcutaneous
LC_{Lo}	153 ppm (10 minutes)	human	inhalation
	8 ppm (4 hours)	rat	inhalation
	24 mg/m ³ (6 hours)	mouse	inhalation
	1570 mg/m ³ (8 hours)	cat	inhalation
	24 mg/m ³ (6 hours)	rabbit	inhalation
	24 mg/m ³ (6 hours)	guinea pig	inhalation

Chronic Toxicity

U.S. Occupational Standard: TWA: 0.1 ppm

Carcinogenicity:

Mutagenicity:

Teratogenicity:

TLV: 0.1 ppm

Other Chronic Effects: Lachrymation
Irritant to respiratory tract
CNS damage

Acrylamide

CHEMICAL NAME

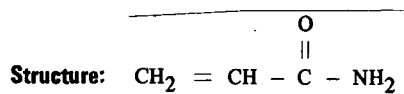
CAS Number: 79-06-1

Chemical Name: Acrylamide

Synonyms: Propenoic acid, amide
Propenamide
Acrylic amide

Molecular

Formula: C₃H₅NO



Chemical Properties

Molecular Weight: 71.08

Physical State: Solid

Vapor Pressure: 1.6 mm at 84.5°C

Vapor Density:

Boiling Point: 125°C at 25 mm

Melting Point: 84 to 85°C

Density: 1.122 at 30°C

Solubility: Very soluble (H₂O)

Octanol/Water Partition Coefficient:

Environmental Persistence

BOD:

Atmospheric Reactivity:

Safety Hazard:

Disaster hazard—moderate—emits acrid fumes

Acrylamide

PRODUCTION DATA

Annual U.S.
Production: 35×10^6 lbs.

Consumption:

Fraction of
Dispersion:

Fraction of Pro-
duction Lost: 0.015

Release Rate (million lbs/yr):

TOXICITY DATA

Acute Toxicity	Dosage	Animal	Route
LD ₅₀	170 mg/kg	rat	oral
	170 mg/kg	mouse	oral
LD _{Lo}	126 mg/kg	rabbit	oral
	1000 mg/kg	rabbit	skin
	252 mg/kg	guinea pig	oral

Non-lethal Acute Effects:
CNS paralysis

Chronic Toxicity

U.S. Occupational Standard: (air) TWA 0.3 mg/m^3

Carcinogenicity:

Mutagenicity:

Teratogenicity:

TLV: 0.3 mg/m^3 (skin)

Other Chronic Effects:

Acrylic acid

CHEMICAL NAME

CAS Number: 79-10-7

Chemical Name: Acrylic acid

Synonyms: Acroleic acid
Propene acid
2-Propenoic acid

**Molecular
Formula:** $C_3H_4O_2$

Structure:
$$H_2C = CH - C \begin{array}{l} \nearrow O \\ \searrow OH \end{array}$$

Chemical Properties

Molecular Weight: 72.06

Physical State: Liquid

Vapor Pressure: 4.61 mm at 25°C

Vapor Density: 2.45

Boiling Point: 141°C

Melting Point: 13°C

Density: 1.0511 at 20°C/4°C

Solubility: Infinite (H₂O)

**Octanol/Water Parti-
tion Coefficient:** 0.43

Environmental Persistence

BOD:

Atmospheric Reactivity: Reacts with oxidizing material

Safety Hazard: Fire—slight
Explosion—moderate, spontaneous polymerization
Disaster hazard—dangerous, toxic fumes

Acrylic acid

PRODUCTION DATA

Annual U.S.
Production: 233 x 10⁶ lbs. (1974) Consumption:
Fraction of
Dispersion: 0.01 Fraction of Pro-
duction Lost: 0.015
Release Rate (million lbs/yr): 2.7

TOXICITY DATA

Acute Toxicity	Dosage	Animal	Route
LD ₅₀	340 mg/kg	rat	oral
	60 mg/kg	mouse	oral
	280 mg/kg	rabbit	skin
LDLo	128 mg/kg	mouse	intraperitoneal
LCLo	6000 ppm (5 hours)	rat	inhalation

Non-lethal Acute Effects:
Irritant to eyes and skin

Chronic Toxicity

U.S. Occupational Standard:

Carcinogenicity:

Mutagenicity:

Teratogenicity:

TLV: unconfirmed

Other Chronic Effects:

Acrylonitrile

CHEMICAL NAME

CAS Number: 107-13-1

Chemical Name: Acrylonitrile

Synonyms: 2-Propenenitrile
Vinyl cyanide

**Molecular
Formula:** C_3H_3N

Structure: $CH_2 = CH - C \equiv N$

Chemical Properties

Molecular Weight: 53.00

Physical State: Liquid

Vapor Pressure: 113.8 mm at 25°C

Vapor Density: 1.83

Boiling Point: 77.5°C

Melting Point: -83 to -84°C

Density: 0.8060 at 20°C/4°C

Solubility: Soluble (H₂O)

**Octanol/Water Parti-
tion Coefficient:** -0.92

Environmental Persistence

BOD:

Atmospheric Reactivity: Reacts with oxidizing materials

Safety Hazard: Fire—dangerous
Explosion—moderate

Acrylonitrile

PRODUCTION DATA

Annual U.S.**Production:** 1411.8 x 10⁶ lbs. (1974)**Consumption:** 511.7 x 10⁶ lbs. (1974)**Fraction of****Dispersion:** 0.01**Fraction of Pro-****duction Lost:** 0.015**Release Rate (million lbs/yr):** 27.4 (1972)

TOXICITY DATA

Acute ToxicityLD₅₀**Dosage**

93 mg/kg

96 mg/kg

93 mg/kg

280 mg/kg

250 mg/kg

15 mg/kg

LD_{Lo}LC₁₀₀LC₅₀LC_{Lo}

784 ppm (1 hour)

576 ppm (4 hours)

500 ppm (4 hours)

120 mg/kg (4 hours)

600 ppm (4 hours)

258 ppm (4 hours)

Animal

rat

rat

rabbit

rabbit

guinea pig

mouse

mouse

guinea pig

rat

dog

cat

rabbit

Route

oral

subcutaneous

oral

skin

skin

intraperitoneal

inhalation

inhalation

inhalation

inhalation

inhalation

inhalation

Chronic Toxicity**U.S. Occupational Standard:** TWA: 20 ppm**Carcinogenicity:****Mutagenicity:****Teratogenicity:**

TLV: 20 ppm (skin)

Other Chronic Effects:

Inhibits respiratory enzyme

Irritation of eyes and nose

Anemia and leucocytosis

Affects liver and kidney

Odor perception: 21.4 ppm

Adipic acid

CHEMICAL NAME

CAS Number: 124-04-9

Chemical Name: Adipic acid

Synonyms: Hexanedioic acid
1,4-Butane dicarboxylic acid
Adipinic acid

**Molecular
Formula:** $C_6H_{10}O_4$

Structure:

$$\begin{array}{c} CH_2 - CH_2 - COOH \\ | \\ CH_2 - CH_2 - COOH \end{array}$$

Chemical Properties

Molecular Weight: 146.14

Physical State: Solid

Vapor Pressure: 1 mm at 159.5°C

Vapor Density: 5.04

Boiling Point: 337.5°C

Melting Point: 153°C

Density: 1.360 at 25°C/4°C

Solubility: Slightly, (15 gm/l of H_2O)

**Octanol/Water Parti-
tion Coefficient:** -0.14

Environmental Persistence

BOD:

Atmospheric Reactivity: Reacts with oxidizing materials

Safety Hazard: Fire-slight

Adipic acid

PRODUCTION DATA

Annual U.S.**Production:** 1478.4 x 10⁶ lbs. (1974)**Consumption:** 1475 x 10⁶ lbs.**Fraction of****Dispersion:** 0.01**Fraction of Pro-****duction Lost:** 0.015**Release Rate (million lbs/yr):** 37.3

TOXICITY DATA

Acute ToxicityLD₅₀

LDLo

Dosage

1900 mg/kg

3600 mg/kg

Animal

mouse

rat

Route

oral

oral

Chronic Toxicity**U.S. Occupational Standard:****Carcinogenicity:****Mutagenicity:****Teratogenicity:****TLV:****Other Chronic Effects:**

Alachlor

CHEMICAL NAME

CAS Number: 15972-60-8

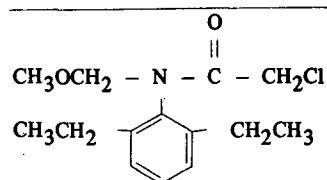
Chemical Name: 2-Chloro-N-(2,6-diethylphenyl)-N-(methoxymethyl)acetamide

Synonyms: 2-Chloro-2',6'-diethyl-N-(methoxymethyl)-acetanilide
Methachlor

Molecular

Formula: $C_{14}H_{20}ClNO_2$

Structure:



Chemical Properties

Molecular Weight: 269.90

Vapor Pressure: 2.2×10^{-5} mm at 25°C

Boiling Point: 100°C

Density:

Octanol/Water Partition Coefficient:

Physical State:

Vapor Density:

Melting Point:

Solubility: 140 ppm H₂O

Environmental Persistence

BOD:

Atmospheric Reactivity:

Safety Hazard:

PRODUCTION DATA**Annual U.S.****Production:** 20×10^6 lbs.**Consumption:** 20×10^6 lbs.**Fraction of****Dispersion:** Estimated as 1.0**Fraction of Pro-****duction Lost:** 0.01**Release Rate (million lbs/yr):** 20.2**TOXICITY DATA****Acute Toxicity**LD₅₀**Dosage**

1200 mg/kg

3000 mg/kg

Animal

rat

mammal

Route

oral

oral

Chronic Toxicity**U.S. Occupational Standard:****Carcinogenicity:****Mutagenicity:****Teratogenicity:****TLV:****Other Chronic Effects:**

Aldrin

CHEMICAL NAME

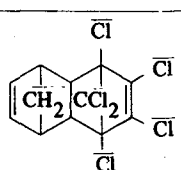
CAS Number: 309-00-2

Chemical Name: 1,2,3,4,10,10-Hexachloro-1,4,4a,5,8,8a-hexahydro-endo,exo-1,4:5,8-dimethanonaphthalene

Synonyms: Octalene
Aldrine
Compound 118

Molecular Formula: $C_{12}H_8Cl_6$

Structure:



Chemical Properties

Molecular Weight: 364.82

Physical State: Solid

Vapor Pressure: 2.31×10^{-5} Torr at 20°C

Vapor Density:

Boiling Point:

Melting Point: 104°C

Density:

Solubility: Insoluble (H₂O)

Octanol/Water Partition Coefficient: 5.00

Environmental Persistence

BOD:

Atmospheric Reactivity: Can be degraded anaerobically
Aldrin → dieldrin in the environment

Safety Hazard:

PRODUCTION DATA

Annual U.S.
Production: 25 x 10⁶ lbs. **Consumption:** 25 x 10⁶ lbs.
Fraction of
Dispersion: 1.0 **Fraction of Pro-**
 duction Lost: 0.01
Release Rate (million lbs/yr): 25.3 (1972)

TOXICITY DATA

Acute Toxicity	Dosage	Animal	Route
LD ₅₀	55 mg/kg	rat	oral
	200 mg/kg	rat	skin
	39 mg/kg	rat	oral
	98 mg/kg	rat	skin
	10 mg/kg	chicken	oral
	7 mg/kg	wild bird	oral
LD _{Lo}	15 mg/kg	rat	skin
	63 mg/kg	mouse	intraperitoneal
	95 mg/kg	dog	unknown

Non-lethal Acute Effects:

Irritability
 Convulsions
 Liver damage

Chronic Toxicity

U.S. Occupational Standard:

Carcinogenicity:

Neoplastic effects

TD_{Lo}: 440 mg/kg (52 weeks)—mouse, oral

Mutagenicity:

Teratogenicity:

TLV: 0.25 mg/m³ (skin)

Other Chronic Effects:

Allyl alcohol

CHEMICAL NAME

CAS Number: 107-18-6

Chemical Name: Allyl alcohol

Synonyms: 2-Propen-1-ol
Vinyl carbinol

**Molecular
Formula:** C₃H₆O

Structure: $\text{CH}_2 = \text{CH} - \text{CH}_2\text{OH}$

Chemical Properties

Molecular Weight: 58.08

Physical State: Liquid

Vapor Pressure: 25.6 mm at 25°C

Vapor Density: 2.00

Boiling Point: 97°C

Melting Point: -129°C

Density: 0.8540 at 20°C/4°C

Solubility: Infinite (H₂O)

**Octanol/Water Parti-
tion Coefficient:** 0.13

Environmental Persistence

BOD: 9.1% of theoretical

Atmospheric Reactivity: Reacts with oxidizing materials

Safety Hazard:

Fire—dangerous

Explosion—moderate

Disaster hazard—dangerous—toxic fumes

Allyl alcohol

PRODUCTION DATA

Annual U.S.
Production:

Consumption:

Fraction of
Dispersion:

Fraction of Pro-
duction Lost:

Release Rate (million lbs/yr):

TOXICITY DATA

Acute Toxicity	Dosage	Animal	Route
LD ₅₀	42 mg/kg	rat	intraperitoneal
	96 mg/kg	mouse	oral
	42 mg/kg	mouse	intraperitoneal
LD _{Lo}	69 mg/kg	rat	oral
	43 mg/kg	dog	oral
	53 mg/kg	rabbit	oral
	53 mg/kg	rabbit	skin
	45 mg/kg	rabbit	parenteral
LC ₅₀	165 ppm (4 hours)	rat	inhalation
LC _{Lo}	1000 ppm (4 hours)	monkey	inhalation
	1000 ppm (4 hours)	rabbit	inhalation

Non-lethal Acute Effects:

Irritant effects:

TC_{Lo} 25 ppm

human

inhalation

Eyes and skin irritated

Respiratory tract irritated

Damage to kidney and liver

Chronic Toxicity

U.S. Occupational Standard: (air) TWA 2 ppm

Carcinogenicity:

Mutagenicity:

Teratogenicity:

TLV: 2 ppm (skin)

Other Chronic Effects:

Allyl chloride

CHEMICAL NAME

CAS Number: 107-05-1

Chemical Name: 3-Chloropropene

Synonyms: 3-Chloroprene 1-Chloro-2-propene
AC 3-Chloropropylene
Chlorallylene 2-Chloropropylene

Molecular

Formula: C_3H_5Cl

Structure: $CH_2 = CH - CH_2 Cl$

Chemical Properties

Molecular Weight: 76.53

Physical State: Liquid

Vapor Pressure: 359 mm at 25°C

Vapor Density: 2.64

Boiling Point: 44.6°C

Melting Point: -134.5°C

Density: 0.938 at 20°C/4°C

Solubility: Soluble (3.3 gm/l H_2O)

Octanol/Water Partition Coefficient:

Environmental Persistence

BOD:

Atmospheric Reactivity: Unreactive to O_2

Reacts vigorously with oxidizing materials

Reactive with O_3 , $t_{1/2}$ = 9 hours, yielding formaldehyde and 2-chloroacetaldehyde

Reactive with OH, $t_{1/2}$ = 21 hours, yielding 2-chloroacetaldehyde

Safety Hazard: Fire—highly dangerous
Explosion—moderate

Allyl chloride

PRODUCTION DATA

Annual U.S.
Production: 295.0 x 10⁶ lbs.

Consumption: 295.0 x 10⁶ lbs.

Fraction of
Dispersion: 0

Fraction of Pro-
duction Lost: 0.015

Release Rate (million lbs/yr): 4.4 (1972)

TOXICITY DATA

Acute Toxicity	Dosage	Animal	Route
LD ₅₀	155 mg/kg	mouse	intraperitoneal
	7150 mg/kg	dog	intravenous
	2200 mg/kg	rabbit	skin
LD _{Lo}	64 mg/kg	rat	oral
LC _{Lo}	290 ppm (8 hours)	rat	inhalation
	6360 ppm	guinea pig	inhalation
Pulmonary hemorrhage			

Chronic Toxicity

U.S. Occupational Standard: TWA: 1 ppm

Carcinogenicity:

Mutagenicity:

Teratogenicity:

TLV: 1 ppm

Other Chronic Effects: Irritation of eyes, nose, and throat
Degeneration of liver and kidney

Allylene

CHEMICAL NAME

CAS Number: 74-99-7

Chemical Name: Propyne

Synonyms: Allylene
Methylacetylene
Propine

**Molecular
Formula:** C₃H₄

Structure: $\text{CH}_3 - \text{C} \equiv \text{CH}$

Chemical Properties

Molecular Weight: 40.07

Physical State: Gas

Vapor Pressure: 3876 mm at 20°C

Vapor Density: 1.38

Boiling Point: -23.2°C at 760 mm

Melting Point: -101.5°C

Density: 0.7062 at -50°C/4°C

Solubility: Slightly (H₂O)

**Octanol/Water Parti-
tion Coefficient:**

Environmental Persistence

BOD:

Atmospheric Reactivity: Reacts violently with oxidizing materials

Safety Hazard:

Fire—dangerous

Explosion—moderate

PRODUCTION DATA**Annual U.S.
Production:****Consumption:****Fraction of
Dispersion:****Fraction of Pro-
duction Lost:****Release Rate (million lbs/yr):****TOXICITY DATA****Acute Toxicity****Dosage****Animal****Route****Non-lethal Acute Effects:**

Anesthetic

Asphyxiant in high concentrations

Chronic Toxicity**U.S. Occupational Standard:****Carcinogenicity:****Mutagenicity:****Teratogenicity:****TLV:****Other Chronic Effects:**

Allyl naphthalene

CHEMICAL NAME

CAS Number: 2489-86-3

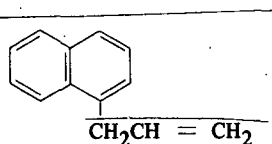
Chemical Name: 1-Allylnaphthalene

Synonyms: 3- α -Naphthylpropene

Molecular

Formula: $C_{13}H_{12}$

Structure:



Chemical Properties

Molecular Weight: 168.24

Physical State: Solid

Vapor Pressure:

Vapor Density:

Boiling Point: 265 to 267°C

Melting Point:

Density: 1.0228 at 20°C/4°C

Solubility:

Octanol/Water Partition Coefficient:

Environmental Persistence

BOD:

Atmospheric Reactivity:

Safety Hazard:

Allyl naphthalene

PRODUCTION DATA

**Annual U.S.
Production:**

Consumption:

**Fraction of
Dispersion:**

**Fraction of Pro-
duction Lost:**

Release Rate (million lbs/yr):

TOXICITY DATA

Acute Toxicity

Dosage

Animal

Route

Chronic Toxicity

U.S. Occupational Standard:

Carcinogenicity:

Mutagenicity:

Teratogenicity:

TLV:

Other Chronic Effects:

Amiben

CHEMICAL NAME

CAS Number: 133-90-4

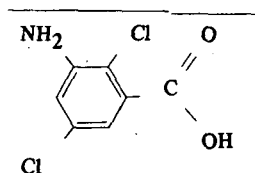
Chemical Name: 3-Amino-2,5-dichlorobenzoic acid

Synonyms: 2,5-Dichloro-3-aminobenzoic acid

Molecular

Formula: $C_7H_5Cl_2NO_2$

Structure:



Chemical Properties

Molecular Weight: 205.96

Physical State: Solid

Vapor Pressure: Negligible at 25°C

Vapor Density:

Boiling Point:

Melting Point: 201°C

Density:

Solubility: 700 ppm (H₂O)

Octanol/Water Partition Coefficient:

Environmental Persistence

BOD:

Atmospheric Reactivity:

Safety Hazard:

Disaster hazard—dangerous—toxic fumes

Amiben

PRODUCTION DATA

Annual U.S.
Production: 20×10^6 lbs.

Consumption: 20×10^6 lbs.

Fraction of
Dispersion: 1.0

Fraction of Pro-
duction Lost: 0.01

Release Rate (million lbs/yr): 20.2

TOXICITY DATA

Acute Toxicity
LD₅₀

Dosage
5620 mg/kg

Animal
rat

Route
oral

Chronic Toxicity

U.S. Occupational Standard:

Carcinogenicity:

Mutagenicity:

Teratogenicity:

TLV:

Other Chronic Effects:

p-Aminobenzoic acid

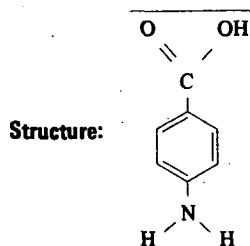
CHEMICAL NAME

CAS Number: 150-13-0

Chemical Name: p-Aminobenzoic acid

Synonyms: PABA
Paraminol

**Molecular
Formula:** $C_7H_7NO_2$



Chemical Properties

Molecular Weight: 137.14

Physical State: Solid

Vapor Pressure:

Vapor Density:

Boiling Point:

Melting Point: 188 to 189°C

Density: 1.374 at 25°C/4°C

Solubility: Soluble (hot H₂O)

**Octanol/Water Parti-
tion Coefficient:** 0.89

Environmental Persistence

BOD:

Atmospheric Reactivity:

Safety Hazard:

p-Aminobenzoic acid

PRODUCTION DATA

**Annual U.S.
Production:**

Consumption:

**Fraction of
Dispersion:**

**Fraction of Pro-
duction Lost:**

Release Rate (million lbs/yr):

TOXICITY DATA

Acute Toxicity
LD₅₀

Dosage
2850 mg/kg
1830 mg/kg
2000 mg/kg
6000 mg/kg
1000 mg/kg

Animal
mouse
rabbit
rabbit
rat
dog

Route
oral
oral
intravenous
oral
oral

Non-lethal Acute Effects:
Methemoglobinemia
Hepatitis

Chronic Toxicity

U.S. Occupational Standard:

Carcinogenicity:

Mutagenicity:

Teratogenicity:

TLV:

Other Chronic Effects:

p-Aminobiphenyl

CHEMICAL NAME

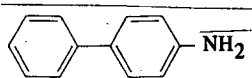
CAS Number: 92-67-1

Chemical Name: 4-Biphenylamine

Synonyms: 4-Aminobiphenyl
Xenylamine
p-Biphenylamine

**Molecular
Formula:** $C_{12}H_{11}N$

Structure:



Chemical Properties

Molecular Weight: 169.23

Physical State: Solid

Vapor Pressure:

Vapor Density:

Boiling Point: 302°C

Melting Point: 53 to 54°C

Density: 1.160 at 20°C/20°C

Solubility: Slightly (H_2O)

**Octanol/Water Parti-
tion Coefficient:**

Environmental Persistence

BOD:

Atmospheric Reactivity:

Safety Hazard:

Disaster hazard—moderate—toxic fumes

p-Aminobiphenyl

PRODUCTION DATA

Annual U.S.
Production:

Consumption:

Fraction of
Dispersion:

Fraction of Pro-
duction Lost:

Release Rate (million lbs/yr):

TOXICITY DATA

Acute Toxicity

LD₅₀

LD_{Lo}

Dosage

500 mg/kg

250 mg/kg

Animal

rat

mouse

Route

oral

intraperitoneal

Chronic Toxicity

U.S. Occupational Standard: (air) Carcinogen

Carcinogenicity:

TD_{Lo}

5200 mg/kg (52 weeks)

rat

oral

4 g/kg

rat

subcutaneous

140 mg/kg (7 days)

mouse

oral

216 mg/kg (3 days)

mouse

subcutaneous

680 mg/kg (3 years)

dog

oral

Mutagenicity:

Teratogenicity:

TLV:

Other Chronic Effects:

Aminoethylethanolamine

CHEMICAL NAME

CAS Number: 111-41-1

Chemical Name: 2[(2-Aminoethyl)amino]ethanol

Synonyms: (2-Hydroxyethyl)-ethylenediamine
N-Aminoethylethanolamine

Molecular
Formula: $C_4H_{12}N_2O$

Structure:
$$H_2N - CH_2 - CH_2 - \overset{H}{N} - CH_2 - CH_2OH$$

Chemical Properties

Molecular Weight: 104.16

Physical State: Liquid

Vapor Pressure: <0.01 mm at 20°C

Vapor Density: 3.59

Boiling Point: 243.7°C at 760 mm

Melting Point:

Density: 0.9556 at 25°C/4°C

Solubility: Very soluble (H_2O)

Octanol/Water Parti-
tion Coefficient:

Environmental Persistence

BOD:

Atmospheric Reactivity: Reacts with oxidizing materials

Safety Hazard:

Fire—moderate

Aminoethylethanolamine

PRODUCTION DATA

Annual U.S.
Production: 13.5×10^6 lbs.

Consumption:

Fraction of
Dispersion: 0.01

Fraction of Pro-
duction Lost: 0.015

Release Rate (million lbs/yr):

TOXICITY DATA

Acute Toxicity
LD₅₀

Dosage
3000 mg/kg
1800 mg/kg

Animal
rat
guinea pig

Route
oral
skin

Chronic Toxicity

U.S. Occupational Standard:

Carcinogenicity:

Mutagenicity:

Teratogenicity:

TLV:

Other Chronic Effects:

Amyl acetate

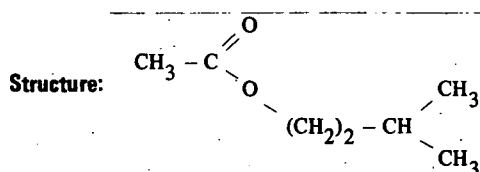
CHEMICAL NAME

CAS Number: 628-63-7

Chemical Name: Acetic acid, pentyl ester

Synonyms: Isoamyl acetate Acetic acid
Banana oil Amylacetate ester
Isopentyl acetate

Molecular
Formula: $C_7H_{14}O_2$



Chemical Properties

Molecular Weight: 130.2

Physical State: Liquid

Vapor Pressure: 9.43 mm at 25°C

Vapor Density: 4.49

Boiling Point: 142.0°C

Melting Point:

Density: 0.876 at 15°C/4°C

Solubility: Slight (H_2O)

Octanol/Water Partition Coefficient:

Environmental Persistence

BOD: 38% of theoretical

Atmospheric Reactivity: Reacts with reducing material

Safety Hazard: Fire—dangerous
Explosion—moderate

Amyl acetate

PRODUCTION DATA

Annual U.S.
Production: 7.6×10^6 lbs. (1964)

Consumption:

Fraction of
Dispersion: 1.0

Fraction of Pro-
duction Lost: 0.015

Release Rate (million lbs/yr): 7.6

TOXICITY DATA

Acute Toxicity	Dosage	Animal	Route
LD ₅₀	7400 mg/kg	rabbit	oral
LDLo	1500 mg/kg	rat	intraperitoneal
LCLo	5200 ppm		

Non-lethal Acute Effects:

Irritant effects—TCLo 188 ppm (30 minutes) human inhalation

Inhalation—1000 ppm (30 minutes) causes headache, fatigue, chest oppression

Irritant—eyes and mucous membranes of nose and throat

Chronic Toxicity

U.S. Occupational Standard: TWA: 125 ppm

Carcinogenicity:

Mutagenicity:

Teratogenicity:

TLV: 100 ppm

Other Chronic Effects:

Amyl alcohol

CHEMICAL NAME

CAS Number: 71-41-0

Chemical Name: Pentyl alcohol

Synonyms: n-Amyl alcohol

Butyl carbinol

1-Pentanol

Primary amyl alcohol

Molecular

Formula: $C_5H_{12}O$

Structure: $CH_3 - CH_2 - CH_2 - CH_2 - CH_2OH$

Chemical Properties

Molecular Weight: 88.15

Physical State: Liquid

Vapor Pressure: 3.16 mm at 25°C

Vapor Density: 3.04

Boiling Point: 137.3°C at 748 mm

Melting Point: -79°C

Density: 0.8144 at 20°C/4°C

Solubility: Insoluble (H_2O)

Octanol/Water Partition Coefficient: 1.56

Environmental Persistence

BOD: 84% of theoretical

Atmospheric Reactivity:

Safety Hazard:

Fire—moderate

Explosion—moderate

Amyl alcohol

PRODUCTION DATA

Annual U.S.
Production:

Consumption:

Fraction of
Dispersion:

Fraction of Pro-
duction Lost:

Release Rate (million lbs/yr):

TOXICITY DATA

Acute Toxicity	Dosage	Animal	Route
LD ₅₀	3030 mg/kg	rat	oral
	200 mg/kg	mouse	oral
LD _{Lo}	490 mg/kg	rat	intraperitoneal
	1390 mg/kg	dog	oral

Non-lethal Acute Effects:

Vapor irritant to eyes and upper respiratory tract

Following ingestion:

Headache

Nausea and vomiting

Delirium

Methemoglobin formation

Chronic Toxicity

U.S. Occupational Standard:

Carcinogenicity:

Mutagenicity:

Teratogenicity:

TLV: 100 ppm

Other Chronic Effects:

Amylamines

CHEMICAL NAME

CAS Number: 110-58-7

Chemical Name: Pentylamine

Synonyms: Amylamine (mixed isomers)
Aminopentane

**Molecular
Formula:** $C_5H_{13}N$

Structure: $CH_3(CH_2)_3CH_2NH_2$ (1 - amyl amine)

Chemical Properties (1-amyamine)

Molecular Weight: 87.17

Physical State: Liquid

Vapor Pressure:

Vapor Density: 3.01

Boiling Point: 104.4°C at 760 mm

Melting Point: -55°C

Density: 0.7547 at 20°C/4°C

Solubility: Infinite (H₂O)

**Octanol/Water Parti-
tion Coefficient:** 1.05

Environmental Persistence

BOD:

Atmospheric Reactivity: Reacts with oxidizing materials

Safety Hazard:

Fire—dangerous

Amylamines

PRODUCTION DATA

Annual U.S.
Production:

Consumption:

Fraction of
Dispersion:

Fraction of Pro-
duction Lost:

Release Rate (million lbs/yr):

TOXICITY DATA

Acute Toxicity
(Mixed isomers)

Dosage

Animal

Route

LD₅₀

470 mg/kg

rat

oral

1120 mg/kg

rabbit

skin

LC_{Lo}

2000 ppm (4 hours)

rat

inhalation

Chronic Toxicity

U.S. Occupational Standard:

Carcinogenicity:

Mutagenicity:

Teratogenicity:

TLV:

Other Chronic Effects:

Strong irritant

Amyl chloride

CHEMICAL NAME

CAS Number: 543-59-7

Chemical Name: 1-Chloropentane

Synonyms: n-Amylchloride
Pentylchloride

**Molecular
Formula:** $C_5H_{11}Cl$

Structure: $CH_3(CH_2)_3CH_2Cl$

Chemical Properties

Molecular Weight: 106.6

Physical State: Liquid

Vapor Pressure: 10 mm at 25°C

Vapor Density: 3.67

Boiling Point: 107.8°C at 760 mm

Melting Point: -99°C

Density: 0.8818 at 20°C/4°C

Solubility: Insoluble (H_2O)

**Octanol/Water Parti-
tion Coefficient:**

Environmental Persistence

BOD:

Atmospheric Reactivity: Reacts with oxidizing materials

Safety Hazard:

Fire—dangerous

Explosion—explosive if mixed with air

Disaster hazard—dangerous—emits toxic fumes (phosgene)

Amyl chloride

PRODUCTION DATA

Annual U.S.
Production:

Consumption:

Fraction of
Dispersion:

Fraction of Pro-
duction Lost:

Release Rate (million lbs/yr):

TOXICITY DATA

Acute Toxicity

Dosage

Animal

Route

Non-lethal Acute Effects:

Narcotic

Local irritant

Chronic Toxicity

U.S. Occupational Standard:

Carcinogenicity: Not tested

Mutagenicity:

Teratogenicity:

TLV:

Other Chronic Effects:

Amylene

CHEMICAL NAME

CAS Number: 109-67-1

Chemical Name: 1-Pentene

Synonyms: Amylene
Propyl ethylene

**Molecular
Formula:** C₅H₁₀

Structure: $\text{CH}_3\text{CH}_2\text{CH}_2\text{CH} = \text{CH}_2$

Chemical Properties

Molecular Weight: 70.14

Physical State: Gas-liquid

Vapor Pressure: 858.0 mm at 25°C

Vapor Density: 2.42

Boiling Point: 29.968°C at 760 mm

Melting Point: -138°C

Density: 0.6405 at 20°C/4°C

Solubility: Insoluble (H₂O)

**Octanol/Water Parti-
tion Coefficient:**

Environmental Persistence

BOD:

Atmospheric Reactivity: Reacts with oxidizing materials

Safety Hazard:

Fire—dangerous toxic fumes

PRODUCTION DATA

**Annual U.S.
Production:**

Consumption:

**Fraction of
Dispersion:**

**Fraction of Pro-
duction Lost:**

Release Rate (million lbs/yr):

TOXICITY DATA

Acute Toxicity

Dosage

Animal

Route

Non-lethal Acute Effects:

Narcotic
Asphyxiant

Chronic Toxicity

U.S. Occupational Standard:

Carcinogenicity:

Mutagenicity:

Teratogenicity:

TLV:

Other Chronic Effects:

Amyl ether

CHEMICAL NAME

CAS Number: 693-65-2

Chemical Name: Pentyl ether

Synonyms: Amyl ether
Amyl oxide
Diamyl ether

**Molecular
Formula:** $C_{10}H_{22}O$

Structure: $CH_3(CH_2)_4 - O - (CH_2)_4 - CH_3$

Chemical Properties

Molecular Weight: 158.29

Physical State: Liquid

Vapor Pressure:

Vapor Density: 5.46

Boiling Point: 190°C at 760 mm

Melting Point: -69°C

Density: 0.7833 at 20°C/4°C

Solubility: Insoluble (H₂O)

**Octanol/Water Parti-
tion Coefficient:**

Environmental Persistence

BOD:

Atmospheric Reactivity: Reacts with oxidizing materials

Safety Hazard:

Fire-moderate

PRODUCTION DATA

**Annual U.S.
Production:**

Consumption:

**Fraction of
Dispersion:**

**Fraction of Pro-
duction Lost:**

Release Rate (million lbs/yr):

TOXICITY DATA

Acute Toxicity

Dosage

Animal

Route

Chronic Toxicity

U.S. Occupational Standard:

Carcinogenicity:

Mutagenicity:

Teratogenicity:

TLV:

Other Chronic Effects:

Amyl mercaptan

CHEMICAL NAME

CAS Number: 110-66-7

Chemical Name: 1-Pentane thiol

Synonyms: Amyl mercaptan Amyl thio alcohol
Amyl hydrosulfide n-Thioamyl alcohol
Amy sulfhydrate

**Molecular
Formula:** $C_5H_{12}S$

Structure: $CH_3CH_2CH_2CH_2CH_2SH$

Chemical Properties

Molecular Weight: 104.22

Physical State: Liquid

Vapor Pressure: 13.8 mm at 25°C

Vapor Density: 3.59

Boiling Point: 112.9°C at 760 mm

Melting Point: -169°C

Density: 0.8327 at 20°C/4°C

Solubility: Insoluble (H_2O)

**Octanol/Water Parti-
tion Coefficient:**

Environmental Persistence

BOD:

Atmospheric Reactivity: Reacts with oxidizing materials

Safety Hazard:
Fire—dangerous

Amyl mercaptan

PRODUCTION DATA

Annual U.S.
Production:

Consumption:

Fraction of
Dispersion:

Fraction of Pro-
duction Lost:

Release Rate (million lbs/yr):

TOXICITY DATA

Acute Toxicity
 LC_{Lo}

Dosage
2000 ppm (4 hours)

Animal
rat

Route
inhalation

Non-lethal Acute Effects:

Irritant to eyes and skin

Chronic Toxicity

U.S. Occupational Standard:

Carcinogenicity:

Mutagenicity:

Teratogenicity:

TLV:

Other Chronic Effects:

Aniline

CHEMICAL NAME

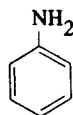
CAS Number: 62-53-3

Chemical Name: Aniline

Synonyms: Benzenamine Benamine
Aminobenzene Phylamine
Aminophen Aniline oil

**Molecular
Formula:** C_6H_7N

Structure:



Chemical Properties

Molecular Weight: 93.12

Physical State: Liquid

Vapor Pressure: 0.67 mm at 25°C

Vapor Density: 3.22

Boiling Point: 184°C

Melting Point: -6.3

Density: 1.02173 at 20°C/4°C

Solubility: Soluble, (36.5 gm/l of H₂O)

**Octanol/Water Parti-
tion Coefficient:** 1.24

Environmental Persistence

BOD:

Atmospheric Reactivity: Reacts with oxidizing materials

Photochemically degraded to n-methylaniline, N,N-dimethylaniline, acetanilide, phenol, and isomeric hydroxyanilines

Safety Hazard: Fire—moderate
Disaster hazard—dangerous, toxic fumes

PRODUCTION DATA

Annual U.S.

Production: 551.2 x 10⁶ lbs. (1974)

Consumption: 480 x 10⁶ lbs.

Fraction of

Dispersion: 0.01

Fraction of Pro-

duction Lost: 0.015

Release Rate (million lbs/yr): 6.15

TOXICITY DATA

Acute Toxicity	Dosage	Animal	Route
LD ₅₀	440 mg/kg	rat	oral
	1400 mg/kg	rat	skin
	420 mg/kg	rat	intraperitoneal
	820 mg/kg	rabbit	skin
	64 mg/kg	rabbit	intravenous
	1290 mg/kg	guinea pig	skin
LD _{Lo}	350 mg/kg	human	oral
	357 mg/kg	human	unreported
	480 mg/kg	mouse	subcutaneous
	1540 mg/kg	dog	skin
	1750 mg/kg	cat	oral
	1540 mg/kg	cat	skin
	200 mg/kg	rabbit	intraperitoneal
	1000 mg/kg	rabbit	subcutaneous
	1750 mg/kg	guinea pig	skin
LC ₅₀	175 ppm (7 hours)	mouse	inhalation
LC _{Lo}	250 ppm (4 hours)	rat	inhalation
	180 ppm (8 hours)	cat	inhalation

Chronic Toxicity

U.S. Occupational Standard: TWA: 5 ppm

Carcinogenicity: Neoplastic effects—papillomas and benign bladder tumors

Mutagenicity:

Teratogenicity:

TLV: 5 ppm (skin)

Odor perception: 0.23 ppm

Other Chronic Effects: Nephrosis
Blood effects
Enzyme and hormone effects

Aniline hydrochloride

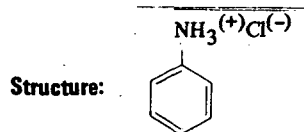
CHEMICAL NAME

CAS Number: 142-04-1

Chemical Name: Aniline hydrochloride

Synonyms: Aniline chloride
Benzenamine hydrochloride
Aniline salt

Molecular Formula: $C_6H_7N \cdot HCl$



Chemical Properties

Molecular Weight: 129.60

Physical State: Solid

Vapor Pressure:

Vapor Density: 1.22 at 4°C

Boiling Point: 245°C at 760 mm

Melting Point: 198°C

Density: 1.2215 at 4°C

Solubility: Very soluble (H₂O)

Octanol/Water Partition Coefficient:

Environmental Persistence

BOD:

Atmospheric Reactivity: Reacts with oxidizing materials

Safety Hazard:

Fire—slight

Disaster hazard—dangerous if heated with acid; aniline and chlorine fumes

Aniline hydrochloride

PRODUCTION DATA

Annual U.S.
Production:

Consumption:

Fraction of
Dispersion:

Fraction of Pro-
duction Lost:

Release Rate (million lbs/yr):

TOXICITY DATA

Acute Toxicity	Dosage	Animal	Route
LD ₅₀	1072 mg/kg	rat	oral
	841 mg/kg	mouse	oral
	750 mg/kg	mouse	intraperitoneal
LD _{Lo}	500 mg/kg	rat	intraperitoneal

Non-lethal Acute Effects:

Depression of CNS
Methemoglobinemia
Hemolysis
Stimulation of bone marrow
Liver damage

Chronic Toxicity

U.S. Occupational Standard:

Carcinogenicity:

Mutagenicity:

Teratogenicity:

TLV:

Other Chronic Effects:

Anisidine (o and p)

CHEMICAL NAME

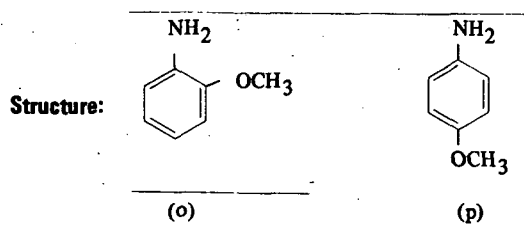
CAS Number: 104-97-9(p); 90-04-0(o)

Chemical Name: p-Anisidine; o-Anisidine

Synonyms: o-Methoxyaniline
o-Aminoanisole

p-Methoxyaniline
p-Aminoanisole
1-Amino-4-methoxybenzene

**Molecular
Formula:** C₇H₉NO



Chemical Properties

Molecular Weight: 123.16

Physical State: Liquid

Vapor Pressure: 1 mm at 61°C

Vapor Density:

Boiling Point: 224°C at 760 mm (o)
243°C at 760 mm (p)

Melting Point: 6.22°C (o)
57.2°C (p)

Density: 1.0923 at 20°C/4°C (o)
1.071 at 57°C/4°C (p)

Solubility: Soluble (H₂O) (p)
Slightly soluble (H₂O) (o)

**Octanol/Water Parti-
tion Coefficient:** 1.65

Environmental Persistence

BOD:

Atmospheric Reactivity:

Safety Hazard:

Disaster hazard—dangerous, toxic fumes

Anisidine (o and p)

PRODUCTION DATA

Annual U.S.
Production: 2.3×10^6 lbs. (total, 1967)

Consumption:

Fraction of
Dispersion:

Fraction of Pro-
duction Lost: 0.015

Release Rate (million lbs/yr):

TOXICITY DATA

Acute Toxicity (p-Anisidine) LD ₅₀	Dosage	Animal	Route
	1400 mg/kg	rat	oral
	1400 mg/kg	rat	intraperitoneal

Non-lethal Acute Effects:
Irritant to skin

Chronic Toxicity

U.S. Occupational Standard: (o and p isomers) (air) TWA 0.5 mg/m^3

Carcinogenicity:

Mutagenicity:

Teratogenicity: TLV: 0.5 mg/m^3 (skin)

Other Chronic Effects:

Anisole

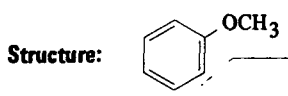
CHEMICAL NAME

CAS Number: 100-66-3

Chemical Name: Anisole

Synonyms: Methoxy benzene
Methylphenyl ether
Phenylmethyl ether

**Molecular
Formula:** C₇H₈O



Chemical Properties

Molecular Weight: 108.15

Physical State: Liquid

Vapor Pressure: 10 mm at 42.2°C

Vapor Density: 3.72

Boiling Point: 155°C

Melting Point: -37.5°C

Density: 0.9961 at 20°C/4°C

Solubility: Insoluble (H₂O)

**Octanol/Water Parti-
tion Coefficient:** 2.27

Environmental Persistence

BOD:

Atmospheric Reactivity: Reacts towards oxidizing materials

Safety Hazard:

Fire—moderate

Anisole

PRODUCTION DATA

Annual U.S.
Production:

Consumption:

Fraction of
Dispersion:

Fraction of Pro-
duction Lost:

Release Rate (million lbs/yr):

TOXICITY DATA

Acute Toxicity
LD₅₀

Dosage
3700 mg/kg
2800 mg/kg

Animal
rat
mouse

Route
oral
oral

Chronic Toxicity

U.S. Occupational Standard:

Carcinogenicity: Not tested

Mutagenicity:

Teratogenicity:

TLV:

Other Chronic Effects:

Anthranilic acid

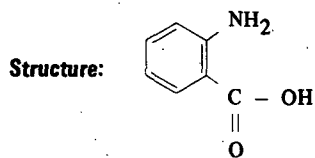
CHEMICAL NAME

CAS Number: 118-92-3

Chemical Name: Anthranilic acid

Synonyms: O-Aminobenzoic acid
2-Aminobenzoic acid

Molecular
Formula: $C_7H_7NO_2$



Chemical Properties

Molecular Weight: 137.14

Vapor Pressure:

Boiling Point: Sublimes

Density: 1.412 at 20°C

Octanol/Water Parti-
tion Coefficient: 1.73

Physical State: Solid

Vapor Density:

Melting Point: 146 to 147°C

Solubility: Soluble (H₂O)
Very soluble (hot H₂O)

Environmental Persistence

BOD:

Atmospheric Reactivity:

Safety Hazard:
Fire—slight

Anthranilic acid

PRODUCTION DATA

Annual U.S.
Production:

Consumption:

Fraction of
Dispersion:

Fraction of Pro-
duction Lost:

Release Rate (million lbs/yr):

TOXICITY DATA

Acute Toxicity
LD₅₀

Dosage
4620 mg/kg

Animal
rat

Route
oral

Chronic Toxicity

U.S. Occupational Standard:

Carcinogenicity:
Neoplastic effects:
TD_{Lo}

16 gm/kg (24 weeks) rat

oral

Mutagenicity:

Teratogenicity:

TLV:

Other Chronic Effects:

Anthraquinone

CHEMICAL NAME

CAS Number: 84-65-1

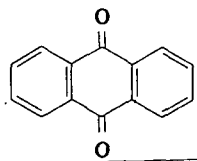
Chemical Name: Anthraquinone

Synonyms: 9,10-Dihydro-9,10-dioxoanthracenedione

Molecular

Formula: $C_{14}H_8O_2$

Structure:



Chemical Properties

Molecular Weight: 208.23

Vapor Pressure: 1 mm at 190°C

Boiling Point: 379.8°C at 760 mm

Density: 1.438 at 4°C

Octanol/Water Partition Coefficient:

Physical State: Solid

Vapor Density: 7.16

Melting Point: 286°C (sublimes)

Solubility: Insoluble (H₂O)

Environmental Persistence

BOD:

Atmospheric Reactivity:

Safety Hazard:

Fire—slight

Anthraquinone

PRODUCTION DATA

Annual U.S.
Production:

Consumption:

Fraction of
Dispersion:

Fraction of Pro-
duction Lost:

Release Rate (million lbs/yr):

TOXICITY DATA

Acute Toxicity

Dosage

Animal

Route

Non-lethal Acute Effects:

Weak sensitizer—dermatitis

Chronic Toxicity

U.S. Occupational Standard:

Carcinogenicity:

Neoplastic effects:

TD_{Lo}

90 gm/kg (90 days)

rat

oral

Mutagenicity:

Teratogenicity:

TLV:

Other Chronic Effects:

Auramine

CHEMICAL NAME

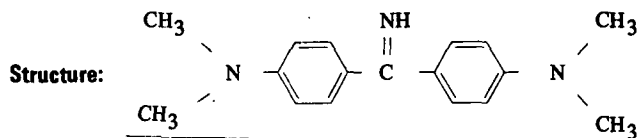
CAS Number: 492-80-8

Chemical Name: 4,4'-Carbonimidoylbis[N,N-dimethyl] benzenamine

Synonyms: 4,4'-(Imidocarbonyl)bis(N,N-dimethyl)aniline Apyonin
Auramine
Bis(p-dimethylamine-phenyl)methyleneimine

Molecular

Formula: $C_{17}H_{21}N_3$



Chemical Properties

Molecular Weight: 267.38

Physical State: Solid

Vapor Pressure:

Vapor Density:

Boiling Point:

Melting Point: 136°C

Density:

Solubility: Insoluble (H₂O)

Octanol/Water Partition Coefficient:

Environmental Persistence

BOD:

Atmospheric Reactivity:

Safety Hazard:

Auramine

PRODUCTION DATA

Annual U.S.
Production:

Consumption:

Fraction of
Dispersion:

Fraction of Pro-
duction Lost:

Release Rate (million lbs/yr):

TOXICITY DATA

Acute Toxicity

Dosage

Animal

Route

Non-lethal Acute Effects:
Dermatitis

Chronic Toxicity

U.S. Occupational Standard:

Carcinogenicity:

Neoplastic effects:

TD_{Lo}

37 gm/kg (87 weeks
continuous)

rat

oral

2625 mg/kg (21 weeks
intermittent)

rat

subcutaneous

Carcinogenic effect:

TD_{Lo}

29 gm/kg (52 weeks
continuous)

mouse

oral

Mutagenicity:

Teratogenicity:

TLV:

Other Chronic Effects:

Azodrin

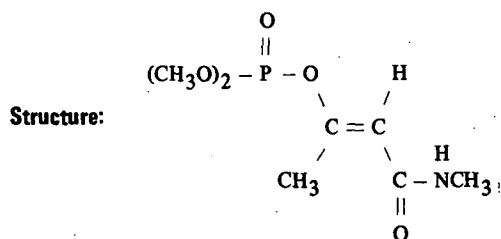
CHEMICAL NAME

CAS Number: 919-44-8

Chemical Name: Phosphoric acid, (Z)dimethyl-1-methyl-3(methylamino)-2-oxo-1-propenyl ester

Synonyms: Phosphoric acid, dimethyl ester with (E)-hydroxyl-N-methylcrotonamide
Dimethyl phosphate of 3-hydroxy-N-methyl-cis-crotonamide
Monocroptophos
O,O-dimethyl-O-(2-methylcarbamoyl-1-methyl-vinyl)-phosphate

Molecular Formula: $C_7H_{14}NO_5P$



Chemical Properties

Molecular Weight: 223.19

Physical State: Solid

Vapor Pressure:

Vapor Density:

Boiling Point: 125°C at 760 mm

Melting Point:

Density:

Solubility: Soluble (H₂O)

Octanol/Water Partition Coefficient:

Environmental Persistence

BOD:

Atmospheric Reactivity:

Safety Hazard:

Fire—dangerous

Disaster hazard—moderate—toxic fumes

PRODUCTION DATA

**Annual U.S.
Production:** 5 x 10⁶ lbs.

Consumption:

**Fraction of
Dispersion:** Estimated as 1.0

**Fraction of Pro-
duction Lost:** 0.015

Release Rate (million lbs/yr): 5

TOXICITY DATA

Acute Toxicity LD ₅₀	Dosage	Animal	Route
	2 mg/kg	rat	oral
	2 mg/kg	mouse	oral
	7 mg/kg	dog	oral
	11 mg/kg	rabbit	oral
	107 mg/kg	rabbit	skin

Chronic Toxicity

U.S. Occupational Standard:

Carcinogenicity:

Mutagenicity:

Teratogenicity:

TLV:

Other Chronic Effects:

Banvel D

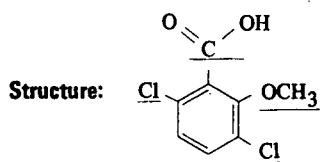
CHEMICAL NAME

CAS Number: 1918-00-9

Chemical Name: 3,6-Dichloro-2-methoxybenzoic acid

Synonyms: 3,6-Dichloro-o-anisic acid
Dicamba

Molecular Formula: $C_8H_6Cl_2O_3$



Chemical Properties

Molecular Weight: 221.04

Physical State: Solid

Vapor Pressure:

Vapor Density:

Boiling Point:

Melting Point: 114 to 116°C

Density:

Solubility: Slightly soluble (H₂O)

Octanol/Water Partition Coefficient:

Environmental Persistence

BOD:

Atmospheric Reactivity:

Safety Hazard:

Banvel D

PRODUCTION DATA

Annual U.S.
Production: 2×10^6 lbs.

Consumption:

Fraction of
Dispersion: Estimated as 1.0

Fraction of Pro-
duction Lost: 0.015

Release Rate (million lbs/yr): 2

TOXICITY DATA

Acute Toxicity
LD₅₀

Dosage
1040 mg/kg
1190 mg/kg
2000 mg/kg
3000 mg/kg
1000 mg/kg

Animal
rat
mouse
rabbit
guinea pig
mammal

Route
oral
oral
oral
oral
unknown

Chronic Toxicity

U.S. Occupational Standard:

Carcinogenicity:

Mutagenicity:

Teratogenicity:

TLV:

Other Chronic Effects:

Banvel T

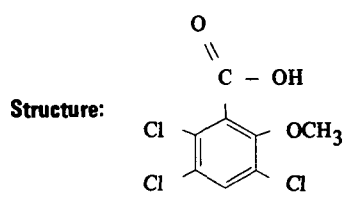
CHEMICAL NAME

CAS Number: 2307-49-5

Chemical Name: 3,5,6-Trichloro-o-anisic acid

Synonyms: Tricamba
2-Methoxy-3,5,6-trichlorobenzoic acid
3,5,6-Trichloro-2-methoxybenzoic acid

Molecular Formula: $C_8H_5O_3Cl_3$



Chemical Properties

Molecular Weight: 255.4

Physical State: Solid

Vapor Pressure:

Vapor Density:

Boiling Point:

Melting Point: 137 to 139°C

Density:

Solubility: Slightly soluble (H₂O)

Octanol/Water Partition Coefficient:

Environmental Persistence

BOD:

Atmospheric Reactivity:

Safety Hazard:

PRODUCTION DATA**Annual U.S.
Production:****Consumption:****Fraction of
Dispersion:****Fraction of Pro-
duction Lost:****Release Rate (million lbs/yr):****TOXICITY DATA****Acute Toxicity**
LD₅₀**Dosage**
300 mg/kg**Animal**
rat**Route**
oral**Chronic Toxicity****U.S. Occupational Standard:****Carcinogenicity:****Mutagenicity:****Teratogenicity:****TLV:****Other Chronic Effects:**

Benzaldehyde

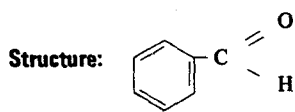
CHEMICAL NAME

CAS Number: 100-52-7

Chemical Name: Benzaldehyde

Synonyms: Benzoic aldehyde Benzenecarbonal
Artificial almond oil

**Molecular
Formula:** C₇H₆O



Chemical Properties

Molecular Weight: 106.12

Physical State: Liquid

Vapor Pressure: 1 mm at 26.2°C

Vapor Density: 3.65

Boiling Point: 179.0°C

Melting Point: -26°C

Density: 1.050 at 15°C/5°C

Solubility: slight (H₂O)

**Octanol/Water Parti-
tion Coefficient:** 2.41

Environmental Persistence

BOD: 50% of theoretical after 10 days

Atmospheric Reactivity: Reacts with oxidizing materials

Safety Hazard: Fire—slight

Benzaldehyde

PRODUCTION DATA

Annual U.S.**Production:** 3.4×10^6 lbs. (1968)**Consumption:****Fraction of****Dispersion:** 0.01**Fraction of Pro-****duction Lost:** 0.015**Release Rate (million lbs/yr):**

TOXICITY DATA

Acute ToxicityLD₅₀**Dosage**

1300 mg/kg

1000 mg/kg

5000 mg/kg

Animal

rat

guinea pig

rat

Route

oral

oral

subcutaneous

Non-lethal Acute Effects:

Dermatitis

Narcosis

Chronic Toxicity**U.S. Occupational Standard:****Carcinogenicity:** Not tested**Mutagenicity:****Teratogenicity:****TLV:****Other Chronic Effects:**

Benzamide

CHEMICAL NAME

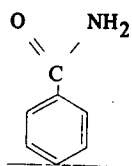
CAS Number: 55-21-0

Chemical Name: Benzamide

Synonyms: Benzoylamide
Benzoic acid, amide

**Molecular
Formula:** C_7H_7NO

Structure:



Chemical Properties

Molecular Weight: 121.14

Vapor Pressure:

Boiling Point: 290°C

Density: 1.0792 at 130°C/4°C

**Octanol/Water Parti-
tion Coefficient:** 0.96

Physical State: Solid

Vapor Density:

Melting Point: 132.5°C

Solubility: Slightly (H₂O)
Very soluble (hot H₂O)

Environmental Persistence

BOD:

Atmospheric Reactivity:

Safety Hazard:

Benzamide

PRODUCTION DATA

**Annual U.S.
Production:**

Consumption:

**Fraction of
Dispersion:**

**Fraction of Pro-
duction Lost:**

Release Rate (million lbs/yr):

TOXICITY DATA

Acute Toxicity
LD₅₀

Dosage
1160 mg/kg

Animal
mouse

Route
oral

Chronic Toxicity

U.S. Occupational Standard:

Carcinogenicity:

Mutagenicity:

Teratogenicity:

TLV:

Other Chronic Effects:

Benzene

CHEMICAL NAME

CAS Number: 71-43-2

Chemical Name: Benzene

Synonyms: Benzol
Phenylhydride
Coal naptha

**Molecular
Formula:** C_6H_6

Structure:



Chemical Properties

Molecular Weight: 78.11

Vapor Pressure: 95.9 mm at 25°C

Boiling Point: 80.1°C at 760 mm

Density: 0.8787 at 20°C/4°C

**Octanol/Water Parti-
tion Coefficient:** 2.28

Physical State: Liquid

Vapor Density: 2.77

Melting Point: 5.5°C

Solubility: Slightly (1.79 gm/l of H_2O)

Environmental Persistence

BOD: 1.9% theoretical

Atmospheric Reactivity:

Activity toward O_2 : slowly reacts with oxidizing materials

Activity toward OH: $t_{1/2}$ = 3 days

Photochemical Degradation: Not photoreactive; is biodegradable; found in air at 0.015-0.057 ppm

Safety Hazard:

Benzene

PRODUCTION DATA

Annual U.S.**Production:** 11,120 x 10⁶ lbs. (1974)**Consumption:** 1,562.8 x 10⁶ lbs.**Fraction of****Dispersion:** 0.01**Fraction of Pro-****duction Lost:** 0.01**Release Rate (million lbs/yr):** 183.6

TOXICITY DATA

Acute ToxicityLD₅₀**Dosage**

3800 mg/kg

4700 mg/kg

468 mg/kg

1150 mg/kg

LD_{Lo}

2000 mg/kg

530 mg/kg

1400 mg/kg

Animal

rat

mouse

mouse

rat

dog

guinea pig

frog

Route

oral

oral

intraperitoneal

intraperitoneal

oral

intraperitoneal

subcutaneous

Non-lethal Acute Effects:Blood effects—TC_{Lo}:

Aplastic anemia

Local irritant—erythema, burning of skin

Narcotic action on CNS

Euphoria

human

inhalation

Chronic Toxicity**U.S. Occupational Standard:** (air) TWA 10 ppm**Carcinogenicity:** TD_{Lo} 1232 mg/kg (52 weeks) mouse skinImplicated in human leukemia, sarcoma**Mutagenicity:**

Positive pulmonary adenomas

Bone marrow cell breaks

Teratogenicity:

No information

TLV: 10 ppm

Other Chronic Effects:

Estrus cycle disorder

Liver, kidney and lung damage

Hormone alteration

Bone marrow hyperplasia

Benzenedisulfonic acid

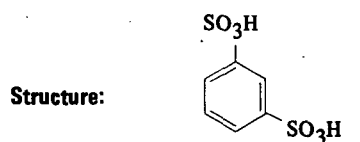
CHEMICAL NAME

CAS Number: 30496-93-6

Chemical Name: m-Benzene disulfonic acid

Synonyms: MBDSA

**Molecular
Formula:** $C_6H_6O_6S_2$



Chemical Properties

Molecular Weight: 238.23

Vapor Pressure:

Boiling Point:

Density:

**Octanol/Water Parti-
tion Coefficient:**

Physical State: Solid

Vapor Density:

Melting Point:

Solubility: Soluble (H₂O)

Environmental Persistence

BOD:

Atmospheric Reactivity:

Safety Hazard:

Benzenedisulfonic acid

PRODUCTION DATA

**Annual U.S.
Production:**

Consumption:

**Fraction of
Dispersion:**

**Fraction of Pro-
duction Lost:**

Release Rate (million lbs/yr):

TOXICITY DATA

Acute Toxicity

Dosage

Animal

Route

Non-lethal Acute Effects:
Irritant

Chronic Toxicity

U.S. Occupational Standard:

Carcinogenicity:

Mutagenicity:

Teratogenicity:

TLV:

Other Chronic Effects:

Benzenesulfonic acid

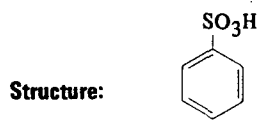
CHEMICAL NAME

CAS Number: 98-11-3

Chemical Name: Benzene sulfonic acid

Synonyms: Phenyl sulfonic acid

**Molecular
Formula:** $C_6H_5O_3S$



Chemical Properties

Molecular Weight: 158.12

Vapor Pressure:

Boiling Point:

Density:

**Octanol/Water Parti-
tion Coefficient:** -2.25

Physical State: Solid

Vapor Density:

Melting Point: 65 to 66°C (anhydrous)

Solubility: Very soluble (H₂O)

Environmental Persistence

BOD:

Atmospheric Reactivity:

Safety Hazard:

Benzenesulfonic acid

PRODUCTION DATA

**Annual U.S.
Production:**

Consumption:

**Fraction of
Dispersion:**

**Fraction of Pro-
duction Lost:**

Release Rate (million lbs/yr):

TOXICITY DATA

Acute Toxicity
LD₅₀

Dosage
2050 mg/kg

Animal
rat

Route
oral

Non-lethal Acute Effects:
Irritant

Chronic Toxicity

U.S. Occupational Standard:

Carcinogenicity:

Mutagenicity:

Teratogenicity:

TLV:

Other Chronic Effects:

Benzidine

CHEMICAL NAME

CAS Number: 92-87-5

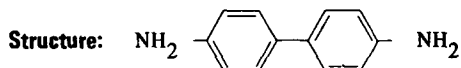
Chemical Name: Benzidine

Synonyms: [1,1'-Biphenyl]-4,4'-diamine
p-Diaminodiphenyl
4,4'-Biphenyldiamine

4,4'-Diaminodiphenyl
4,4'-Diphenylenediamine

Molecular

Formula: C₁₂H₁₂N₂



Chemical Properties

Molecular Weight: 184.24

Physical State: Solid

Vapor Pressure:

Vapor Density:

Boiling Point: 401.7°C

Melting Point: 122 and 125°C—unstable
128°C—stable

Density: 1.250 at 20°C/4°C

Solubility: Slightly soluble (H₂O)

Octanol/Water Partition Coefficient:

Environmental Persistence

BOD: 1.9 for 20 days at 20°C

Atmospheric Reactivity:

Reacts rapidly with RO₂ (t_{1/2} = 4 days)

O₃ (t_{1/2} = 1 day)

OH (t_{1/2} = 1 day)

Safety Hazard:

Benzidine

PRODUCTION DATA

Annual U.S.
Production: 10.4×10^6 lbs (1972)

Consumption:

Fraction of
Dispersion: 0.01

Fraction of Pro-
duction Lost: 0.015

Release Rate (million lbs/yr): .02

TOXICITY DATA

Acute Toxicity	Dosage	Animal	Route
LD ₅₀	309 mg/kg	rat	oral
	214 mg/kg	mouse	oral
LD _{Lo}	200 mg/kg	dog	oral
	400 mg/kg	dog	subcutaneous

Non-lethal Acute Effects:

Bone marrow depression
Liver and kidney damage
Blood hemolysis
Epicardial petechial hemorrhage

Chronic Toxicity

U.S. Occupational Standard:

Carcinogenicity:

TD _{Lo}	4500 mg/kg (30 days)	rat	oral
	2100 mg/kg (35 weeks)	rat	subcutaneous
	8 gm/kg (35 weeks)	mouse	subcutaneous
	75 gm/kg (3 years)	hamster	oral
TC _{Lo}	10 mg/m ³ (56 weeks)	rat	inhalation
	18 mg/m ³ (13 years)	human	inhalation

Mutagenicity:

Teratogenicity:

TLV:

Other Chronic Effects:

Benzil

CHEMICAL NAME

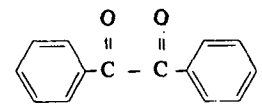
CAS Number: 134-81-6

Chemical Name: Benzil

Synonyms: Dibenzoyl
Diphenyl-ethanedione
Diphenylglyoxal

**Molecular
Formula:** $C_{14}H_{10}O_2$

Structure:



Chemical Properties

Molecular Weight: 210.2

Physical State: Solid

Vapor Pressure: 1 mm at 128.4°C

Vapor Density:

Boiling Point: 346 to 348°C (decomposes)

Melting Point: 95°C

Density: 1.084 at 102°C/4°C

Solubility: Insoluble (H_2O)

Octanol/Water Partition Coefficient:

Environmental Persistence

BOD:

Atmospheric Reactivity:

Safety Hazard:

PRODUCTION DATA**Annual U.S.
Production:****Consumption:****Fraction of
Dispersion:****Fraction of Pro-
duction Lost:****Release Rate (million lbs/yr):****TOXICITY DATA****Acute Toxicity**
LD₅₀**Dosage**
2710 mg/kg**Animal**
rat**Route**
oral**Chronic Toxicity****U.S. Occupational Standard:****Carcinogenicity:****Mutagenicity:****Teratogenicity:****TLV:****Other Chronic Effects:**

Benzilic acid

CHEMICAL NAME

CAS Number: 76-93-7

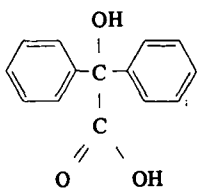
Chemical Name: Benzilic acid

Synonyms: Diphenyl glycolic acid
Diphenyl hydroxyacetic acid
 α -Hydroxy- α -phenylbenzene acetic acid

Molecular

Formula: $C_{14}H_{12}O_3$

Structure:



Chemical Properties

Molecular Weight: 228.24

Physical State: Solid

Vapor Pressure:

Vapor Density:

Boiling Point: 180°C (decomposes)

Melting Point: 148 to 151°C

Density:

Solubility: Slightly soluble (H_2O)

Octanol/Water Partition Coefficient: 2.32

Environmental Persistence

BOD:

Atmospheric Reactivity: Reacts with oxidizing materials

Safety Hazard:

Fire—slight

Benzilic acid

PRODUCTION DATA

**Annual U.S.
Production:**

Consumption:

**Fraction of
Dispersion:**

**Fraction of Pro-
duction Lost:**

Release Rate (million lbs/yr):

TOXICITY DATA

Acute Toxicity

Dosage

Animal

Route

Chronic Toxicity

U.S. Occupational Standard:

Carcinogenicity:

Mutagenicity:

Teratogenicity:

TLV:

Other Chronic Effects:

Benzoic acid

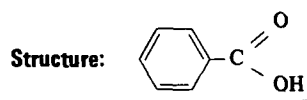
CHEMICAL NAME

CAS Number: 65-85-0

Chemical Name: Benzoic Acid

Synonyms: Phenyl formic acid
Benzene carboxylic acid
Carboxybenzene

Molecular
Formula: C₇H₆O₂



Chemical Properties

Molecular Weight: 122.12

Vapor Pressure: 1 mm at 96.0°C

Boiling Point: 249°C

Density: 1.2659 15°C/4°C

Octanol/Water Parti-
tion Coefficient: 2.00

Physical State: Solid

Vapor Density: 4.21

Melting Point: 122.4°C

Solubility: Slightly (H₂O)

Environmental Persistence

BOD: 46% of theoretical after 10 days

Atmospheric Reactivity: Reacts with oxidizing materials

Safety Hazard: Fire—slight

Benzoic acid

PRODUCTION DATA

Annual U.S. Production:	79 x 10 ⁶ lbs. (1974)	Consumption:	70 x 10 ⁶ lbs. (1975) (excludes phenol production) ca. 90 x 10 ⁶ lbs. (phenol production)
Fraction of Dispersion:	0.01	Fraction of Pro- duction Lost:	0.015
Release Rate (million lbs/yr):			

TOXICITY DATA

Acute Toxicity	Dosage	Animal	Route
LD ₅₀	3040 mg/kg	rat	oral
	2370 mg/kg	mouse	oral
	1460 mg/kg	mouse	intraperitoneal
Non-lethal Acute Effects:			
Skin effects-TD _{Lo}	6 mg/kg	human	skin
<u>Gastrointestinal effects</u>			

Chronic Toxicity

U.S. Occupational Standard:

Carcinogenicity:

Mutagenicity:

Teratogenicity:

TLV:

Other Chronic Effects:

Benzoin

CHEMICAL NAME

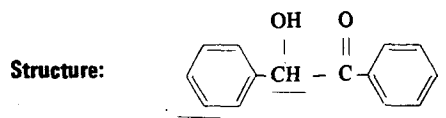
CAS Number: 119-53-9

Chemical Name: Benzoin

Synonyms: Phenyl α -hydroxybenzylketone
Phenylbenzoylcarbinol

Benzoylphenylcarbinol
2-Hydroxy-1,2-diphenylethanone

**Molecular
Formula:** $C_{14}H_{12}O_2$



Chemical Properties (dl mixture)

Molecular Weight: 212.24

Physical State: Solid

Vapor Pressure: 1 mm at 135.6°C

Vapor Density:

Boiling Point: 344°C at 768 mm

Melting Point: 137°C

Density: 1.310 at 20°C/4°C

Solubility: Slightly (hot H_2O)

Octanol/Water Partition Coefficient:

Environmental Persistence

BOD:

Atmospheric Reactivity:

Safety Hazard:
Fire—slight

Benzoin

PRODUCTION DATA

**Annual U.S.
Production:**

Consumption:

**Fraction of
Dispersion:**

**Fraction of Pro-
duction Lost:**

Release Rate (million lbs/yr):

TOXICITY DATA

Acute Toxicity

Dosage

Animal

Route

Chronic Toxicity

U.S. Occupational Standard:

Carcinogenicity:

Mutagenicity:

Teratogenicity:

TLV:

Other Chronic Effects:

Benzonitrile

CHEMICAL NAME

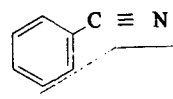
CAS Number: 100-47-0

Chemical Name: Benzonitrile

Synonyms: Phenyl cyanide
Benzoic acid, nitrile

**Molecular
Formula:** C₇H₅N

Structure:



Chemical Properties

Molecular Weight: 103.13

Vapor Pressure: 1 mm at 28.2°C

Boiling Point: 190.7°C

Density: 1.246 at 20°C/4°C

**Octanol/Water Parti-
tion Coefficient:** 1.56

Physical State: Liquid

Vapor Density:

Melting Point: -13.1°C

Solubility: Slightly (hot H₂O)

Environmental Persistence

BOD: 40% of theoretical

Atmospheric Reactivity:

Safety Hazard:

Benzonitrile

PRODUCTION DATA

**Annual U.S.
Production:**

Consumption:

**Fraction of
Dispersion:**

**Fraction of Pro-
duction Lost:**

Release Rate (million lbs/yr):

TOXICITY DATA

Acute Toxicity

LD₅₀

LD_{Lo}

LC_{Lo}

Dosage

1200 mg/kg

720 mg/kg

180 mg/kg

950 ppm (8 hours)

Animal

rat

rat

mouse

rat

Route

skin

oral

subcutaneous

inhalation

Chronic Toxicity

U.S. Occupational Standard:

Carcinogenicity: Not tested

Mutagenicity:

Teratogenicity:

TLV:

Other Chronic Effects:

Benzophenone

CHEMICAL NAME

CAS Number: 119-61-9

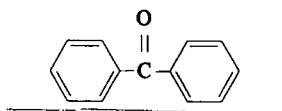
Chemical Name: Benzophenone

Synonyms: Phenyl ketone
Diphenyl ketone
Diphenyl methanone

Benzoylbenzene

**Molecular
Formula:** $C_{13}H_{10}O$

Structure:



Chemical Properties

Molecular Weight: 182.21

Physical State: Solid

Vapor Pressure: 1 mm at 108.2°C

Vapor Density:

Boiling Point: 305.9°C at 760 mm

Melting Point: 47.5°C (α)

Density: α form 1.0976 at 50°C/50°C

Solubility: Insoluble (H_2O)

**Octanol/Water Parti-
tion Coefficient:**

Environmental Persistence

BOD:

Atmospheric Reactivity: Reacts with oxidizing materials

Safety Hazard:

Fire—slight

Benzophenone

PRODUCTION DATA

**Annual U.S.
Production:**

Consumption:

**Fraction of
Dispersion:**

**Fraction of Pro-
duction Lost:**

Release Rate (million lbs/yr):

TOXICITY DATA

Acute Toxicity

Dosage

Animal

Route

Chronic Toxicity

U.S. Occupational Standard:

Carcinogenicity: Under test

Mutagenicity:

Teratogenicity:

TLV:

Other Chronic Effects:

Benzoquinone

CHEMICAL NAME

CAS Number: 106-51-4

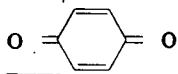
Chemical Name: p-Benzoquinone

Synonyms: Quinone
Chinone

1,4-Benzoquinone
Paraquinone

Molecular Formula: $C_6H_4O_2$

Structure:



Chemical Properties

Molecular Weight: 108.09

Physical State: Solid

Vapor Pressure:

Vapor Density:

Boiling Point: Sublimes

Melting Point: 115.7°C

Density: 1.318 at 20°C/4°C

Solubility: Soluble (H₂O)

Octanol/Water Partition Coefficient:

Environmental Persistence

BOD:

Atmospheric Reactivity:

Safety Hazard:

Disaster hazard—dangerous—toxic fumes

Benzoquinone

PRODUCTION DATA

Annual U.S.
Production:

Consumption:

Fraction of
Dispersion:

Fraction of Pro-
duction Lost:

Release Rate (million lbs/yr):

TOXICITY DATA

Acute Toxicity	Dosage	Animal	Route
LD ₅₀	130 mg/kg	rat	oral
	8500 mg/kg	mouse	intraperitoneal
	25 mg/kg	rat	intravenous
LC _{Lo}	320 mg/m ³	mouse	inhalation

Non-lethal Acute Effects:

Irritation of skin and mucous membranes
Necrosis, if in eyes
Kidney damage
Dermatitis

Chronic Toxicity

U.S. Occupational Standard:

Carcinogenicity:

TD_{Lo}

2000 mg/kg (28 weeks)

mouse

skin

Mutagenicity:

Teratogenicity:

TLV: 0.1 mg/m³

Other Chronic Effects:

Benzotrichloride

CHEMICAL NAME

CAS Number: 98-07-7

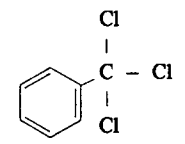
Chemical Name: α,α,α -Trichlorotoluene

Synonyms: Phenylchloroform

Molecular

Formula: $C_7H_5Cl_3$

Structure:



Chemical Properties

Molecular Weight: 195.48

Vapor Pressure: 1 mm at 45.8°C

Boiling Point: 220.6°C at 760 mm

Density: 1.38 at 15.5°C/15.5°C

Octanol/Water Partition Coefficient:

Physical State: Liquid

Vapor Density: 6.77

Melting Point: -4.75°C

Solubility: Insoluble (H_2O)

Environmental Persistence

BOD:

Atmospheric Reactivity:

Safety Hazard:

Benzotrichloride

PRODUCTION DATA

Annual U.S.
Production:

Consumption:

Fraction of
Dispersion:

Fraction of Pro-
duction Lost:

Release Rate (million lbs/yr):

TOXICITY DATA

Acute Toxicity
LD_{Lo}
LC_{Lo}

Dosage
2150 mg/kg
125 ppm (4 hours)

Animal
frog
rat

Route
unreported
inhalation

Chronic Toxicity

U.S. Occupational Standard:

Carcinogenicity: Not tested

Mutagenicity:

Teratogenicity:

TLV:

Other Chronic Effects:

Benzoyl chloride

CHEMICAL NAME

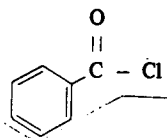
CAS Number: 98-88-4

Chemical Name: Benzoyl chloride

Synonyms: Benzene carbonyl chloride
 α -Chlorobenzaldehyde

**Molecular
Formula:** C_7H_5ClO

Structure:



Chemical Properties

Molecular Weight: 140.57

Physical State: Liquid

Vapor Pressure: 1 mm at 32.1°C

Vapor Density: 4.88

Boiling Point: 197.2°C at 760 mm

Melting Point: -0.5°C

Density: 1.2120 at 20°C/4°C

Solubility: Slightly (decomposes) (H_2O)

**Octanol/Water Parti-
tion Coefficient:**

Environmental Persistence

BOD: 40% of theoretical

Atmospheric Reactivity:

Safety Hazard:

Disaster hazard—dangerous—fumes of phosgene

Benzoyl chloride

PRODUCTION DATA

Annual U.S. Production: Estimated as 16.8×10^6 lbs. (1975) **Consumption:**

Fraction of Dispersion: **Fraction of Production Lost:** 0.015

Release Rate (million lbs/yr):

TOXICITY DATA

Acute Toxicity	Dosage	Animal	Route
LD ₅₀	2460 mg/kg	rat	oral
	790 mg/kg	rabbit	skin
TCLo	16 ppm	human	inhalation

Non-lethal Acute Effects:
Irritant to skin, eyes, and mucous membranes
Lachrymatory

Chronic Toxicity

U.S. Occupational Standard:

Carcinogenicity: Not tested

Mutagenicity:

Teratogenicity:

TLV:

Other Chronic Effects:

Benzyl alcohol

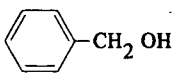
CHEMICAL NAME

CAS Number: 100-51-6

Chemical Name: Benzyl alcohol

Synonyms: Hydroxytoluene Benzenemethanol
Phenylcarbinol
 α -Hydroxytoluene

**Molecular
Formula:** C₇H₈O

Structure: 

Chemical Properties

Molecular Weight: 108.13

Physical State: Solid

Vapor Pressure: 1 mm at 58.0°C

Vapor Density: 3.72

Boiling Point: 205.35°C at 760 mm

Melting Point: -15.3°C

Density: 1.0419 at 20°C/4°C

Solubility: Soluble (H₂O)

**Octanol/Water Parti-
tion Coefficient:** 1.10

Environmental Persistence

BOD: 63% of theoretical

Atmospheric Reactivity: Reacts with oxidizing materials

Safety Hazard: Fire—slight

Benzyl alcohol

PRODUCTION DATA

Annual U.S.**Production:** 10.4×10^6 lbs. (1972)**Consumption:** 11.7×10^6 lbs. (1974)**Fraction of****Dispersion:** 0.01**Fraction of Pro-****duction Lost:** 0.015**Release Rate (million lbs/yr):**

TOXICITY DATA

Acute ToxicityLD₅₀**Dosage**

1230 mg/kg

64 mg/kg

1580 mg/kg

480 mg/kg

1940 mg/kg

100 mg/kg

LD_{Lo}

400 mg/kg

50 mg/kg

9 mg/kg

400 mg/kg

LC₅₀

1000 pm (8 hours)

Animal

rat

rat

mouse

mouse

rabbit

wild bird

rat

dog

dog

guinea pig

Route

oral

intravenous

oral

intravenous

oral

oral

intraperitoneal

intravenous

parenteral

intraperitoneal

Non-lethal Acute Effects:

Irritant—skin, mucous membranes

Headache, vertigo, nausea, vomiting, and diarrhea

Chronic Toxicity**U.S. Occupational Standard:****Carcinogenicity:****Mutagenicity:****Teratogenicity:****Other Chronic Effects:****TLV:**

Benzylamine

CHEMICAL NAME

CAS Number: 100-46-9

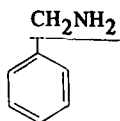
Chemical Name: Benzenemethanamine

Synonyms: Aminotoluene

Molecular

Formula: C_7H_9N

Structure:



Chemical Properties

Molecular Weight: 107.2

Physical State: Liquid

Vapor Pressure: 1 mm at 29°C

Vapor Density:

Boiling Point: 184.5°C

Melting Point:

Density: 0.983 at 19°C/4°C

Solubility: Very soluble (H₂O)

Octanol/Water Partition Coefficient: 1.14

Environmental Persistence

BOD:

Atmospheric Reactivity:

Safety Hazard:

Disaster hazard—moderate—toxic fumes

Benzylamine

PRODUCTION DATA

**Annual U.S.
Production:**

Consumption:

**Fraction of
Dispersion:**

**Fraction of Pro-
duction Lost:**

Release Rate (million lbs/yr):

TOXICITY DATA

Acute Toxicity

Dosage

Animal

Route

Chronic Toxicity

U.S. Occupational Standard:

Carcinogenicity: Not tested

Mutagenicity:

Teratogenicity:

TLV:

Other Chronic Effects:

Benzylbenzene

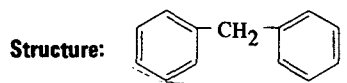
CHEMICAL NAME

CAS Number: 101-81-5

Chemical Name: Diphenyl methane

Synonyms: Benzylbenzene
Ditan

**Molecular
Formula:** $C_{13}H_{12}$



Chemical Properties

Molecular Weight: 168.23

Physical State: Solid

Vapor Pressure: 1 mm at 76.0°C

Vapor Density: 5.79

Boiling Point: 264.7°C at 760 mm

Melting Point: 26.5°C

Density: 1.0056 at 20°C/4°C

Solubility: Insoluble (H₂O)

**Octanol/Water Parti-
tion Coefficient:**

Environmental Persistence

BOD:

Atmospheric Reactivity: Reacts with oxidizing materials

Safety Hazard:
Fire—slight

Benzylbenzene

PRODUCTION DATA

**Annual U.S.
Production:**

Consumption:

**Fraction of
Dispersion:**

**Fraction of Pro-
duction Lost:**

Release Rate (million lbs/yr):

TOXICITY DATA

Acute Toxicity
LD_{Lo}

Dosage
5000 mg/kg

Animal
rat

Route
oral

Chronic Toxicity

U.S. Occupational Standard:

Carcinogenicity: Not tested

Mutagenicity:

Teratogenicity:

TLV:

Other Chronic Effects:

Benzy benzoate

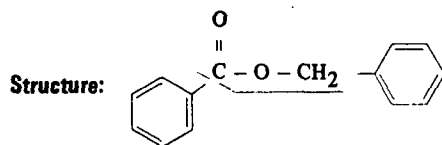
CHEMICAL NAME

CAS Number: 120-51-4

Chemical Name: Benzoic acid, benzyl ester

Synonyms: Benzyl benzoate
Benzyl benzene carboxylate

**Molecular
Formula:** $C_{14}H_{12}O_2$



Chemical Properties

Molecular Weight: 212.25

Physical State: Liquid

Vapor Pressure: <10 mm at 25°C

Vapor Density: 7.3

Boiling Point: 323 to 324°C at 760 mm

Melting Point: 21°C

Density: 1.1121 at 25°C/4°C

Solubility: Insoluble (H₂O)

**Octanol/Water Parti-
tion Coefficient:** 3.97

Environmental Persistence

BOD:

Atmospheric Reactivity: Reacts with oxidizing materials

Safety Hazard:
Fire—slight

Benzyl benzoate

PRODUCTION DATA

Annual U.S.
Production:

Consumption:

Fraction of
Dispersion:

Fraction of Pro-
duction Lost:

Release Rate (million lbs/yr):

TOXICITY DATA

Acute Toxicity
LD₅₀

Dosage
1700 mg/kg
1400 mg/kg
2240 mg/kg
1800 mg/kg
1000 mg/kg

Animal
rat
mouse
cat
rabbit
guinea pig

Route
oral
oral
oral
oral
oral

Chronic Toxicity

U.S. Occupational Standard:

Carcinogenicity:

Mutagenicity:

Teratogenicity:

TLV:

Other Chronic Effects:

Benzyl chloride

CHEMICAL NAME

CAS Number: 100-44-7

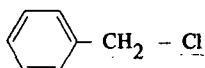
Chemical Name: α -Chlorotoluene

Synonyms: α -Tolylchloride

Molecular

Formula: C_7H_7Cl

Structure:



Chemical Properties

Molecular Weight: 126.5

Physical State: Liquid

Vapor Pressure: 1.4 mm at 25°C

Vapor Density: 4.36

Boiling Point: 179°C at 760 mm

Melting Point: -39°C

Density: 1.1026 at 18°C/4°C

Solubility: Insoluble (H_2O)

Octanol/Water Partition Coefficient:

Environmental Persistence

BOD:

Atmospheric Reactivity: Reacts with oxidizing agents

Safety Hazard: Fire—moderate
Explosion—moderate (violent with metals)
Disaster hazard—dangerous fumes

Benzyl chloride

PRODUCTION DATA

Annual U.S.
Production: 80.4×10^6 lbs. (1972)

Consumption: 90×10^6 lbs. (1974)

Fraction of
Dispersion: 0.01

Fraction of Pro-
duction Lost: 0.015

Release Rate (million lbs/yr):

TOXICITY DATA

Acute Toxicity	Dosage	Animal	Route
LD ₅₀	1231 mg/kg	rat	oral
	1000 mg/kg	rat	subcutaneous
	1624 mg/kg	mouse	oral

Non-lethal Acute Effects:

Unspecified effects:			
TCLo	16 ppm	human	inhalation
Irritant—intensely to eyes, skin, and mucous membranes			

Chronic Toxicity

U.S. Occupational Standard:

Carcinogenicity:

Neoplastic effects:

TDLo	2100 mg/kg (51 weeks)	rat	subcutaneous
------	-----------------------	-----	--------------

Mutagenicity:

Teratogenicity:

TLV:

Other Chronic Effects: CNS depression

Benzylidichloride

CHEMICAL NAME

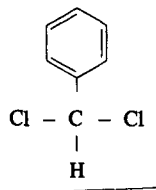
CAS Number: 98-87-3

Chemical Name: α,α -Dichlorotoluene

Synonyms: Benzyl dichloride (Dichloromethyl)-benzene
Benzal chloride
Benzylidene chloride

Molecular Formula: $C_7H_6Cl_2$

Structure:



Chemical Properties

Molecular Weight: 161.03

Vapor Pressure: <10 mm at 25°C

Boiling Point: 205.2°C at 760 mm

Density: 1.295 at 16°C

Octanol/Water Partition Coefficient:

Physical State: Liquid

Vapor Density:

Melting Point: -16.1°C

Solubility: Insoluble (H₂O)

Environmental Persistence

BOD:

Atmospheric Reactivity:

Safety Hazard:

Benzyldichloride

PRODUCTION DATA

**Annual U.S.
Production:**

Consumption:

**Fraction of
Dispersion:**

**Fraction of Pro-
duction Lost:**

Release Rate (million lbs/yr):

TOXICITY DATA

Acute Toxicity

LD₅₀

Dosage

3249 mg/kg

467 mg/kg

LC₅₀

200 mg/m³

Animal

rat

mouse

mouse

Route

oral

oral

inhalation

Non-lethal Acute Effects:

Irritant—strong lachrymator

Chronic Toxicity

U.S. Occupational Standard:

Carcinogenicity: Not tested

Mutagenicity:

Teratogenicity:

TLV:

Other Chronic Effects:

CNS depression

Bis (chloromethyl) ether

CHEMICAL NAME

CAS Number: 542-88-1

Chemical Name: Oxybis(chloromethane)

Synonyms: Dichloromethyl ether 1,1'-Dichlorodimethyl ether
Dichlorinated methyloxiide
BCME

**Molecular
Formula:** $C_2H_4Cl_2O$

Structure: $ClCH_2 - O - CH_2Cl$

Chemical Properties

Molecular Weight: 114.96

Physical State: Liquid (volatile)

Vapor Pressure:

Vapor Density: 4.0

Boiling Point: 104°C at 760 mm

Melting Point: -41.5°C

Density: 1.328 at 15°C/4°C

Solubility: Decomposes (H_2O)

**Octanol/Water Parti-
tion Coefficient:**

Environmental Persistence

BOD:

Atmospheric Reactivity:

Safety Hazard:

Bis (chloromethyl) ether

PRODUCTION DATA

Annual U.S.
Production:

Consumption:

Fraction of
Dispersion:

Fraction of Pro-
duction Lost:

Release Rate (million lbs/yr):

TOXICITY DATA

Acute Toxicity

LD₅₀

TD_{Lo}

Dosage

210 mg/kg

280 mg/kg

315 mg/kg (245 days,
intermittent)

11 gm/kg (47 weeks,
intermittent)

Animal

rat

rabbit

rat

mouse

Route

oral

skin

subcutaneous

skin

Chronic Toxicity

U.S. Occupational Standard:

Carcinogenicity: Carcinogen, produces tumors

Mutagenicity:

Teratogenicity:

TLV:

Other Chronic Effects:

Bisphenol A

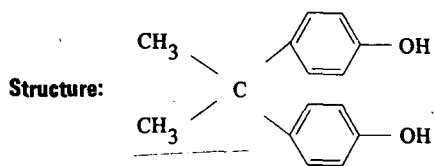
CHEMICAL NAME

CAS Number: 80-05-7

Chemical Name: 4,4'-Isopropylidenediphenol

Synonyms: Bisphenol A
2,2-Bis (4-hydroxyphenyl) propane
4,4'-Isopropylidenediphenol

Molecular Formula: C₁₅H₁₆O₂



Chemical Properties

Molecular Weight: 228.29

Vapor Pressure: 1 mm at 193°C

Boiling Point: 250 to 252°C

Density:

Octanol/Water Partition Coefficient:

Physical State: Solid

Vapor Density:

Melting Point: 153°C

Solubility: Insoluble (H₂O)

Environmental Persistence

BOD:

Atmospheric Reactivity:

Safety Hazard:

Bisphenol A

PRODUCTION DATA

**Annual U.S.
Production:** 370.4×10^6 lbs. (1974)

Consumption: 370.4×10^6 lbs.

**Fraction of
Dispersion:** 0.01

**Fraction of Pro-
duction Lost:** 0.015

Release Rate (million lbs/yr):

TOXICITY DATA

Acute Toxicity
LD₅₀

Dosage
150 mg/kg

Animal
rat

Route
intraperitoneal

Chronic Toxicity

U.S. Occupational Standard: (air) TWA: 0.5 ppm

Carcinogenicity:

Mutagenicity:

Teratogenicity:

Other Chronic Effects:

TLV:
(odorless)

Bromacil

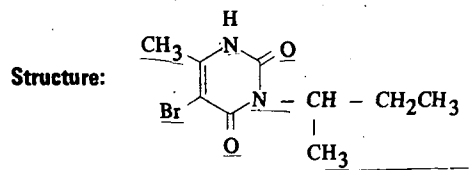
CHEMICAL NAME

CAS Number: 314-42-1

Chemical Name: 5-Bromo-3-sec butyl-6-methyluracil

Synonyms: Borea

Molecular
Formula: $C_9H_{13}BrN_2O_2$



Chemical Properties

Molecular Weight: 247.12

Physical State: Solid

Vapor Pressure:

Vapor Density:

Boiling Point: Sublimes

Melting Point: 158 to 159°C

Density:

Solubility: 815 ppm at 25°C

Octanol/Water Parti-
tion Coefficient:

Environmental Persistence

BOD:

Atmospheric Reactivity:

Safety Hazard:

Bromacil

PRODUCTION DATA

Annual U.S.**Production:** 8×10^6 lbs.**Consumption:****Fraction of****Dispersion:** Estimated as 1.0**Fraction of Pro-****duction Lost:** 0.015**Release Rate (million lbs/yr):** 8

TOXICITY DATA

Acute ToxicityLD₅₀LD_{Lo}**Dosage**

3400 mg/kg

3750 mg/kg

Animal

rat

mouse

Route

oral

oral

Chronic Toxicity**U.S. Occupational Standard:****Carcinogenicity:****Mutagenicity:****Teratogenicity:****TLV:****Other Chronic Effects:**

Bromobenzene

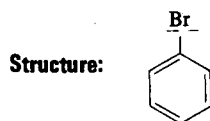
CHEMICAL NAME

CAS Number: 108-86-1

Chemical Name: Bromobenzene

Synonyms: Phenylbromide

Molecular
Formula: C_6H_5Br



Chemical Properties

Molecular Weight: 157.02

Physical State: Liquid

Vapor Pressure: 48 mm at 25°C

Vapor Density: 5.41

Boiling Point: 156.2°C at 760 mm

Melting Point: -30.5°C

Density: 1.4950 at 20°C/4°C

Solubility: Insoluble (H_2O)

Octanol/Water Parti-
tion Coefficient: 2.99

Environmental Persistence

BOD:

Atmospheric Reactivity: Reacts with oxidizing materials

Safety Hazard:

Fire—moderate

Bromobenzene

PRODUCTION DATA

**Annual U.S.
Production:**

Consumption:

**Fraction of
Dispersion:**

**Fraction of Pro-
duction Lost:**

Release Rate (million lbs/yr):

TOXICITY DATA

Acute Toxicity

Dosage

Animal

Route

Non-lethal Acute Effects:

At high concentrations, may cause paralysis and skin irritation

Chronic Toxicity

U.S. Occupational Standard:

Carcinogenicity: Not tested

Mutagenicity:

Teratogenicity:

TLV:

Other Chronic Effects:

Bromonaphthalene

CHEMICAL NAME

CAS Number: 580-13-2

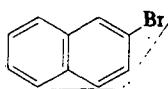
Chemical Name: 2-Bromonaphthalene

Synonyms:

Molecular

Formula: C₁₀H₇Br

Structure:



Chemical Properties

Molecular Weight: 207.08

Physical State: Solid

Vapor Pressure: 1 mm at 84.2°C

Vapor Density:

Boiling Point: 281 to 282°C at 760 mm

Melting Point: 59°C

Density: 1.605 at 0°C

Solubility: Insoluble (H₂O)

Octanol/Water Partition Coefficient:

Environmental Persistence

BOD:

Atmospheric Reactivity:

Safety Hazard:

Bromonaphthalene

PRODUCTION DATA

**Annual U.S.
Production:**

Consumption:

**Fraction of
Dispersion:**

**Fraction of Pro-
duction Lost:**

Release Rate (million lbs/yr):

TOXICITY DATA

Acute Toxicity

Dosage

Animal

Route

Chronic Toxicity

U.S. Occupational Standard:

Carcinogenicity:

Mutagenicity:

Teratogenicity:

TLV:

Other Chronic Effects:

1, 3-Butadiene

CHEMICAL NAME

CAS Number: 106-99-0

Chemical Name: 1,3-Butadiene

Synonyms:	Erythrene	Butadiene	<u>Pyrrolylene</u>
	Biethylene	α - β -Butadiene	Vinylethylene
	Bivinyll	Divinyll	

Molecular

Formula: C₄H₆

Structure: CH₂ = CH - CH = CH₂

Chemical Properties

Molecular Weight: 54.09

Physical State: Gas

Vapor Pressure: 1690 mm at 25°C

Vapor Density: 1.87

Boiling Point: -4.5°C

Melting Point: -109.91°C

Density: 0.621 at 20°C/4°C

Solubility: Insoluble (H₂O)

Octanol/Water Partition Coefficient:

Environmental Persistence

BOD:

Atmospheric Reactivity: Reacts with oxidizing materials

Safety Hazard: Fire—dangerous
Explosion—moderate—forms explosive peroxides with air
Disaster hazard—moderate—toxic fumes

1, 3-Butadiene

PRODUCTION DATA

Annual U.S.**Production:** 3682.1×10^6 lbs. (1974)**Consumption:** 4208×10^6 lbs.**Fraction of****Dispersion:** 0.01**Fraction of Pro-****duction Lost:** 0.015**Release Rate (million lbs/yr):** 56.9 (1972)

TOXICITY DATA

Acute ToxicityLD_{Lo}**Dosage**

250,000 ppm (25 minutes)

Animal

human

Route

inhalation

Non-lethal Acute Effects:

Irritant to eyes and mucous membranes

Unconsciousness due to inhalation**Chronic Toxicity****U.S. Occupational Standard:** (air) TWA: 1000 ppm**Carcinogenicity:****Mutagenicity:****Teratogenicity:**

TLV: 1000 ppm (skin)

Other Chronic Effects:Slight irritation of eyes, upper respiratory tract: 8000 ppm (8 hours)

Butane

CHEMICAL NAME

CAS Number: 106-97-8

Chemical Name: Butane

Synonyms: Methylethylmethane
Butylhydride
Diethyl

n-Butane

Molecular
Formula: C₄H₁₀

Structure: CH₃CH₂CH₂CH₃

Chemical Properties

Molecular Weight: 58.13

Physical State: Gas

Vapor Pressure: ca. 2

Vapor Density: 2.046

Boiling Point: -0.5°C

Melting Point: -138.35°C

Density: 0.6012 at 0°C/4°

Solubility: 85.98 gm/l of H₂O

Octanol/Water Parti-
tion Coefficient:

Environmental Persistence

BOD:

Atmospheric Reactivity: Reacts with oxidizing materials

Safety Hazard:

Fire—dangerous

Explosion—moderate

Disaster hazard—moderate—toxic fumes

Butane

PRODUCTION DATA

Annual U.S. Production:	2331.1 x 10 ⁶ lbs.	Consumption:	2331.1 x 10 ⁶ lbs.
Fraction of Dispersion:	0	Fraction of Production Lost:	0.01
Release Rate (million lbs/yr):	23.3		

TOXICITY DATA

Acute Toxicity	Dosage	Animal	Route
LD ₅₀	658 mg/m ³	rat	inhalation

Non-lethal Acute Effects:

Drowsiness
Narcotic
Simple asphyxiant

Chronic Toxicity

U.S. Occupational Standard: (air) TWA 500 ppm

Carcinogenicity:

Mutagenicity:

Teratogenicity:

TLV:

Other Chronic Effects:

1-Butene

CHEMICAL NAME

CAS Number: 106-98-9

Chemical Name: 1-Butene

Synonyms: 1-Butylene
α-Butylene
Ethylethylene

**Molecular
Formula:** C₄H₈

Structure: CH₃ - CH₂ - CH = CH₂

Chemical Properties

Molecular Weight: 56.12

Physical State: Gas

Vapor Pressure: 3480 at 21°C

Vapor Density: 1.93

Boiling Point: -6.3°C at 760 mm

Melting Point: -185.35°C

Density: 0.5951 at 20°C/4°C

Solubility: Insoluble (H₂O)

**Octanol/Water Parti-
tion Coefficient:**

Environmental Persistence

BOD:

Atmospheric Reactivity: Reacts with oxidizing materials

Safety Hazard:

Fire—dangerous

Explosion—moderate

1-Butene

PRODUCTION DATA

Annual U.S.**Production:** 2478 x 10⁶ lbs. (total butenes)**Consumption:** 3960 x 10⁶ lbs.**Fraction of
Dispersion:****Fraction of Pro-
duction Lost:****Release Rate (million lbs/yr):**

TOXICITY DATA

Acute Toxicity**Dosage****Animal****Route****Non-lethal Acute Effects:**

Asphyxiant

Chronic Toxicity**U.S. Occupational Standard:****Carcinogenicity:****Mutagenicity:****Teratogenicity:****TLV:****Other Chronic Effects:**

2-Butene (cis and trans)

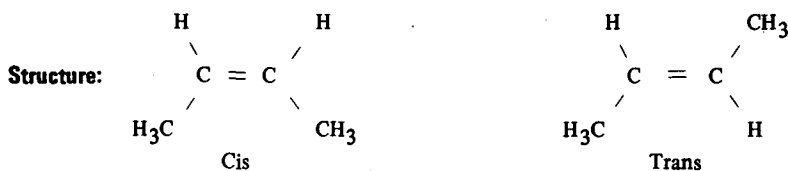
CHEMICAL NAME

CAS Number: 107-01-7

Chemical Name: 2-Butene

Synonyms: 2-Butene (cis and trans)
Dimethylethylene
Pseudobutylene

**Molecular
Formula:** C₄H₈



Chemical Properties

Molecular Weight: 56.12

Physical State: Gas

Vapor Pressure: 1410 mm at 21°C (cis)
1592 mm at 21°C (trans)

Vapor Density: 1.9 (cis and trans)

Boiling Point: 3.7°C at 760 mm (cis)
0.88°C at 760 mm (trans)

Melting Point: -138.91°C (cis)
-105.55°C (trans)

Density: 0.6213 at 20°C/4°C (cis)
0.6042 at 20°C/4°C (trans)

Solubility: Insoluble (H₂O)

**Octanol/Water Parti-
tion Coefficient:**

Environmental Persistence

BOD:

Atmospheric Reactivity: Reacts with oxidizing materials

Safety Hazard:

Fire—dangerous

2-Butene (cis and trans)

PRODUCTION DATA

Annual U.S. Production: $2,478 \times 10^6$ lbs. (total butenes) **Consumption:** $3,600 \times 10^6$ lbs.
Fraction of Dispersion: **Fraction of Production Lost:** 0.015
Release Rate (million lbs/yr):

TOXICITY DATA

Acute Toxicity	Dosage	Animal	Route
-----------------------	---------------	---------------	--------------

Non-lethal Acute Effects:
Simple asphyxiant

Chronic Toxicity

U.S. Occupational Standard:

Carcinogenicity:

Mutagenicity:

Teratogenicity:

TLV:

Other Chronic Effects:

2-Butoxyethanol

CHEMICAL NAME

CAS Number: 111-76-2

Chemical Name: 2-Butoxyethanol

Synonyms:	N-butoxyethanol	Butylglycol	3-Oxa-1-heptanol
	2-Butoxy-1-ethanol	Butyl oxitol	Butyl cellosolve
	Ethylene glycol, monobutyl ether	o-Butyl ethylene glycol	Glycol, n-butyl ether
	Monobutyl glycol ether		

Molecular

Formula: C₆H₁₄O₂

Structure: OH - CH₂ - CH₂ - O CH₂ - CH₂ - CH₂ - CH₃

Chemical Properties

Molecular Weight: 118.18

Physical State: Liquid

Vapor Pressure: 0.88 mm at 25°C

Vapor Density: 4.07

Boiling Point: 171.2°C

Melting Point:

Density:

Solubility: 45 gm/l of H₂O

Octanol/Water Partition Coefficient:

Environmental Persistence

BOD:

Atmospheric Reactivity:

Safety Hazard: Fire—moderate

2-Butoxyethanol

PRODUCTION DATA

Annual U.S.**Production:** 133.3 x 10⁶ lbs.**Consumption:** 124.3 x 10⁶ lbs.**Fraction of****Dispersion:** 0.97**Fraction of Pro-****duction Lost:** 0.015**Release Rate (million lbs/yr):** 122.6

TOXICITY DATA

Acute Toxicity**LD₅₀****Dosage****Animal****Route**

1480 mg/kg

rat

oral

550 mg/kg

rat

intraperitoneal

340 mg/kg

rat

intravenous

1230 mg/kg

mouse

oral

536 mg/kg

mouse

intraperitoneal

1130 mg/kg

mouse

intravenous

320 mg/kg

rabbit

oral

560 mg/kg

rabbit

skin

280 mg/kg

rabbit

intravenous

1200 mg/kg

guinea pig

oral

230 mg/kg

guinea pig

skin

LC₅₀

700 ppm (7 hours)

mouse

inhalation

LC_{Lo}

500 ppm (4 hours)

rat

inhalation

Irritant Effects:**TD_{Lo}**

195 ppm (8 hours)

human

inhalation

Chronic Toxicity**U.S. Occupational Standard:****Carcinogenicity:****Mutagenicity:****Teratogenicity:****TLV:****Other Chronic Effects:**

n-Butyl acetate

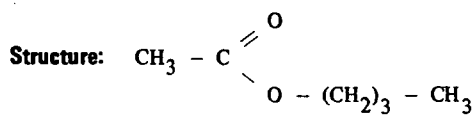
CHEMICAL NAME

CAS Number: 123-86-4

Chemical Name: Acetic acid, butyl ester

Synonyms: Butyl acetate
Butyl ethanoate
Butyl ester of acetic acid

**Molecular
Formula:** $C_6H_{12}O_2$



Chemical Properties

Molecular Weight: 116.16

Physical State: Liquid

Vapor Pressure: 15 mm at 25°C

Vapor Density:

Boiling Point: 125°C

Melting Point: -77.9

Density: 0.882 at 20°/20°C

Solubility: 6.29 gm/l of H₂O

**Octanol/Water Parti-
tion Coefficient:**

Environmental Persistence

BOD: 1.83 for 20 days at 20°C

Atmospheric Reactivity: Reacts with oxidizing materials

Safety Hazard: Fire—moderate
Explosion—moderate

n-Butyl acetate

PRODUCTION DATA

Annual U.S.**Production:** 77.2×10^6 lbs. (1974)**Consumption:** 77.2×10^6 lbs.**Fraction of
Dispersion:** 1.0**Fraction of Pro-
duction Lost:** 0.015**Release Rate (million lbs/yr):** 97.1 (1972)

TOXICITY DATA

Acute ToxicityLD₅₀LD_{Lo}**Dosage**

1230 mg/kg

1500 mg/kg

3200 mg/kg

4700 mg/kg

Animal

mouse

rat

rabbit

guinea pig

Route

intraperitoneal

intraperitoneal

oral

oral

Non-lethal Acute Effects:Irritant effects—TC_{Lo} 200 ppmEyes and respiratory tract

human

inhalation

Chronic Toxicity**U.S. Occupational Standard:** (air) TWA: 150 ppm**Carcinogenicity:** Not tested**Mutagenicity:****Teratogenicity:****TLV:****Other Chronic Effects:** Narcosis

sec-Butyl acetate

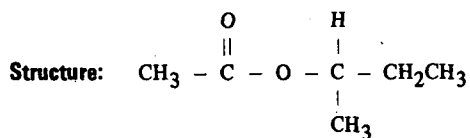
CHEMICAL NAME

CAS Number: 105-46-4

Chemical Name: Acetic acid, secondary butyl ester

Synonyms: sec-Butyl acetate
2-Butonal acetate

**Molecular
Formula:** C₆H₁₂O₂



Chemical Properties

Molecular Weight: 116.16

Physical State: Liquid

Vapor Pressure:

Vapor Density: 4.0

Boiling Point: 112.2°C at 760 mm

Melting Point:

Density: 0.8758 at 16°C/4°C

Solubility: Insoluble (H₂O)

**Octanol/Water Parti-
tion Coefficient:**

Environmental Persistence

BOD:

Atmospheric Reactivity: Reacts with oxidizing materials

Safety Hazard:

Fire—dangerous

Explosion—moderate

sec-Butyl acetate

PRODUCTION DATA

Annual U.S.

Production:

Consumption:

**Fraction of
Dispersion:**

**Fraction of Pro-
duction Lost:**

Release Rate (million lbs/yr):

TOXICITY DATA

Acute Toxicity

Dosage

Animal

Route

Chronic Toxicity

U.S. Occupational Standard: (air) TWA 200 ppm

Carcinogenicity:

Mutagenicity:

Teratogenicity:

TLV:

Other Chronic Effects:

Minor allergen and irritant

Moderately toxic if inhaled or ingested

tert-Butyl acetate

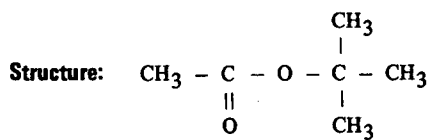
CHEMICAL NAME

CAS Number: 540-88-5

Chemical Name: Acetic acid, 1,1-dimethylethyl ester

Synonyms: tert-Butyl acetate
Acetic acid, tert-butyl ester

**Molecular
Formula:** C₆H₁₂O₂



Chemical Properties

Molecular Weight: 116.16

Physical State: Liquid

Vapor Pressure:

Vapor Density:

Boiling Point: 97 to 98°C at 760 mm

Melting Point:

Density: 0.8665 at 20°C/4°C

Solubility: Slightly (H₂O)

**Octanol/Water Parti-
tion Coefficient:**

Environmental Persistence

BOD:

Atmospheric Reactivity:

Safety Hazard:

tert-Butyl acetate

PRODUCTION DATA

**Annual U.S.
Production:**

Consumption:

**Fraction of
Dispersion:**

**Fraction of Pro-
duction Lost:**

Release Rate (million lbs/yr):

TOXICITY DATA

Acute Toxicity

Dosage

Animal

Route

Chronic Toxicity

U.S. Occupational Standard: (air) TWA 200 ppm

Carcinogenicity:

Mutagenicity:

Teratogenicity:

TLV:

Other Chronic Effects:

Minor allergen and irritant

Moderately toxic if inhaled or ingested

n-Butyl acrylate

CHEMICAL NAME

CAS Number: 141-32-2

Chemical Name: Acrylic acid, butyl ester

Synonyms: n-Butyl acrylate
2-Propenoic acid, butyl ester

**Molecular
Formula:** C₇H₁₂O₂

Structure:
$$\text{CH}_2 = \text{CH} - \overset{\text{O}}{\underset{||}{\text{C}}} - \text{O}(\text{CH}_2)_3\text{CH}_3$$

Chemical Properties

Molecular Weight: 128.17

Physical State: Liquid

Vapor Pressure: 5.63 mm at 25°C

Vapor Density: 4.42

Boiling Point: 146 to 148°C at 760 mm

Melting Point: -64.6°C

Density: 0.8898 at 20°C/4°C

Solubility: Insoluble (H₂O)

**Octanol/Water Parti-
tion Coefficient:**

Environmental Persistence

BOD:

Atmospheric Reactivity: Reacts with oxidizing materials

Safety Hazard:

Fire—moderate

n-Butyl acrylate

PRODUCTION DATA

**Annual U.S.
Production:**

Consumption:

**Fraction of
Dispersion:**

**Fraction of Pro-
duction Lost:**

Release Rate (million lbs/yr):

TOXICITY DATA

Acute Toxicity

LD₅₀

Dosage

3730 mg/kg

2000 mg/kg

Animal

rat

rabbit

Route

oral

skin

LC_{Lo}

1000 ppm (4 hours)

rat

inhalation

Chronic Toxicity

U.S. Occupational Standard:

Carcinogenicity: Not tested

Mutagenicity:

Teratogenicity:

TLV:

Other Chronic Effects:

n-Butyl alcohol

CHEMICAL NAME

CAS Number: 71-36-3

Chemical Name: Butyl alcohol

Synonyms: Butanol
n-Butanol
l-Butanol
Butyl alcohol

Butyl hydroxide
1-Hydroxybutane
Methylolpropane

Propylcarbinol
n-Propylcarbinol

Molecular
Formula: C₃H₁₀O

Structure: CH₃ - CH₂ - CH₂ - CH₂ OH

Chemical Properties

Molecular Weight: 74.12

Physical State: Liquid

Vapor Pressure: 7.69 mm at 25°C

Vapor Density: 2.55

Boiling Point: 117 to 118°C at 760 mm

Melting Point: -89.0°C

Density: 0.80978 at 20°C/4°C

Solubility: Soluble (84.92 gm/l H₂O)

Octanol/Water Parti-
tion Coefficient: 1.00

Environmental Persistence

BOD: 2.4 for 20 days at 20°C

Atmospheric Reactivity: Reacts with oxidizing materials

Safety Hazard: Fire—dangerous
Explosion—moderate
Disaster hazard—moderate—toxic fumes

PRODUCTION DATA**Annual U.S.****Production:** 557.6×10^6 lbs.**Consumption:** 350×10^6 lbs. (1975)**Fraction of
Dispersion:** 0.23**Fraction of Pro-
duction Lost:** 0.015**Release Rate (million lbs/yr):** 144.6**TOXICITY DATA****Acute Toxicity**LD₅₀
LD_{Lo}**Dosage**
790 mg/kg
970 mg/kg
4250 mg/kg**Animal**
rat
rat
rabbit**Route**
oral
intraperitoneal
oral**Non-lethal Acute Effects:**Pulmonary effects—TC_{Lo} 25 ppm

human

inhalation

Irritant—eyes, corneal inflammation, headache, dizziness

Slight irritant—nose and throat

Dermatis—fingernails and finger

Keratitis—reported**Chronic Toxicity****U.S. Occupational Standard:** (air) TWA: 100 ppm**Carcinogenicity:****Mutagenicity:****Teratogenicity:**

TLV: 100 ppm

Other Chronic Effects:Eye irritation: 50 ppm
Mild irritation: 25 ppm
Headaches: 50 ppm

sec-Butyl alcohol

CHEMICAL NAME

CAS Number: 78-92-2

Chemical Name: sec-Butyl alcohol

Synonyms:	Butan-2-ol	Butylene hydrate	s-Butanol
	Ethyl methyl carbinol	2-Butanol	2-Hydroxybutane
	2-Butyl alcohol	Methyl ethyl carbinol	

**Molecular
Formula:** C₄H₁₀O

Structure:

$$\begin{array}{c} \text{CH}_3 - \text{CH}_2 - \text{CH} - \text{CH}_3 \\ | \\ \text{OH} \end{array}$$

Chemical Properties

Molecular Weight: 74.12

Physical State: Liquid

Vapor Pressure: 17.5 mm at 25°C

Vapor Density: 2.55

Boiling Point: 99.5

Melting Point: -114.7°C

Density: 0.808 at 20°C/4°C

Solubility: 101 gm/l of H₂O

**Octanol/Water Parti-
tion Coefficient:**

Environmental Persistence

BOD: 2.0 for 50 days at 20°C

Atmospheric Reactivity: Reacts with oxidizing materials

Safety Hazard: Fire-dangerous

sec-Butyl alcohol

PRODUCTION DATA

Annual U.S.

Production: 417×10^6 lbs.

Consumption: 417×10^6 lbs.

Fraction of

Dispersion: 0.05

Fraction of Pro-

duction Lost: 0.015

Release Rate (million lbs/yr): 27.1

TOXICITY DATA

Acute Toxicity

LD₅₀

LD_{Lo}

Dosage

771 mg/kg

6000 mg/kg

Animal

mouse

rabbit

Route

intraperitoneal

oral

Chronic Toxicity

U.S. Occupational Standard: (air) TWA: 150 ppm

Carcinogenicity: Not tested

Mutagenicity:

Teratogenicity:

TLV: 150 ppm

Other Chronic Effects:

tert-Butyl alcohol

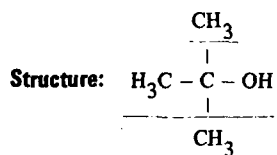
CHEMICAL NAME

CAS Number: 75-65-0

Chemical Name: tert-Butyl alcohol

Synonyms: 2-Methyl-2 propanol
1,1-Dimethyl ethanol

**Molecular
Formula:** C₄H₁₀O



Chemical Properties

Molecular Weight: 74.12

Vapor Pressure: 41.54 at 25°C

Boiling Point: 82.2

Density: 0.7887 at 20°C/4°C

**Octanol/Water Parti-
tion Coefficient:** 0.61

Physical State: Liquid

Vapor Density: 2.55

Melting Point: 25.5°C

Solubility: Infinite (H₂O)

Environmental Persistence

BOD:

Atmospheric Reactivity:

Safety Hazard: Fire—dangerous
Explosion—moderate

tert-Butyl alcohol

PRODUCTION DATA

Annual U.S.**Production:** 1000 x 10⁶ lbs.**Consumption:** 1000 x 10⁶ lbs.**Fraction of****Dispersion:** 0.7**Fraction of Pro-****duction Lost:** 0.015**Release Rate (million lbs/yr):** 715

TOXICITY DATA

Acute ToxicityLD₅₀**Dosage**

3500 mg/kg

933 mg/kg

LD_{Lo}

4500 mg/kg

Animal

rat

mouse

rabbit

Route

oral

intraperitoneal

oral

Chronic Toxicity**U.S. Occupational Standard:** (air) TWA: 100 ppm**Carcinogenicity:****Mutagenicity:****Teratogenicity:****TLV:****Other Chronic Effects:**Human skin contact: slight erythema and hyperemia

n-Butylaldehyde

CHEMICAL NAME

CAS Number: 123-72-8

Chemical Name: Butyraldehyde

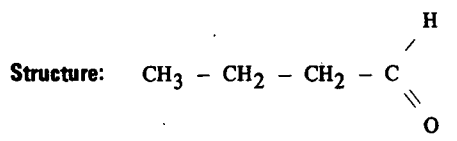
Synonyms: n-Butyl aldehyde

Butanal

Butyric aldehyde

Molecular

Formula: C₄H₈O



Chemical Properties

Molecular Weight: 72.12

Physical State: Liquid

Vapor Pressure: <10 mm at 25°C

Vapor Density: 2.5

Boiling Point: 75.7°C at 760 mm

Melting Point: -99°C

Density: 0.8170 at 20°C/4°C

Solubility: Soluble (H₂O)

Octanol/Water Partition Coefficient: 1.18

Environmental Persistence

BOD:

Atmospheric Reactivity: Reacts with oxidizing materials

Safety Hazard:

Fire—dangerous

n-Butylaldehyde

PRODUCTION DATA

Annual U.S.

Production:

Consumption:

**Fraction of
Dispersion:**

**Fraction of Pro-
duction Lost:**

Release Rate (million lbs/yr):

TOXICITY DATA

Acute Toxicity

LD₅₀

Dosage

2490 mg/kg

2700 mg/kg

Animal

rat

mouse

Route

oral

subcutaneous

Non-lethal Acute Effects:

Irritant

TC_{Lo}

580 mg/kg

human

inhalation

Chronic Toxicity

U.S. Occupational Standard:

Carcinogenicity: Not tested

Mutagenicity:

Teratogenicity:

TLV:

Other Chronic Effects:

sec-Butylamine

CHEMICAL NAME

CAS Number: 513-49-5

Chemical Name: (S)2-Butanamine

Synonyms: sec-Butylamine
2-Aminobutane

**Molecular
Formula:** C₄H₁₁N

Structure:
$$\begin{array}{ccccccc} & & & & \text{NH}_2 & & \\ & & & & | & & \\ \text{CH}_3 & - & \text{CH}_2 & - & \text{CH} & - & \text{CH}_3 \end{array}$$

Chemical Properties

Molecular Weight: 73.14

Physical State: Liquid

Vapor Pressure: 169.8 mm at 25°C

Vapor Density:

Boiling Point: 63.0°C at 760 mm

Melting Point: -104.5°C

Density: 0.724 at 20°C/4°C

Solubility: Soluble (H₂O)

**Octanol/Water Parti-
tion Coefficient:**

Environmental Persistence

BOD:

Atmospheric Reactivity: Reacts with oxidizing materials

Safety Hazard:
Fire—dangerous

sec-Butylamine

PRODUCTION DATA

**Annual U.S.
Production:**

Consumption:

**Fraction of
Dispersion:**

**Fraction of Pro-
duction Lost:**

Release Rate (million lbs/yr):

TOXICITY DATA

Acute Toxicity
LD₅₀

Dosage
380 mg/kg

Animal
rat

Route
oral

Chronic Toxicity

U.S. Occupational Standard:

Carcinogenicity:

Mutagenicity:

Teratogenicity:

TLV:

Other Chronic Effects:

tert-Butylamine

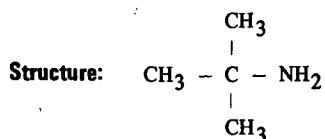
CHEMICAL NAME

CAS Number: 75-64-9

Chemical Name: t-Butylamine

Synonyms: 2-Aminoisobutane
Trimethylaminomethane

**Molecular
Formula:** C₄H₁₁N



Chemical Properties

Molecular Weight: 73.14

Physical State: Liquid

Vapor Pressure: 169.8 mm at 25°C

Vapor Density: 2.5

Boiling Point: 44 to 46°C

Melting Point:

Density: 0.700 at 15°C

Solubility: Miscible (H₂O)

**Octanol/Water Parti-
tion Coefficient:**

Environmental Persistence

BOD: 27% of theoretical

Atmospheric Reactivity: Reacts with oxidizing materials

Safety Hazard:

Fire—moderate

tert-Butylamine

PRODUCTION DATA

**Annual U.S.
Production:**

Consumption:

**Fraction of
Dispersion:**

**Fraction of Pro-
duction Lost:**

Release Rate (million lbs/yr):

TOXICITY DATA

Acute Toxicity
LD₅₀

Dosage
180 mg/kg
900 mg/kg

Animal
rat
mouse

Route
oral
oral

Chronic Toxicity

U.S. Occupational Standard:

Carcinogenicity:

Mutagenicity:

Teratogenicity:

TLV:

Other Chronic Effects:

Butylbenzoic acid

CHEMICAL NAME

CAS Number: 1320-16-7

Chemical Name: (1,1-Dimethylethyl)benzoic acid

Synonyms: p-t-BBA

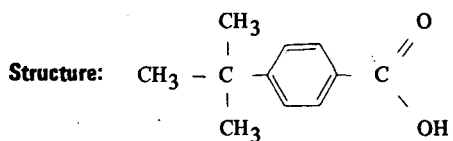
TBBA

p-tert-Butylbenzoic acid

4-tert-Butylbenzoic acid

Molecular

Formula: $C_{11}H_{14}O_2$



Chemical Properties

Molecular Weight: 178.23

Vapor Pressure:

Boiling Point:

Density: 1.142 at 20°C/4°C

Octanol/Water Partition Coefficient:

Physical State: Solid

Vapor Density:

Melting Point: 164.5 to 165.5°C

Solubility: Insoluble (H₂O)

Environmental Persistence

BOD:

Atmospheric Reactivity:

Safety Hazard:

Fire—slight

Butylbenzoic acid

PRODUCTION DATA

Annual U.S.
Production:

Consumption:

Fraction of
Dispersion:

Fraction of Pro-
duction Lost:

Release Rate (million lbs/yr):

TOXICITY DATA

Acute Toxicity
LD₅₀

Dosage
735 mg/kg

Animal
rat

Route
oral

Non-lethal Acute Effects:
Local irritant

Chronic Toxicity

U.S. Occupational Standard:

Carcinogenicity:

Mutagenicity:

Teratogenicity:

TLV:

Other Chronic Effects:

Butylene

CHEMICAL NAME

CAS Number: 115-11-7

Chemical Name: 1-Methylpropene

Synonyms: Isobutylene
Isobutene

Molecular
Formula: C_4H_8

Structure:
$$\begin{array}{c} \text{CH}_3 \\ | \\ \text{H}_3\text{C} - \text{C} = \text{CH}_2 \end{array}$$

Chemical Properties

Molecular Weight: 56.11

Physical State: Gas

Vapor Pressure: 3290 mm at 40.5°C

Vapor Density: 1.94

Boiling Point: -6.9°C

Melting Point: -140.35°C

Density: 0.5942 at 20°C/4°C (liquid)

Solubility: Insoluble (H_2O)

Octanol/Water Partition Coefficient:

Environmental Persistence

BOD:

Atmospheric Reactivity: Reacts with oxidizing materials

Safety Hazard:

Fire—dangerous

Explosion—dangerous

Butylene

PRODUCTION DATA

**Annual U.S.
Production:**

Consumption:

**Fraction of
Dispersion:**

**Fraction of Pro-
duction Lost:**

Release Rate (million lbs/yr):

TOXICITY DATA

Acute Toxicity

Dosage

Animal

Route

Non-lethal Acute Effects:

Simple asphyxiant with narcotic action

Chronic Toxicity

U.S. Occupational Standard:

Carcinogenicity:

Mutagenicity:

Teratogenicity:

TLV:

Other Chronic Effects:

1, 3-Butylene glycol

CHEMICAL NAME

CAS Number: 107-88-0

Chemical Name: 1,3-Butanediol

Synonyms: 1,3-Butylene glycol
 β -Butyleneglycol

**Molecular
Formula:** $C_4H_{10}O_2$

Structure:
$$\begin{array}{ccccccc} & & \text{OH} & & & & \\ & & | & & & & \\ \text{CH}_3 & - & \text{CH} & - & \text{CH}_2 & - & \text{CH}_2\text{OH} \end{array}$$

Chemical Properties

Molecular Weight: 90.12

Physical State: Liquid

Vapor Pressure: 0.06 mm at 20°C

Vapor Density: 3.2

Boiling Point: 207.5°C

Melting Point:

Density: 1.006 at 20°C/20°C

Solubility: Infinite (H_2O)

**Octanol/Water Parti-
tion Coefficient:** -1.10

Environmental Persistence

BOD:

Atmospheric Reactivity: Reacts with oxidizing materials

Safety Hazard:
Fire—slight

1, 3-Butylene glycol

PRODUCTION DATA

**Annual U.S.
Production:**

Consumption:

**Fraction of
Dispersion:**

**Fraction of Pro-
duction Lost:**

Release Rate (million lbs/yr):

TOXICITY DATA

Acute Toxicity

Dosage

Animal

Route

Chronic Toxicity

U.S. Occupational Standard:

Carcinogenicity: Not tested

Mutagenicity:

Teratogenicity:

TLV:

Other Chronic Effects:

n-Butyl glycidyl ether

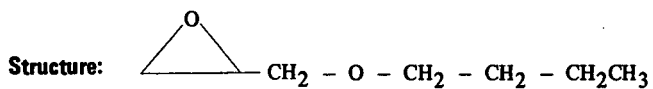
CHEMICAL NAME

CAS Number: 2426-08-6

Chemical Name: (Butoxymethyl)oxirane

Synonyms: n-Butyl glycidyl ether
BGE
1-Butoxy-2,3-epoxypropane

**Molecular
Formula:** C₇H₁₄O₂



Chemical Properties

Molecular Weight: 130.21

Physical State: Liquid

Vapor Pressure:

Vapor Density:

Boiling Point: 164°C

Melting Point:

Density: 0.91

Solubility: Slightly (H₂O)

**Octanol/Water Parti-
tion Coefficient:**

Environmental Persistence

BOD:

Atmospheric Reactivity:

Safety Hazard:
Explosion—dangerous

n-Butyl glycidyl ether

PRODUCTION DATA

Annual U.S.
Production:

Consumption:

Fraction of
Dispersion:

Fraction of Pro-
duction Lost:

Release Rate (million lbs/yr):

TOXICITY DATA

Acute Toxicity
LD₅₀

Dosage
2050 mg/kg
1190 mg/kg
1520 mg/kg
700 mg/kg
2520 mg/kg
670 ppm

Animal
rat
rat
mouse
mouse
rabbit
rat

Route
oral
intraperitoneal
intraperitoneal
intraperitoneal
skin
inhalation

Non-lethal Acute Effects:
Irritant to skin

Chronic Toxicity

U.S. Occupational Standard: (air) TWA 50 ppm

Carcinogenicity:

Mutagenicity:

Teratogenicity:

TLV: 50 ppm or 27 mg/m³

Other Chronic Effects:

Butylmercaptan

CHEMICAL NAME

CAS Number: 109-71-5

Chemical Name: 1-Butanethiol

Synonyms: n-Butylmercaptan
n-Butyl thioalcohol

**Molecular
Formula:** C₄H₁₀S

Structure: CH₃CH₂CH₂CH₂SH

Chemical Properties

Molecular Weight: 90.19

Physical State: Liquid

Vapor Pressure: <10 mm at 25°C

Vapor Density: 3.1

Boiling Point: 98.46°C

Melting Point: -115.67°C

Density: 0.8299 at 20°C/4°C

Solubility: Slightly (H₂O)

**Octanol/Water Parti-
tion Coefficient:** 2.28

Environmental Persistence

BOD:

Atmospheric Reactivity: Reacts with oxidizing materials

Safety Hazard:
Fire—dangerous

Butylmercaptan

PRODUCTION DATA

Annual U.S.
Production:

Consumption:

Fraction of
Dispersion:

Fraction of Pro-
duction Lost:

Release Rate (million lbs/yr):

TOXICITY DATA

Acute Toxicity	Dosage	Animal	Route
LD ₅₀	1500 mg/kg	rat	oral
	399 mg/kg	rat	intratracheal
LC ₅₀	4020 ppm (4 hours)	rat	inhalation
	2500 ppm (4 hours)	mouse	inhalation
LC _{Lo}	700 ppm (30 minutes)	dog	inhalation
Non-lethal Acute Effects:			
TC _{Lo}	10 mg/m ³ (3 hours)	human	inhalation
<u>Paralysis if concentrated vapor is inhaled</u>			

Chronic Toxicity

U.S. Occupational Standard: (air) TWA 10 ppm

Carcinogenicity:

Mutagenicity:

Teratogenicity:

TLV:

Other Chronic Effects:

tert-Butylphenol

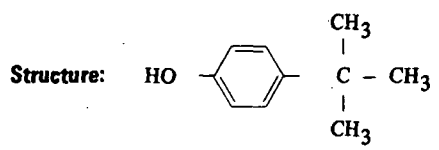
CHEMICAL NAME

CAS Number: 98-54-4

Chemical Name: p-tert-Butylphenol

Synonyms: Hydroxy-4-tert-butyl benzene

**Molecular
Formula:** C₁₀H₁₄O



Chemical Properties

Molecular Weight: 150.22

Vapor Pressure: 1 mm at 70°C

Boiling Point: 239.5°C at 760 mm

Density: 0.908 at 80°C/4°C

**Octanol/Water Parti-
tion Coefficient:** 3.31

Physical State: Solid

Vapor Density: 5.1

Melting Point: 101°C

Solubility: Soluble (hot H₂O)

Environmental Persistence

BOD:

Atmospheric Reactivity:

Safety Hazard:

tert-Butylphenol

PRODUCTION DATA

Annual U.S.
Production:

Consumption:

Fraction of
Dispersion:

Fraction of Pro-
duction Lost:

Release Rate (million lbs/yr):

TOXICITY DATA

Acute Toxicity
LD₅₀

Dosage
3250 mg/kg
2520 mg/kg

Animal
rat
rabbit

Route
oral
skin

Non-lethal Acute Effects:
Irritant

Chronic Toxicity

U.S. Occupational Standard:

Carcinogenicity:

Mutagenicity:

Teratogenicity:

TLV:

Other Chronic Effects:

tert-Butyl toluene

CHEMICAL NAME

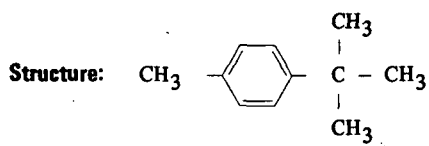
CAS Number: 98-51-1

Chemical Name: p-tert-Butyl toluene

Synonyms: p-Methyl-tert-butylbenzene

Molecular

Formula: $C_{11}H_{16}$



Chemical Properties

Molecular Weight: 148.25

Physical State: Liquid

Vapor Pressure: <10 mm at 25°C

Vapor Density:

Boiling Point: 193°C at 760 mm

Melting Point: -52°C

Density: 0.8612 at 20°C/4°C

Solubility: Insoluble

Octanol/Water Partition Coefficient:

Environmental Persistence

BOD:

Atmospheric Reactivity: Reacts with oxidizing materials

Safety Hazard:

Fire—moderate

Disaster hazard—dangerous—toxic fumes

tert-Butyl toluene

PRODUCTION DATA

**Annual U.S.
Production:**

Consumption:

**Fraction of
Dispersion:**

**Fraction of Pro-
duction Lost:**

Release Rate (million lbs/yr):

TOXICITY DATA

Acute Toxicity	Dosage	Animal	Route
LD₅₀	1500 mg/kg	rat	oral
	900 mg/kg	mouse	oral
	2000 mg/kg	rabbit	oral
LC₅₀	1500 mg/m ³ (4 hours)	rat	inhalation
	248 ppm (2 hours)	mouse	inhalation

Non-lethal Acute Effects:

Irritant:

TC_{Lo}	10 ppm (3 minutes)	human	inhalation
------------------------	--------------------	-------	------------

Central nervous system effects:

TC_{Lo}	20 ppm (5 minutes)	human	inhalation
------------------------	--------------------	-------	------------

Chronic Toxicity

U.S. Occupational Standard: (air) TWA 10 ppm

Carcinogenicity:

Mutagenicity:

Teratogenicity:

TLV: 10 ppm

Other Chronic Effects:

Liver and kidneys

n-Butyric acid

CHEMICAL NAME

CAS Number: 107-92-6

Chemical Name: Butyric acid

Synonyms: Butanoic acid
Ethyl acetic acid
Propyl formic acid

**Molecular
Formula:** $C_4H_8O_2$

Structure: $CH_3CH_2CH_2 - \underset{\begin{array}{c} || \\ O \end{array}}{C} - OH$

Chemical Properties

Molecular Weight: 88.12

Physical State: Liquid

Vapor Pressure: 1 mm at 25.5°C

Vapor Density: 3.04

Boiling Point: 163.53°C at 760 mm

Melting Point: -4.26°C

Density: 0.9577 at 20°C/4°C

Solubility: Infinite (H_2O)

**Octanol/Water Parti-
tion Coefficient:** 0.97

Environmental Persistence

BOD:

Atmospheric Reactivity: Reacts with oxidizing materials

Safety Hazard:
Fire-moderate

n-Butyric acid

PRODUCTION DATA

Annual U.S.
Production:

Consumption:

Fraction of
Dispersion:

Fraction of Pro-
duction Lost:

Release Rate (million lbs/yr):

TOXICITY DATA

Acute Toxicity	Dosage	Animal	Route
LD ₅₀	2400 mg/kg	rat	oral
	3180 mg/kg	mouse	intraperitoneal
	3180 mg/kg	mouse	subcutaneous
	800 mg/kg	mouse	intravenous
LD _{Lo}	500 mg/kg	mouse	oral
	3600 mg/kg	rabbit	oral

Non-lethal Acute Effects:

Irritant to eyes, skin and respiratory tract

Chronic Toxicity

U.S. Occupational Standard:

Carcinogenicity:

Mutagenicity:

Teratogenicity:

TLV:

Other Chronic Effects:

n-Butyric anhydride

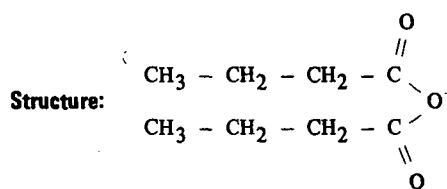
CHEMICAL NAME

CAS Number: 106-31-0

Chemical Name: Butyric anhydride

Synonyms: n-Butyric anhydride
Butanoic anhydride

**Molecular
Formula:** C₈H₁₄O₃



Chemical Properties

Molecular Weight: 158.20

Physical State: Liquid

Vapor Pressure: <10 mm at 25°C

Vapor Density: 5.4

Boiling Point: 199.4 to 201.4° at 760 mm

Melting Point: -75°C

Density: 0.9668 at 20°C/4°C

Solubility: Decomposes (H₂O)

**Octanol/Water Parti-
tion Coefficient:**

Environmental Persistence

BOD:

Atmospheric Reactivity: Reacts with oxidizing materials

Safety Hazard:
Fire—moderate

n-Butyric anhydride

PRODUCTION DATA

**Annual U.S.
Production:**

Consumption:

**Fraction of
Dispersion:**

**Fraction of Pro-
duction Lost:**

Release Rate (million lbs/yr):

TOXICITY DATA

Acute Toxicity

Dosage

Animal

Route

Chronic Toxicity

U.S. Occupational Standard:

Carcinogenicity: Not tested

Mutagenicity:

Teratogenicity:

TLV:

Other Chronic Effects:

Butyronitrile

CHEMICAL NAME

CAS Number: 109-74-0

Chemical Name: Butyronitrile

Synonyms: Butane nitrile
N-propyl cyanide

**Molecular
Formula:** C_4H_7N

Structure: $CH_3 - CH_2CH_2C \equiv N$

Chemical Properties

Molecular Weight: 69.11

Physical State: Liquid

Vapor Pressure: 20 mm at 25.7°C

Vapor Density:

Boiling Point: 118°C at 760 mm

Melting Point: -112°C

Density: 0.7936 at 20°C/4°C

Solubility: Slightly (H_2O)

**Octanol/Water Parti-
tion Coefficient:**

Environmental Persistence

BOD:

Atmospheric Reactivity: Reacts with oxidizing materials

Safety Hazard:

Fire—dangerous

Disaster hazard—dangerous—toxic fumes—chlorides

Butyronitrile

PRODUCTION DATA

Annual U.S.
Production:

Consumption:

Fraction of
Dispersion:

Fraction of Pro-
duction Lost:

Release Rate (million lbs/yr):

TOXICITY DATA

Acute Toxicity	Dosage	Animal	Route
LD ₅₀	500 mg/kg	rabbit	skin
LD _{Lo}	50 mg/kg	rat	oral
	100 mg/kg	mouse	oral
	100 mg/kg	guinea pig	skin
LC _{Lo}	400 ppm	rat	inhalation

Chronic Toxicity

U.S. Occupational Standard:

Carcinogenicity: Not tested

Mutagenicity:

Teratogenicity:

TLV:

Other Chronic Effects:

Buxten

CHEMICAL NAME

CAS Number: 2282-34-0 (a and b)

Chemical Name: 3-(1-Methylbutyl)phenol methylcarbamate

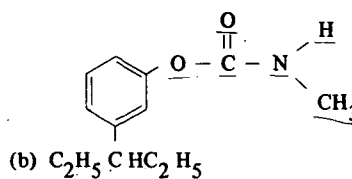
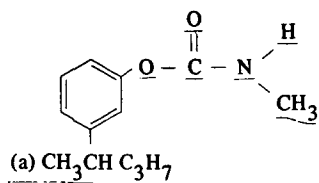
Synonyms: Bux

Mixture of (a) m-(1-Methylbutyl)-phenyl methylcarbamate + (b) m-(1-Ethyl propyl)phenyl methylcarbamate

Molecular

Formula: $C_{13}H_{19}NO_2$

Structure:



Chemical Properties

Molecular Weight: 221.33

Physical State: Solid

Vapor Pressure:

Vapor Density:

Boiling Point:

Melting Point: 26.4°C

Density:

Solubility: Insoluble (H_2O)

Octanol/Water Partition Coefficient:

Environmental Persistence

BOD:

Atmospheric Reactivity:

Safety Hazard:

PRODUCTION DATA**Annual U.S.****Production:** 6 x 10⁶ lbs.**Consumption:****Fraction of****Dispersion:** Estimated as 1.0**Fraction of Pro-****duction Lost:** 0.015**Release Rate (million lbs/yr):** 6**TOXICITY DATA****Acute Toxicity**LD₅₀

3:1 mixture (a:b)

LD₅₀

4:1 mixture (a:b)

LD₅₀**Dosage**

1050 mg/kg

170 mg/kg

1400 mg/kg

400 mg/kg

87 mg/kg

242 mg/kg

1400 mg/kg

400 mg/kg

Animal

rat

rat

dog

rabbit

rat

rat

dog

rabbit

Route

oral

oral

skin

skin

oral

skin

skin

skin

Chronic Toxicity**U.S. Occupational Standard:****Carcinogenicity:****Mutagenicity:****Teratogenicity:****TLV:****Other Chronic Effects:**

Cacodylic acid

CHEMICAL NAME

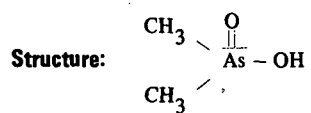
CAS Number: 75-60-5

Chemical Name: Hydroxydimethyl arsine oxide

Synonyms: Dimethyl arsenic acid

Molecular

Formula: $C_2H_7AsO_2$



Chemical Properties

Molecular Weight: 138.01

Physical State: Solid

Vapor Pressure:

Vapor Density:

Boiling Point:

Melting Point: 200°C

Density:

Solubility: Very soluble (H₂O)

Octanol/Water Partition Coefficient:

Environmental Persistence

BOD:

Atmospheric Reactivity:

Safety Hazard: Disaster hazard—hazardous when water solution is in contact with active metals; i.e., Fe, Al, and Zn.

Cacodylic acid

PRODUCTION DATA

Annual U.S.**Production:** 2×10^6 lbs.**Consumption:****Fraction of****Dispersion:** 1.0**Fraction of Pro-****duction Lost:** 0.015**Release Rate (million lbs/yr):** 2

TOXICITY DATA

Acute ToxicityLD₅₀LD_{Lo}**Dosage**

1350 mg/kg

185 mg/kg

500 mg/kg

1000 mg/kg

Animal

rat

mouse

mouse

dog

Route

oral

intraperitoneal

unknown

subcutaneous

Chronic Toxicity**U.S. Occupational Standard:****Carcinogenicity:** Carcinogenic**Mutagenicity:****Teratogenicity:****TLV:****Other Chronic Effects:**

Calcium propionate

CHEMICAL NAME

CAS Number: 4075-81-4

Chemical Name: Calcium propionate

Synonyms: Propionic acid, calcium salt

Molecular

Formula: C₆H₁₀O₄·Ca

Structure:
$$\begin{array}{c} (\text{CH}_3 - \text{CH}_2 - \text{C} - \text{O}^{(-)})_2 \text{Ca}^{(++)} \\ \parallel \\ \text{O} \end{array}$$

Chemical Properties

Molecular Weight: 186

Vapor Pressure:

Boiling Point:

Density:

Octanol/Water Partition Coefficient:

Physical State: Solid

Vapor Density:

Melting Point:

Solubility: Soluble (55.8 gm/l H₂O)

Environmental Persistence

BOD:

Atmospheric Reactivity:

Safety Hazard:

Calcium propionate

PRODUCTION DATA

Annual U.S.**Production:** 21.4×10^6 lbs.**Consumption:** 21.4×10^6 lbs.**Fraction of****Dispersion:** 1.0**Fraction of Pro-****duction Lost:** 0.03**Release Rate (million lbs/yr):** 22.0

TOXICITY DATA

Acute Toxicity**Dosage****Animal****Route****Chronic Toxicity****U.S. Occupational Standard:****Carcinogenicity:****Mutagenicity:****Teratogenicity:****TLV:****Other Chronic Effects:**

Calcium stearate

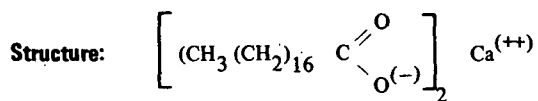
CHEMICAL NAME

CAS Number: 1592-23-0

Chemical Name: Stearic acid, calcium salt

Synonyms: Calcium salt octadecanoic acid
Calcium stearate

**Molecular
Formula:** $(C_{18}H_{35}O_2)_2 \cdot Ca$



Chemical Properties

Molecular Weight: 681.48

Vapor Pressure:

Boiling Point:

Density:

**Octanol/Water Parti-
tion Coefficient:**

Physical State:

Vapor Density:

Melting Point: 156 to 160°C

Solubility: Insoluble (H₂O)

Environmental Persistence

BOD: 80% of theoretical for 5 days at 20°C

Atmospheric Reactivity:

Safety Hazard:

Calcium stearate

PRODUCTION DATA

Annual U.S.**Production:** 37.8×10^6 lbs.**Consumption:** 37.8×10^6 lbs.**Fraction of****Dispersion:** 1.0**Fraction of Pro-****duction Lost:** 0.03**Release Rate (million lbs/yr):** 38.9

TOXICITY DATA

Acute Toxicity**Dosage****Animal****Route****Chronic Toxicity****U.S. Occupational Standard:****Carcinogenicity:****Mutagenicity:****Teratogenicity:****TLV:****Other Chronic Effects:**

Camphor

CHEMICAL NAME

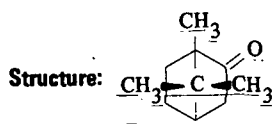
CAS Number: 76-22-2

Chemical Name: Camphor

Synonyms: 2-Camphone

Molecular

Formula: $C_{10}H_{16}O$



Chemical Properties

Molecular Weight: 152.23

Vapor Pressure: 1 mm at 41.5°C

Boiling Point: 204°C (sublimes)

Density: 0.992 at 25°C/4°C

Octanol/Water Partition Coefficient:

Physical State: Solid

Vapor Density: 5.24

Melting Point: 174 to 179°C

Solubility: Insoluble (H_2O)

Environmental Persistence

BOD:

Atmospheric Reactivity:

Safety Hazard: Fire—moderate
Explosion—moderate

Camphor

PRODUCTION DATA

Annual U.S.
Production:

Consumption:

Fraction of
Dispersion:

Fraction of Pro-
duction Lost:

Release Rate (million lbs/yr):

TOXICITY DATA

Acute Toxicity

LD₅₀
LD_{Lo}

Dosage
900 mg/kg
2200 mg/kg

Animal
rat
mouse

Route
intraperitoneal
subcutaneous

Non-lethal Acute Effects:

Irritant
Convulsions

Chronic Toxicity

U.S. Occupational Standard: (air) 2 ppm (TWA)

Carcinogenicity: Neoplastic effect

TD_{Lo}

84 mg/kg (1 week)

mouse

skin

Mutagenicity:

Teratogenicity:

TLV: 5 mg/m³

Other Chronic Effects:

Caprolactam

CHEMICAL NAME

CAS Number: 105-60-2

Chemical Name: Hexahydro-2H-azepin-2-one

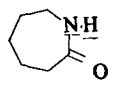
Synonyms: 2-Oxohexamethylenimine
6-Aminohexanoic acid, cyclic lactam

2-Azacycloheptanone

Molecular

Formula: $C_6H_{11}NO$

Structure:



Chemical Properties

Molecular Weight: 113.16

Vapor Pressure: 6 mm at 120°C

Boiling Point: 139°C at 12 mm

Density:

Octanol/Water Partition Coefficient:

Physical State: Solid

Vapor Density:

Melting Point: 69 to 71°C

Solubility: Soluble (H₂O)

Environmental Persistence

BOD:

Atmospheric Reactivity:

Safety Hazard:

Caprolactam

PRODUCTION DATA

Annual U.S. Production: 950 x 10⁶ lbs. (1976) capacity **Consumption:** 660 x 10⁶ lbs.

Fraction of Dispersion: **Fraction of Production Lost:** 0.015

Release Rate (million lbs/yr):

TOXICITY DATA

Acute Toxicity	Dosage	Animal	Route
LD ₅₀	2140 mg/kg	rat	oral
LD _{Lo}	1410 mg/kg	rabbit	skin
	950 mg/kg	guinea pig	subcutaneous

Non-lethal Acute Effects:

Irritant: TD _{Lo}	7 ppm	human	inhalation
----------------------------	-------	-------	------------

Chronic Toxicity

U.S. Occupational Standard:

Carcinogenicity:

Mutagenicity:

Teratogenicity:

TLV:

Other Chronic Effects:

Captan

CHEMICAL NAME

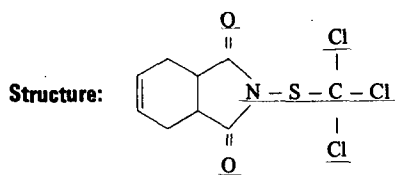
CAS Number: 133-06-2

Chemical Name: N-(Trichloromethyl)thio-4-cyclohexene-1,2-dicarboximide

Synonyms: Cis-N[(trichloromethyl)thio]-4-cyclohexene-1,2-dicarboximide
Tetrahydrophthalimide
Orthocide

Molecular

Formula: $C_9H_8Cl_3NO_2S$



Chemical Properties

Molecular Weight: 300.62

Vapor Pressure: 0.01μ at $25^\circ C$

Boiling Point:

Density: 1.74

Octanol/Water Partition Coefficient: 2.35

Physical State: Solid

Vapor Density:

Melting Point: $175^\circ C$

Solubility: Insoluble (H_2O)

Environmental Persistence

BOD:

Atmospheric Reactivity:

Safety Hazard:

PRODUCTION DATA**Annual U.S.****Production:** 18×10^6 lbs.**Consumption:** 18×10^6 lbs.**Fraction of
Dispersion:** 1.0**Fraction of Pro-
duction Lost:** 0.01**Release Rate (million lbs/yr):** 18.1**TOXICITY DATA****Acute Toxicity**LD₅₀LD_{Lo}**Dosage**

480 mg/kg

9 mg/kg

Animal

rat

mouse

Route

oral

intraperitoneal

Chronic Toxicity**U.S. Occupational Standard:** (air) TWA 5 mg/m³**Carcinogenicity:****Mutagenicity:****Teratogenicity:** 3000 mg/kg pregnant rat oral TLV:
38 mg/kg pregnant rabbit oral
300 mg/kg pregnant hamster oral**Other Chronic Effects:**

Carbaryl

CHEMICAL NAME

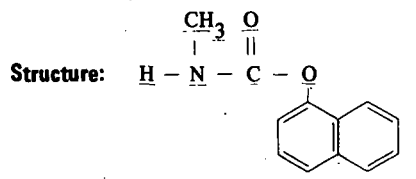
CAS Number: 63-25-2

Chemical Name: Methylcarbamic acid, 1-naphthyl ester

Synonyms: 1-Naphthyl-N-methyl carbamate
Sevin

Molecular

Formula: $C_{12}H_{11}NO_2$



Chemical Properties

Molecular Weight: 201.24

Physical State: Solid

Vapor Pressure:

Vapor Density:

Boiling Point:

Melting Point: 142°C

Density: 1.232 at 20°C/20°C

Solubility:

Octanol/Water Partition Coefficient:

Environmental Persistence

BOD:

Atmospheric Reactivity:

Safety Hazard:

Carbaryl

PRODUCTION DATA

Annual U.S.**Production:** 55 x 10⁶ lbs.**Consumption:** 55 x 10⁶ lbs.**Fraction of****Dispersion:** 1.0**Fraction of Pro-****duction Lost:** 0.015**Release Rate (million lbs/yr):** 55

TOXICITY DATA

Acute ToxicityLD₅₀**Dosage**89 mg/kg
48 mg/kg
500 mg/kg
396 mg/kg
710 mg/kg
280 mg/kg
197 mg/kg
56 mg/kg**Animal**rat
rat
rat
mouse
rabbit
guinea pig
chicken
wild bird**Route**oral
intraperitoneal
unknown
intraperitoneal
oral
oral
oral
oral**Non-lethal Acute Effects:**

Central nervous system—cholinesterase inhibitor

TD_{Lo}

2800 µg/kg

man

oral

Chronic Toxicity**U.S. Occupational Standard:** (air) TWA 5 mg/m³**Carcinogenicity:**TD_{Lo}57000 mg/kg (95 wks)
80 mg/kgrat
ratoral
implant**Mutagenicity:****Teratogenicity:**388 mg/kg pregnant dog
300 mg/kg pregnant guinea pig
50 mg/kg pregnant ratoral
oral
oral**TVL:****Other Chronic Effects:**

Carbofuran

CHEMICAL NAME

CAS Number: 1563-66-2

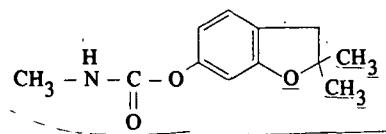
Chemical Name: Methyl carbamic acid

Synonyms: 2,3-Dihydro-2,2-dimethyl-7-benzofuranyl
Carbofuran

Methyl carbamate
Furadan

**Molecular
Formula:** $C_{12}H_{15}NO_3$

Structure:



Chemical Properties

Molecular Weight: 2213

Physical State:

Vapor Pressure: 2×10^{-5} mm at 33°C

Vapor Density:

Boiling Point:

Melting Point: 150 to 152°C

Density: 1.180 at 20°C/20°C

Solubility: Slightly (H_2O)

**Octanol/Water Parti-
tion Coefficient:**

Environmental Persistence

BOD:

Atmospheric Reactivity:

Safety Hazard:

Carbofuran

PRODUCTION DATA

Annual U.S.**Production:** 8×10^6 lbs.**Consumption:****Fraction of****Dispersion:** Estimated as 1.0**Fraction of Pro-****duction Lost:** 0.015**Release Rate (million lbs/yr):** 8.0

TOXICITY DATA

Acute ToxicityLD₅₀**Dosage**

11 mg/kg

10 mg/kg

5300 µg/kg

120 mg/kg

2 mg/kg

19 mg/kg

885 mg/kg

6 mg/kg

415 µg/kg

LC₅₀85 mg/m³52 mg/m³32 mg/m³**Animal**

human

human

rat

rat

mouse

dog

rabbit

chicken

duck

rat

dog

guinea pig

Route

oral

skin

oral

skin

oral

oral

skin

skin

inhalation

inhalation

inhalation

Chronic Toxicity**U.S. Occupational Standard:** (air) TWA 50 µg/m³**Carcinogenicity:****Mutagenicity:****Teratogenicity:****TLV:****Other Chronic Effects:**

Carbon disulfide

CHEMICAL NAME

CAS Number: 75-15-0

Chemical Name: Carbondisulfide

Synonyms: Carbonbisulfide
Carbondisulphide
Weeviltox

**Molecular
Formula:** CS₂

Structure: S = C = S

Chemical Properties

Molecular Weight: 76.13

Physical State: Liquid

Vapor Pressure:

Vapor Density:

Boiling Point: 46.25°C at 760 mm

Melting Point: -111.53°C

Density: 1.2632 at 20°C/4°C

Solubility: Slightly soluble (H₂O)

**Octanol/Water Parti-
tion Coefficient:**

Environmental Persistence

BOD:

Atmospheric Reactivity:

Safety Hazard:

Fire—dangerous

Explosion—dangerous

Carbon disulfide

PRODUCTION DATA

Annual U.S.
Production: 767.8 x 10⁶ lbs. Consumption: 752 x 10⁶ lbs. (1966)

Fraction of
Dispersion: Fraction of Pro-
duction Lost: 0.015

Release Rate (million lbs/yr):

TOXICITY DATA

Acute Toxicity	Dosage	Animal	Route
LD _{Lo}	300 mg/kg	rabbit	subcutaneous
	400 mg/kg	rat	intraperitoneal
LC _{Lo}	4000 ppm (30 minutes)	human	inhalation
TC _{Lo}	50 mg/m ³	human	inhalation

Chronic Toxicity

U.S. Occupational Standard:

(air) TWA	20 ppm
Ceiling concentration	30 ppm
Peak concentration	100 ppm (30 minutes/8 hours)

Carcinogenicity:

Mutagenicity:

Teratogenicity:

TLV:

Other Chronic Effects:

Carbontetrabromide

CHEMICAL NAME

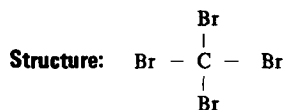
CAS Number: 558-13-4

Chemical Name: Carbon tetrabromide

Synonyms: Tetrabromomethane

Molecular

Formula: CBr₄



Chemical Properties

Molecular Weight: 331.67

Physical State: Solid

Vapor Pressure: 44 mm at 96.3°C

Vapor Density:

Boiling Point: 189.5°C

Melting Point: 90.1°C

Density: 3.42

Solubility: Soluble (H₂O) (785 mg/l at 20°C)

Octanol/Water Partition Coefficient:

Environmental Persistence

BOD: No oxygen consumption in standard testing

Atmospheric Reactivity: Reacts with oxidizing materials.

Safety Hazard: Disaster hazard—dangerous—toxic fumes

Carbontetrabromide

PRODUCTION DATA

Annual U.S.

Production: 1000×10^6 lbs. (1970)

Consumption: 1000×10^6 lbs.

**Fraction of
Dispersion:** 0.01

**Fraction of Pro-
duction Lost:** 0.015

Release Rate (million lbs/yr): 60

TOXICITY DATA

Acute Toxicity

LD₅₀

LD_{Lo}

Dosage

298 mg/kg

1000 mg/kg

Animal

mouse

rat

Route

subcutaneous

oral

Non-lethal Acute Effects:

Narcotic in high concentrations

Chronic Toxicity

U.S. Occupational Standard: (air) TWA 100 ppb

Carcinogenicity:

Mutagenicity:

Teratogenicity:

TLV:

Other Chronic Effects:

Carbon tetrachloride

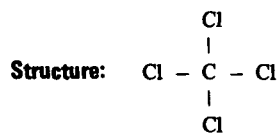
CHEMICAL NAME

CAS Number: 56-23-5

Chemical Name: Carbon tetrachloride

Synonyms: Tetrachloromethane
Perchloromethane
Methane tetrachloride

**Molecular
Formula:** CCl₄



Chemical Properties

Molecular Weight: 153.82

Physical State: Liquid

Vapor Pressure: 115.2 mm at 25°C

Vapor Density: 5.32

Boiling Point: 76.54°C at 760 mm

Melting Point: -22.99°C

Density: 1.5940 at 20°C/4°C

Solubility: Insoluble (H₂O)

**Octanol/Water Parti-
tion Coefficient:** 2.64

Environmental Persistence

BOD: No O₂ consumption in standard BOD test

Atmospheric Reactivity:

Photodegradation: t_{1/2} = 10 to 33 weeks

Safety Hazard:

Carbon tetrachloride

PRODUCTION DATA

Annual U.S. Production: 1,047 x 10 ⁶ lbs. (1973)	Consumption: 989.3 x 10 ⁶ lbs.
Fraction of Dispersion: 0.05	Fraction of Production Lost: 0.015
Release Rate (million lbs/yr): 65	

TOXICITY DATA

Acute Toxicity	Dosage	Animal	Route
LD ₅₀	1770 mg/kg	rat	oral
	4620 mg/kg	mouse	intraperitoneal
LD _{Lo}	6380 mg/kg	rabbit	oral
	3200 mg/kg	mouse	subcutaneous
	1000 mg/kg	dog	oral
	125 mg/kg	dog	intravenous
	4785 mg/kg	cat	subcutaneous
	3000 mg/kg	rabbit	subcutaneous
LC ₅₀	9526 ppm (8 hours)	mouse	inhalation
LC _{Lo}	38,110 ppm (2 hours)	cat	inhalation
	550 ppm	rabbit	inhalation
	20,000 ppm (2 hours)	guinea pig	inhalation
	1,000 ppm	human	inhalation
	4,000 ppm (4 hours)	rat	inhalation

Chronic Toxicity

U.S. Occupational Standard:

(air) TWA:	10 ppm
Ceiling level:	25 ppm
Peak concentration:	200 ppm/5 minutes/4 hours

Carcinogenicity:

Neoplastic effects:

TD _{Lo}	133 gm/kg	rat	subcutaneous
	(25 weeks, intermittent)		

Carcinogenicity:

TD _{Lo}	4800 mg/kg	mouse	oral
	(88 days, intermittent)		

Mutagenicity:

Positive

Teratogenicity:

TLV: 10 ppm
Odor perception: 77 ppm

Other Chronic Effects:

Castor oil

CHEMICAL NAME

CAS Number: 8001-79-4

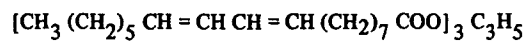
Chemical Name: Castor oil

Synonyms: Ricinus oil
Ricinolein
Glycerides of ricinoleic and isoricinoleic acids

Molecular

Formula: (not given in CAS Registry Number Handbook, since the composition varies)

Structure:



Chemical Properties

Molecular Weight: (Varies)

Physical State: Liquid

Vapor Pressure:

Vapor Density:

Boiling Point: 313°C (decomposes)

Melting Point: -10°C

Density: 0.96 at 25° C/25° C

Solubility: <5.0 gm/l H₂O (Insoluble)

Octanol/Water Partition Coefficient:

Environmental Persistence

BOD:

Atmospheric Reactivity:

Safety Hazard: Fire—slight

Castor oil

PRODUCTION DATA

Annual U.S.**Production:** 100×10^6 lbs.**Consumption:** 198×10^6 lbs.**Fraction of****Dispersion:** 0.64**Fraction of Pro-****duction Lost:** 0.03**Release Rate (million lbs/yr):** 129.7 (1972)

TOXICITY DATA

Acute Toxicity**Dosage****Animal****Route**(Low toxicity)**Chronic Toxicity****U.S. Occupational Standard:****Carcinogenicity:****Mutagenicity:****Teratogenicity:****TLV:****Other Chronic Effects:**

Cellulose acetate

CHEMICAL NAME

CAS Number: 9004-35-7

Chemical Name: Cellulose acetate

Synonyms: CA Acetyl cellulose
Acetate ester of cellulose
Acetic acid, cellulose ester

Molecular Formula: (Exact composition unknown or undetermined)

Structure: (Cellulose ester in which the cellulose is not completely esterified by the acetic acid.)

Chemical Properties

Molecular Weight: (undetermined)

Physical State: Solid

Vapor Pressure: Negligible

Vapor Density:

Boiling Point:

Melting Point: 260°C (decomposes)

Density: 1.27 to 1.34

Solubility: Insoluble (H₂O)

Octanol/Water Partition Coefficient:

Environmental Persistence

BOD:

Atmospheric Reactivity:

Safety Hazard: Fire - moderate

Cellulose acetate

PRODUCTION DATA

Annual U.S.**Production:** 807.1×10^6 lbs.**Consumption:** 820.4×10^6 lbs.**Fraction of****Dispersion:** 0**Fraction of Pro-****duction Lost:** 0.03**Release Rate (million lbs/yr):** 24.6

TOXICITY DATA

Acute Toxicity
(Nontoxic)**Dosage****Animal****Route****Chronic Toxicity****U.S. Occupational Standard:****Carcinogenicity:****Mutagenicity:****Teratogenicity:****TLV:****Other Chronic Effects:**

Chloranil

CHEMICAL NAME

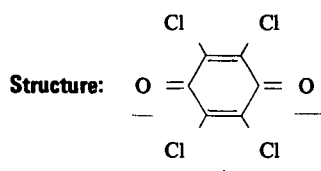
CAS Number: 118-75-2

Chemical Name: 2,3,5,6-Tetrachloro-p-benzoquinone

Synonyms: Tetrachloroquinone
Dow seed disinfectant No. 5

Molecular

Formula: $C_6Cl_4O_2$



Chemical Properties

Molecular Weight: 245.9

Vapor Pressure: 1 mm at 70.7°C

Boiling Point: Sublimes

Density:

Octanol/Water Partition Coefficient:

Physical State: Solid

Vapor Density:

Melting Point: 290°C (sealed tube)

Solubility: Insoluble (H₂O)

Environmental Persistence

BOD:

Atmospheric Reactivity:

Safety Hazard: Disaster hazard—dangerous toxic fumes—chloride

Chloranil

PRODUCTION DATA

**Annual U.S.
Production:**

Consumption:

**Fraction of
Dispersion:**

**Fraction of Pro-
duction Lost:**

Release Rate (million lbs/yr):

TOXICITY DATA

Acute Toxicity

LD₅₀

Dosage
4000 mg/kg

Animal

rat

Route

oral

500 mg/kg

rat

intraperitoneal

LD_{Lo}

4 mg/kg

mouse

intraperitoneal

Chronic Toxicity

U.S. Occupational Standard:

Carcinogenicity:

Mutagenicity:

Teratogenicity:

TLV:

Other Chronic Effects:

Chlordane

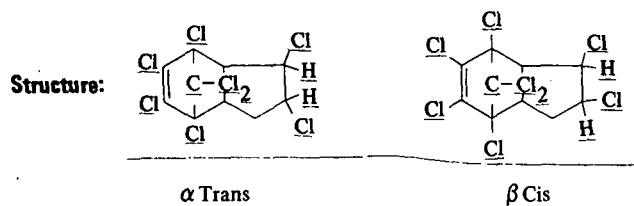
CHEMICAL NAME

CAS Number: 57-74-9

Chemical Name: 1,2,4,5,6,7,8,8-Octachloro-3a,4,7,7a-tetrahydro-4,7-methanoindane

Synonyms: Chlordane
Octaklor
Gama-chlordan

**Molecular
Formula:** $C_{10}H_8Cl_8$



Chemical Properties

Molecular Weight: 409.76

Physical State: Liquid

Vapor Pressure: 1×10^{-5} mm at 25°C

Vapor Density:

Boiling Point: 175°C

Melting Point:

Density: 1.57 to 1.63 at 15.5°C/15.5°C

Solubility: Insoluble (H_2O)

**Octanol/Water Parti-
tion Coefficient:**

Environmental Persistence

BOD:

Atmospheric Reactivity:

Safety Hazard:

Chlordane

PRODUCTION DATA

Annual U.S.**Production:** 25×10^6 lbs.**Consumption:** 25×10^6 lbs.**Fraction of****Dispersion:** 1.0**Fraction of Pro-****duction Lost:** 0.01**Release Rate (million lbs/yr):** 25.3

TOXICITY DATA

Acute Toxicity**LD₅₀****Dosage**

283 mg/kg

700 mg/kg

200 mg/kg

430 mg/kg

100 mg/kg

220 mg/kg

40 mg/kg

Animal

rat

rat

rat

mouse

rabbit

chicken

human

Route

oral

skin

intraperitoneal

oral

oral

oral

oral

LD_{Lo}**Non-lethal Acute Effects:**CNS stimulant**Chronic Toxicity****U.S. Occupational Standard:** (air) TWA 500 $\mu\text{g}/\text{m}^3$ **Carcinogenicity:****Mutagenicity:****Teratogenicity:****TLV:** 0.5 mg/m³**Other Chronic Effects:**

Liver damage

Kidney damage

Chloroacetaldehyde

CHEMICAL NAME

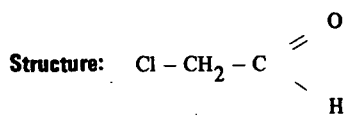
CAS Number: 107-20-0

Chemical Name: Chloroacetaldehyde

Synonyms: Chloroaldehyde
2-Chloro-1-ethanal

Molecular

Formula: C_2H_3ClO



Chemical Properties

Molecular Weight: 78.5

Vapor Pressure: 100 mm at 45°C

Boiling Point: 85 to 85.5°C at 748 mm

Density: 1.19 at 25°C/25°C

Octanol/Water Partition Coefficient:

Physical State: Liquid

Vapor Density:

Melting Point:

Solubility: Soluble (H_2O)

Environmental Persistence

BOD:

Atmospheric Reactivity:

Safety Hazard: Fire—moderate

Chloroacetaldehyde

PRODUCTION DATA

Annual U.S.

Production:

Consumption:

Fraction of

Dispersion:

Fraction of Pro-

duction Lost:

Release Rate (million lbs/yr):

TOXICITY DATA

Acute Toxicity

LD₅₀

Dosage

23 mg/kg

2 mg/kg

21 mg/kg

2 mg/kg

67 mg/kg

1390 mg/kg

636 mg/kg

Animal

rat

rat

mouse

mouse

rabbit

rabbit

guinea pig

Route

oral

intraperitoneal

oral

intraperitoneal

skin

intraperitoneal

intraperitoneal

Non-lethal Acute Effects:

Irritant to eyes, skin respiratory organs and mucous membranes

Chronic Toxicity

U.S. Occupational Standard: (air) TWA 1 ppm

Carcinogenicity:

Mutagenicity:

Teratogenicity:

TLV:

Other Chronic Effects:

Chloroacetic acid

CHEMICAL NAME

CAS Number: 79-11-8

Chemical Name: Chloroacetic acid

Synonyms: Monochloroacetic acid

Molecular

Formula: $\text{C}_2\text{H}_3\text{ClO}_2$

Structure:

$$\begin{array}{c} \text{Cl} \quad \text{O} \\ | \quad || \\ \text{CH}_2 - \text{C} - \text{OH} \end{array}$$

Chemical Properties

Molecular Weight: 94.50

Vapor Pressure: 1 mm at 23°C

Boiling Point: 187.85°C

Density: 1.4043 at 40°C/4°C

Octanol/Water Partition Coefficient: 1.83

Physical State: Solid

Vapor Density: 3.26

Melting Point: 63.0°C

Solubility: Very soluble (H₂O)

Environmental Persistence

BOD:

Atmospheric Reactivity:

Safety Hazard: Disaster hazard—dangerous—toxic fumes—phosgene and chlorides

Chloroacetic acid

PRODUCTION DATA

Annual U.S.**Production:** 84×10^6 lbs. (capacity)**Consumption:** 96×10^6 lbs. (1975)**Fraction of****Dispersion:** 0.01**Fraction of Pro-****duction Lost:** 0.015**Release Rate (million lbs/yr):**

TOXICITY DATA

Acute ToxicityLD₅₀**Dosage**

76 mg/kg

5 mg/kg

500 mg/kg

Animal

rat

rat

mouse

Route

oral

subcutaneous

intraperitoneal

Non-lethal Acute Effects:Irritant**Chronic Toxicity****U.S. Occupational Standard:****Carcinogenicity:****Mutagenicity:****Teratogenicity:****TLV:****Other Chronic Effects:**

Chloroacetophenone

CHEMICAL NAME

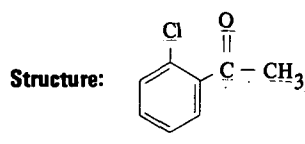
CAS Number: 532-27-4

Chemical Name: 2-Chloroacetophenone

Synonyms: o-Chlorophenyl methyl ketone

Molecular

Formula: C_8H_7ClO



Chemical Properties

Molecular Weight: 154.60

Physical State: Solid or liquid

Vapor Pressure: 1 mm at 48°C

Vapor Density:

Boiling Point: 227 to 228°C at 738 mm

Melting Point:

Density: 1.2016 at 17°C/4°C

Solubility:

Octanol/Water Partition Coefficient: 2.08

Environmental Persistence

BOD:

Atmospheric Reactivity:

Safety Hazard:

Chloroacetophenone

PRODUCTION DATA

Annual U.S.
Production:

Consumption:

Fraction of
Dispersion:

Fraction of Pro-
duction Lost:

Release Rate (million lbs/yr):

TOXICITY DATA

Acute Toxicity

Dosage

Animal

Route

Non-lethal Acute Effects:

TC_{Lo}

Irritant

93 mg/m³ (3 minutes)

20 mg/m³ (3 minutes)

human

human

inhalation

inhalation

Chronic Toxicity

U.S. Occupational Standard: (air) Ceiling concentration 0.05 ppm

Carcinogenicity: Not tested

Mutagenicity:

Teratogenicity:

TLV:

Other Chronic Effects:

m-Chloroaniline

CHEMICAL NAME

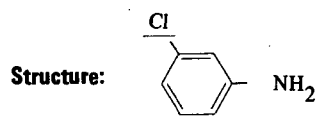
CAS Number: 108-42-9

Chemical Name: m-Chloroaniline

Synonyms: 3-Chloroaniline

Molecular

Formula: C_6H_5ClN



Chemical Properties

Molecular Weight: 127.52

Vapor Pressure: 1 mm at 63.5°C

Boiling Point: 230.5°C

Density: 1.223 at 15°C/15°C

Octanol/Water Partition Coefficient:

Physical State: Liquid

Vapor Density:

Melting Point: -10.6°C

Solubility: Insoluble (H_2O)

Environmental Persistence

BOD:

Atmospheric Reactivity:

Safety Hazard:

m-Chloroaniline

PRODUCTION DATA

**Annual U.S.
Production:**

Consumption:

**Fraction of
Dispersion:**

**Fraction of Pro-
duction Lost:**

Release Rate (million lbs/yr):

TOXICITY DATA

Acute Toxicity
LD₅₀
LD_{Lo}

Dosage
880 mg/kg
310 mg/kg

Animal
rat
cat

Route
oral
subcutaneous

Chronic Toxicity

U.S. Occupational Standard:

Carcinogenicity:

Mutagenicity:

Teratogenicity:

TLV:

Other Chronic Effects:

o-Chloroaniline

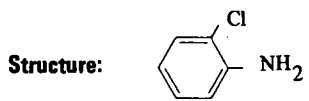
CHEMICAL NAME

CAS Number: 95-51-2

Chemical Name: o-Chloroaniline

Synonyms: 2-Chloroaniline

**Molecular
Formula:** C₆H₆ClN



Chemical Properties

Molecular Weight: 127.52

Physical State: Liquid

Vapor Pressure: 1 mm at 63.5°C

Vapor Density:

Boiling Point: 208.84

Melting Point: 2.3°C

Density:

Solubility: Insoluble (H₂O)

**Octanol/Water Parti-
tion Coefficient:** 1.2125 at 20°C/4°C

Environmental Persistence

BOD:

Atmospheric Reactivity:

Safety Hazard:

o-Chloroaniline

PRODUCTION DATA

**Annual U.S.
Production:**

Consumption:

**Fraction of
Dispersion:**

**Fraction of Pro-
duction Lost:**

Release Rate (million lbs/yr):

TOXICITY DATA

Acute Toxicity
LD₅₀
LD_{Lo}

Dosage
256 mg/kg
310 mg/kg

Animal
mouse
cat

Route
oral
subcutaneous

Chronic Toxicity

U.S. Occupational Standard:

Carcinogenicity:

Mutagenicity:

Teratogenicity:

TLV:

Other Chronic Effects:

p-Chloroaniline

CHEMICAL NAME

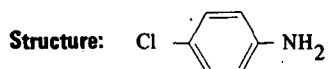
CAS Number: 106-47-8

Chemical Name: p-Chloroaniline

Synonyms: 4-Chloroaniline

Molecular

Formula: C_6H_5ClN



Chemical Properties

Molecular Weight: 127.52

Vapor Pressure: 1 mm at 63.5°C

Boiling Point: 232°C at 760mm

Density: 1.429 at 19°C/4°C

Octanol/Water Partition Coefficient:

Physical State: Solid

Vapor Density:

Melting Point: 69.5°C

Solubility: Soluble (hot H₂O)

Environmental Persistence

BOD:

Atmospheric Reactivity:

Safety Hazard:

p-Chloroaniline

PRODUCTION DATA

**Annual U.S.
Production:**

Consumption:

**Fraction of
Dispersion:**

**Fraction of Pro-
duction Lost:**

Release Rate (million lbs/yr):

TOXICITY DATA

Acute Toxicity

LD₅₀

Dosage

300 mg/kg

420 mg/kg

100 mg/kg

100 mg/kg

LD_{Lo}

125 mg/kg

36 mg/kg

Animal

rat

rat

mouse

wild bird

cat

rabbit

Route

oral

intraperitoneal

oral

oral

subcutaneous

skin

Chronic Toxicity

U.S. Occupational Standard:

Carcinogenicity:

Mutagenicity:

Teratogenicity:

TLV:

Other Chronic Effects:

o-Chlorobenzaldehyde

CHEMICAL NAME

CAS Number: 89-98-5

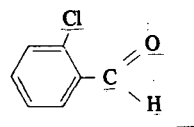
Chemical Name: o-Chlorobenzaldehyde

Synonyms: 2-Chlorobenzaldehyde

Molecular

Formula: C_7H_5ClO

Structure:



Chemical Properties

Molecular Weight: 140.57

Vapor Pressure: 10 mm at 25°C

Boiling Point: 211.9°C at 760 mm

Density: 1.2483 at 20°C/4°C

Octanol/Water Partition Coefficient:

Physical State: Liquid

Vapor Density:

Melting Point: 12.39°C

Solubility: Slightly soluble (H₂O)

Environmental Persistence

BOD:

Atmospheric Reactivity:

Safety Hazard:

o-Chlorobenzaldehyde

PRODUCTION DATA

**Annual U.S.
Production:**

Consumption:

**Fraction of
Dispersion:**

**Fraction of Pro-
duction Lost:**

Release Rate (million lbs/yr):

TOXICITY DATA

Acute Toxicity
LD_{Lo}

Dosage
10 mg/kg

Animal
mouse

Route
intraperitoneal

Chronic Toxicity

U.S. Occupational Standard:

Carcinogenicity: Not tested

Mutagenicity:

Teratogenicity:

TLV:

Other Chronic Effects:

p-Chlorobenzaldehyde

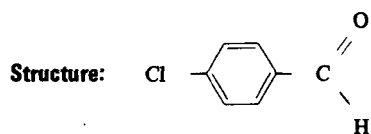
CHEMICAL NAME

CAS Number: 104-88-1

Chemical Name: p-Chlorobenzaldehyde

Synonyms:

**Molecular
Formula:** C_7H_5ClO



Chemical Properties

Molecular Weight: 140.57

Vapor Pressure:

Boiling Point: 213 to 214°C at 760 mm

Density: 1.196 at 61°C/4°C

**Octanol/Water Parti-
tion Coefficient:**

Physical State: Solid

Vapor Density:

Melting Point: 47.5°C

Solubility: Soluble (H_2O)

Environmental Persistence

BOD:

Atmospheric Reactivity:

Safety Hazard:

p-Chlorobenzaldehyde

PRODUCTION DATA

**Annual U.S.
Production:**

Consumption:

**Fraction of
Dispersion:**

**Fraction of Pro-
duction Lost:**

Release Rate (million lbs/yr):

TOXICITY DATA

Acute Toxicity

Dosage

Animal

Route

Chronic Toxicity

U.S. Occupational Standard:

Carcinogenicity:

Mutagenicity:

Teratogenicity:

TLV:

Other Chronic Effects:

Chlorobenzene

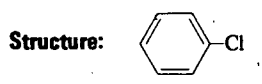
CHEMICAL NAME

CAS Number: 108-90-7

Chemical Name: Chlorobenzene

Synonyms: Phenyl chloride
Monochlorobenzene
Chlorobenzol

**Molecular
Formula:** C_6H_5Cl



Chemical Properties

Molecular Weight: 112.56

Physical State: Liquid

Vapor Pressure: 12.14 mm at 25°C

Vapor Density: 3.88

Boiling Point: 131.7°C at 760 mm

Melting Point: -45.6°C

Density: 1.1058 at 20°C/4°C

Solubility: Insoluble (H₂O)

**Octanol/Water Parti-
tion Coefficient:** 2.84

Environmental Persistence

BOD: 1% of theoretical

Atmospheric Reactivity: Reacts with oxidizing materials

Safety Hazard: Fire—dangerous
Explosion—moderate

Chlorobenzene

PRODUCTION DATA

Annual U.S.**Production:** 690 x 10⁶ lbs. (1976 capacity)**Consumption:****Fraction of****Dispersion:****Fraction of Pro-****duction Lost:** 0.015**Release Rate (million lbs/yr):**

TOXICITY DATA

Acute ToxicityLD₅₀**Dosage**

2910 mg/kg

2830 mg/kg

4000 mg/kg

Animal

rat

rabbit

rat

Route

oral

oral

subcutaneous

Non-lethal Acute Effects:Strong narcotic**Chronic Toxicity****U.S. Occupational Standard:** (air) TWA 75 ppm**Carcinogenicity:****Mutagenicity:****Teratogenicity:****TLV:** 75 ppm**Other Chronic Effects:**

Kidney damage

Liver damage

Chlorobenzilate

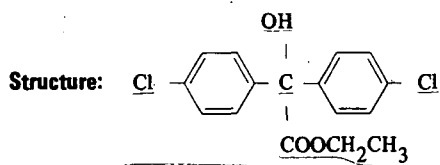
CHEMICAL NAME

CAS Number: 510-15-6

Chemical Name: 4,4'-Dichlorobenzilic acid, ethyl ester

Synonyms: Acaradben
Benzilic acid, 4,4'-dichloroethyl ester
Chlorobenzilate

Molecular Formula: $C_{16}H_{14}Cl_2O_3$



Chemical Properties

Molecular Weight: 325.20

Physical State: Liquid

Vapor Pressure:

Vapor Density:

Boiling Point: 148°C at 0.04 mm

Melting Point:

Density:

Solubility:

Octanol/Water Partition Coefficient:

Environmental Persistence

BOD:

Atmospheric Reactivity:

Safety Hazard:

Chlorobenzilate

PRODUCTION DATA

Annual U.S.
Production: 2×10^6 lbs.

Consumption:

Fraction of
Dispersion: 1.0

Fraction of Pro-
duction Lost: 0.015

Release Rate (million lbs/yr): 2.0

TOXICITY DATA

Acute Toxicity

LD₅₀

Dosage

700 mg/kg

724 mg/kg

LC₅₀

500 ppm

Animal

rat

mouse

mammal

Route

oral

oral

inhalation

Chronic Toxicity

U.S. Occupational Standard:

Carcinogenicity: TD_{Lo}

71 mg/kg (82 weeks) mouse

oral

Mutagenicity:

Teratogenicity:

TLV:

Other Chronic Effects:

Chlorobenzoic acid

CHEMICAL NAME

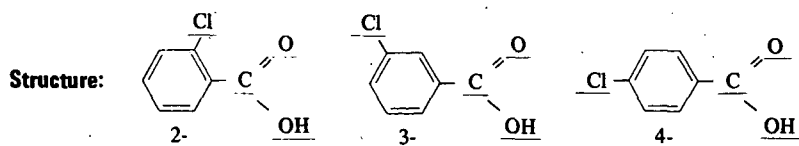
CAS Number: 2-: 118-91-2
3-: 535-80-8
4-: 74-11-3

Chemical Name: Chlorobenzoic acid (2-, 3- and 4-)

Synonyms: Chlorobenzoic acid (o-, m-, and p-)

Molecular

Formula: $C_7H_5ClO_2$



Chemical Properties

Molecular Weight: 156.57

Vapor Pressure:

Boiling Point: (2-) and (3-) sublime

Density: (2-) 1.544 gm/ml (20°C)
(3-) 1.496 gm/mg (25°C/4°C)

Octanol/Water Partition Coefficient:

Physical State: Solid

Vapor Density:

Melting Point: (2-) 142°C (3-) 158°C
(4-) 243°C

Solubility: (2-) Slightly soluble in hot water
(3-) Very soluble in hot water
(4-) Slightly soluble in hot water

Environmental Persistence

BOD:

Atmospheric Reactivity:

Safety Hazard:

Chlorobenzoic acid

PRODUCTION DATA

**Annual U.S.
Production:**

Consumption:

**Fraction of
Dispersion:**

**Fraction of Pro-
duction Lost:**

Release Rate (million lbs/yr):

TOXICITY DATA

Acute Toxicity

Dosage

Animal

Route

Chronic Toxicity

U.S. Occupational Standard:

Carcinogenicity:

Mutagenicity:

Teratogenicity:

TLV:

Other Chronic Effects:

o-Chlorobenzoyl chloride

CHEMICAL NAME

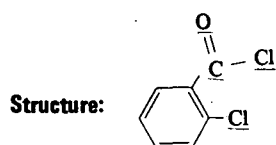
CAS Number: 609-65-4

Chemical Name: o-Chlorobenzoyl chloride

Synonyms: 2-Chlorobenzoic acid chloride

Molecular

Formula: C₇H₄Cl₂O



Chemical Properties

Molecular Weight: 175.02

Vapor Pressure:

Boiling Point: 238°C at 760 mm

Density: 1.270 to 1.280 at 25°C/15°C

Octanol/Water Partition Coefficient:

Physical State: Liquid

Vapor Density: 5.55

Melting Point: -4 to -6°C

Solubility: Decomposes (H₂O)

Environmental Persistence

BOD:

Atmospheric Reactivity:

Safety Hazard:

o-Chlorobenzoyl chloride

PRODUCTION DATA

**Annual U.S.
Production:**

Consumption:

**Fraction of
Dispersion:**

**Fraction of Pro-
duction Lost:**

Release Rate (million lbs/yr):

TOXICITY DATA

Acute Toxicity

Dosage

Animal

Route

Chronic Toxicity

U.S. Occupational Standard:

Carcinogenicity: Not tested

Mutagenicity:

Teratogenicity:

TLV:

Other Chronic Effects:

o-Chlorobenzylidene malonitrile

CHEMICAL NAME

CAS Number: 18270-61-6

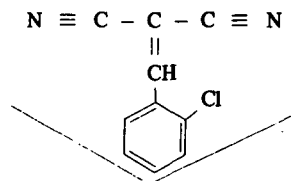
Chemical Name: o-Chlorobenzylidene malonitrile

Synonyms: OCBM

Molecular

Formula: $C_{10}H_5ClN_2$

Structure:



Chemical Properties

Molecular Weight: 188.62

Physical State: Solid

Vapor Pressure:

Vapor Density:

Boiling Point: 313°C

Melting Point:

Density:

Solubility:

Octanol/Water Partition Coefficient:

Environmental Persistence

BOD:

Atmospheric Reactivity:

Safety Hazard:

o-Chlorobenzylidene malononitrile

PRODUCTION DATA

Annual U.S.

Production:

Consumption:

Fraction of

Dispersion:

Fraction of Pro-

duction Lost:

Release Rate (million lbs/yr):

TOXICITY DATA

Acute Toxicity

LD₅₀

Dosage

178 mg/kg

48 mg/kg

35 mg/kg

LD_{Lo}

282 mg/kg

50 mg/kg

8 mg/kg

Animal

rat

rat

rat

mouse

mouse

rabbit

Route

oral

intraperitoneal

intravenous

oral

intraperitoneal

intravenous

Non-lethal Acute Effects:

Central nervous system—TC_{Lo} 1500 mg/m³ (90 minutes) human inhalation

Chronic Toxicity

U.S. Occupational Standard:

Carcinogenicity:

Mutagenicity:

Teratogenicity:

TLV:

Other Chronic Effects:

Chlorodifluoroethane

CHEMICAL NAME

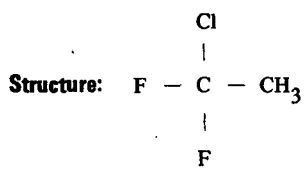
CAS Number: 75-68-3

Chemical Name: Chlorodifluoroethane

Synonyms: 1 Chloro-1, 1-difluoroethane

Molecular

Formula: C₂H₃ClF₂



Chemical Properties

Molecular Weight: 100.50

Vapor Pressure:

Boiling Point: -9.2°C

Density:

Octanol/Water Partition Coefficient:

Physical State: Gas

Vapor Density:

Melting Point: -130.8°C

Solubility: Insoluble (H₂O)

Environmental Persistence

BOD:

Atmospheric Reactivity:

Safety Hazard:

Chlorodifluoroethane

PRODUCTION DATA

**Annual U.S.
Production:** 0.2×10^6 lbs. (1972)

Consumption:

**Fraction of
Dispersion:** 0.01

**Fraction of Pro-
duction Lost:** 0.015

Release Rate (million lbs/yr):

TOXICITY DATA

Acute Toxicity

Dosage

Animal

Route

Asphyxiant at high concentrations

Chronic Toxicity

U.S. Occupational Standard:

Carcinogenicity:

Mutagenicity:

Teratogenicity:

TLV:

Other Chronic Effects:

Chlorodifluoromethane

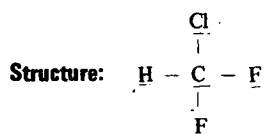
CHEMICAL NAME

CAS Number: 75-45-6

Chemical Name: Chlorodifluoromethane

Synonyms: Freon-22
Fluorocarbon-22

**Molecular
Formula:** CHClF_2



Chemical Properties

Molecular Weight: 86.5

Vapor Pressure: 2210.6 mm at 25°C

Boiling Point: -40.8°C

Density: 3.87 that of air at 0°C

**Octanol/Water Parti-
tion Coefficient:**

Physical State: Gas

Vapor Density:

Melting Point: -160°C

Solubility: Partly soluble (H_2O)

Environmental Persistence

BOD:

Atmospheric Reactivity:

Safety Hazard:

Chlorodifluoromethane

PRODUCTION DATA

**Annual U.S.
Production:** 80 x 10⁶ lbs. (1972)

Consumption:

**Fraction of
Dispersion:**

**Fraction of Pro-
duction Lost:** 0.015

Release Rate (million lbs/yr):

TOXICITY DATA

Acute Toxicity

Dosage

Animal

Route

Asphyxiant at high concentrations

Chronic Toxicity

U.S. Occupational Standard:

Carcinogenicity:

Mutagenicity:

Teratogenicity:

TLV:

Other Chronic Effects:

2-Chloro-4-ethylamino-6-isopropylamino-s-triazine

CHEMICAL NAME

CAS Number: 1912-24-9

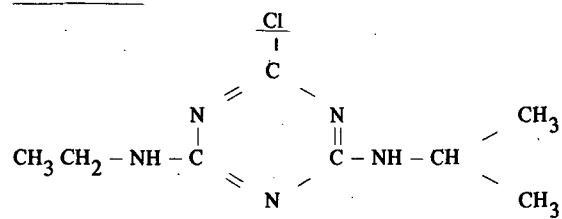
Chemical Name: 2-Chloro-4-(ethylamino)-6-isopropylamino-s-triazine

Synonyms: Atrazine

Molecular

Formula: $C_8H_{14}ClN_5$

Structure:



Chemical Properties

Molecular Weight: 215.72

Physical State: Solid

Vapor Pressure: 3.0×10^{-7} Torrs at 20°C

Vapor Density:

Boiling Point:

Melting Point: 175°C

Density:

Solubility: Slightly (hot H₂O)

Octanol/Water Partition Coefficient:

Environmental Persistence

BOD:

Atmospheric Reactivity:

Safety Hazard:

2-Chloro-4-ethylamino-6-isopropylamino-s-triazine

PRODUCTION DATA

Annual U.S. Production:	110 x 10 ⁶ lbs.	Consumption:	110 x 10 ⁶ lbs.
Fraction of Dispersion:	1.0	Fraction of Production Lost:	0.01
Release Rate (million lbs/yr): 111.1			

TOXICITY DATA

Acute Toxicity	Dosage	Animal	Route
LD ₅₀	1750 mg/kg	rat	oral
	1750 mg/kg	mouse	oral
	3080 mg/kg	rat	unknown

Chronic Toxicity

U.S. Occupational Standard:

Carcinogenicity:

Mutagenicity:

Teratogenicity:

TLV:

Other Chronic Effects:

Chloroform

CHEMICAL NAME

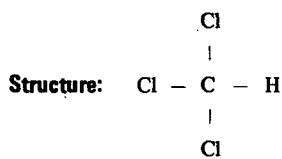
CAS Number: 67-66-3

Chemical Name: Chloroform

Synonyms: Trichloromethane
Methenyl chloride

Molecular

Formula: CHCl_3



Chemical Properties

Molecular Weight: 119.39

Physical State: Liquid

Vapor Pressure: 200 mm at 25.9°C

Vapor Density: 4.12

Boiling Point: 61.26°C

Melting Point: -63.5°C

Density: 1.49845 at 15°C

Solubility: Soluble (H_2O) (8.0 gm/l)

Octanol/Water Partition Coefficient: 1.17

Environmental Persistence

BOD: 4% of theoretical for 5 days at 20°C
Total theoretical oxygen demand gm/gm=0.19

Atmospheric Reactivity: Activity toward OH: $t_{1/2} = >1$ yr.

Safety Hazard: Fire-Slight

Disaster hazard—dangerous—toxic fumes—phosgene

Chloroform

PRODUCTION DATA

Annual U.S.
Production: 234.7×10^6 lbs. Consumption: 234.7

Fraction of
Dispersion: 0.70 Fraction of Pro-
duction Lost: 0.015

Release Rate (million lbs/yr): 167.8

TOXICITY DATA

Acute Toxicity	Dosage	Animal	Route
LD ₅₀	800 mg/kg		
	704 mg/kg		
LD _{Lo}	2500 mg/kg	mouse	oral
	1000 mg/kg	dog	oral
	75 mg/kg	dog	intravenous
	800 mg/kg	rabbit	subcutaneous
LC ₅₀	28 ppm	mouse	inhalation
	100 ppm	dog	inhalation
	59 ppm	rabbit	inhalation
LC _{Lo}	8000 ppm (4 hours)	rat	inhalation
	20000 ppm (2 hours)	guinea pig	inhalation

Non-lethal Acute Effects:

Systemic effects			
(Liver or kidney)	10 ppm (1 year)	human	inhalation
Motor reflexes—rabbit			
Kidney, liver and vascular lesions			

Chronic Toxicity

U.S. Occupational Standard: (air) TWA 50 ppm, but NIOSH has recommended TWA 10 ppm

Carcinogenicity: TD _{Lo}	18 gm/kg (120 days)	mouse	oral
-----------------------------------	---------------------	-------	------

Mutagenicity:

Teratogenicity:

TLV:

Other Chronic Effects: Irritation of eyes and mucous membranes
CNS paralysis after prolonged exposure
Placental lesions
Alteration in neuroendocrine system

Chloronaphthalene

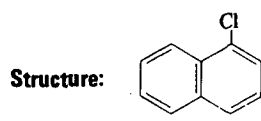
CHEMICAL NAME

CAS Number: 90-13-1

Chemical Name: 1-Chloronaphthalene

Synonyms: α -Chloronaphthalene
Chloronaphthalene

Molecular
Formula: C₁₀H₇Cl



Chemical Properties

Molecular Weight: 162.61

Vapor Pressure: 1 mm at 80.6°C

Boiling Point: 259°C at 753mm

Density: 1.1938 at 20°C/4°C

Octanol/Water Parti-
tion Coefficient:

Physical State: Liquid

Vapor Density:

Melting Point: -2.3°C

Solubility: Insoluble (H₂O)

Environmental Persistence

BOD:

Atmospheric Reactivity:

Safety Hazard: Fire—slight

Chloronaphthalene

PRODUCTION DATA

Annual U.S.**Production:** 5×10^6 lbs. (1972)**Consumption:****Fraction of****Dispersion:** 0.01**Fraction of Pro-****duction Lost:** 0.015**Release Rate (million lbs/yr):**

TOXICITY DATA

Acute ToxicityLD₅₀**Dosage**

1540 mg/kg

1091 mg/kg

Animal

rat

mouse

Route

oral

oral

Non-lethal Acute Effects:

Conjunctivitis

Jaundice

Vomiting

Highly toxic by ingestion, inhalation, and skin absorption

Strong irritant

Chronic Toxicity**U.S. Occupational Standard:**(Tolerance: 2 mg/m³)**Carcinogenicity:** Not tested**Mutagenicity:****Teratogenicity:****TLV:****Other Chronic Effects:**

m-Chloronitrobenzene

CHEMICAL NAME

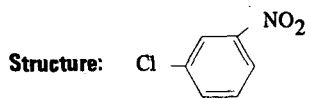
CAS Number: 88-73-3

Chemical Name: 1-Chloro-3-nitrobenzene

Synonyms: m-Chloronitrobenzene

Molecular

Formula: $C_6H_4ClNO_2$



Chemical Properties

Molecular Weight: 157.6

Physical State: Solid

Vapor Pressure: <10 mm at 25°C

Vapor Density:

Boiling Point: 236°C

Melting Point: 44°C

Density: 1.534 at 20°C/4°C

Solubility: Insoluble (H₂O)

Octanol/Water Partition Coefficient: 2.41

Environmental Persistence

BOD:

Atmospheric Reactivity: Reacts with oxidizing materials

Safety Hazard: Fire—slight

Disaster hazard—dangerous—toxic fumes—nitrate and phosgene

m-Chloronitrobenzene

PRODUCTION DATA

Annual U.S. Production: 142×10^6 lbs. (1967) **Consumption:**

Fraction of Dispersion: 0.01 **Fraction of Production Lost:** 0.015

Release Rate (million lbs/yr):

TOXICITY DATA

Acute Toxicity	Dosage	Animal	Route
LD ₅₀	288 mg/kg	rat	oral
	135 mg/kg	mouse	oral
Non-lethal Acute Effects:			
Eye TC _{Lo}	12 mg/m ³	human	inhalation
Intoxication			
Methemoglobin, cyanosis and blood changes			
Poisoning by pulmonary route			
<u>Irritation of nose and throat</u>			

Chronic Toxicity

U.S. Occupational Standard:

Carcinogenicity: Not tested

Mutagenicity:

Teratogenicity: TLV: 1 ppm

Other Chronic Effects:

o-Chloronitrobenzene

CHEMICAL NAME

CAS Number: 88-73-3

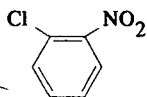
Chemical Name: 1-Chloro-2-nitrobenzene

Synonyms: 1-Chloro-2-nitrobenzene

Molecular

Formula: $C_6H_4ClNO_2$

Structure:



Chemical Properties

Molecular Weight: 157.6

Physical State: Liquid

Vapor Pressure:

Vapor Density:

Boiling Point: 245 to 246°C at 760 mm

Melting Point: 33.5 to 35.0°C

Density: 1.305

Solubility: Insoluble (H₂O)

Octanol/Water Partition Coefficient:

Environmental Persistence

BOD:

Atmospheric Reactivity: Reacts with oxidizing materials

Safety Hazard: Fire—slight

Disaster hazard—dangerous—toxic fumes—nitrate and phosgene

o-Chloronitrobenzene

PRODUCTION DATA

Annual U.S. Production: 142×10^6 lbs. (1967) **Consumption:**

Fraction of Dispersion: 0.01 **Fraction of Production Lost:** 0.015

Release Rate (million lbs/yr):

TOXICITY DATA

Acute Toxicity	Dosage	Animal	Route
LD ₅₀	288 mg/kg	rat	oral
	135 mg/kg	mouse	oral

Non-lethal Acute Effects:

Anemia
Irritation of eyes, nose, and throat

Chronic Toxicity

U.S. Occupational Standard:

Carcinogenicity:

Mutagenicity:

Teratogenicity:

TLV: 1 ppm

Other Chronic Effects:

p-Chloronitrobenzene

CHEMICAL NAME

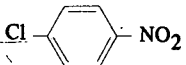
CAS Number: 100-00-5

Chemical Name: 1-Chloro-4-nitrobenzene

Synonyms: p-Chloronitrobenzene
p-Nitrochlorobenzene

Molecular

Formula: $C_6H_4ClNO_2$

Structure: 

Chemical Properties

Molecular Weight: 157.6

Physical State: Solid

Vapor Pressure: 10 mm at 25°C

Vapor Density: 5.43

Boiling Point: 242°C

Melting Point: 83°C

Density: 1.520 at 22°C/4°C

Solubility: Insoluble (H₂O)

Octanol/Water Partition Coefficient: 2.41

Environmental Persistence

BOD:

Atmospheric Reactivity:

Safety Hazard: Fire—slight

Disaster hazard—dangerous—toxic fumes—nitrate and chlorides

p-Chloronitrobenzene

PRODUCTION DATA

**Annual U.S.
Production:** 142×10^6 lbs. (1967)

Consumption:

**Fraction of
Dispersion:** 0.01

**Fraction of Pro-
duction Lost:** 0.015

Release Rate (million lbs/yr):

TOXICITY DATA

Acute Toxicity
LD₅₀

Dosage
1414 mg/kg
420 mg/kg

Animal
mouse
mammal

Route
oral
intraperitoneal

Chronic Toxicity

U.S. Occupational Standard: (air) TWA 1 mg/m³

Carcinogenicity:

Mutagenicity:

Teratogenicity:

TLV:

Other Chronic Effects:

1-Chloro-1-nitropropane

CHEMICAL NAME

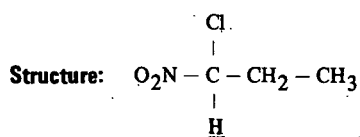
CAS Number: 600-25-9

Chemical Name: 1-Chloro-1-nitropropane

Synonyms:

Molecular

Formula: $C_3H_6ClNO_2$



Chemical Properties

Molecular Weight: 123.5

Physical State: Liquid

Vapor Pressure:

Vapor Density: 4.26

Boiling Point: 139.5°C

Melting Point:

Density: 1.209 at 20°C/20°C

Solubility: Slightly soluble (H_2O)

Octanol/Water Partition Coefficient:

Environmental Persistence

BOD:

Atmospheric Reactivity:

Safety Hazard: Fire—moderate
Explosion—dangerous
Disaster hazard—toxic fumes

1-Chloro-1-nitropropane

PRODUCTION DATA

Annual U.S.
Production:

Consumption:

Fraction of
Dispersion:

Fraction of Pro-
duction Lost:

Release Rate (million lbs/yr):

TOXICITY DATA

Acute Toxicity

LD₅₀

Dosage

510 mg/kg

Animal

mouse

Route

oral

LD_{Lo}

165 mg/kg

mouse

subcutaneous

50 mg/kg

rabbit

oral

Non-lethal Acute Effects:

Irritant

Severe pulmonary edema after inhalation or ingestion

Chronic Toxicity

U.S. Occupational Standard: (air) 20 ppm

Carcinogenicity:

Mutagenicity:

Teratogenicity:

TLV:

Other Chronic Effects:

m-Chlorophenol

CHEMICAL NAME

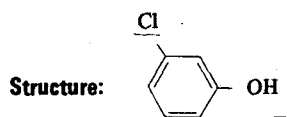
CAS Number: 108-43-0

Chemical Name: m-Chlorophenol

Synonyms: 3-Chlorophenol

Molecular

Formula: C_6H_5ClO



Chemical Properties

Molecular Weight: 128.6

Physical State: Solid

Vapor Pressure: 1 mm at 44.2°C

Vapor Density:

Boiling Point: 214°C

Melting Point: 33°C

Density: 1.268 at 25°C

Solubility: Slightly soluble (H_2O)

Octanol/Water Partition Coefficient:

Environmental Persistence

BOD:

Atmospheric Reactivity:

Safety Hazard: Disaster hazard—toxic fumes

m-Chlorophenol

PRODUCTION DATA

**Annual U.S.
Production:**

Consumption:

**Fraction of
Dispersion:**

**Fraction of Pro-
duction Lost:**

Release Rate (million lbs/yr):

TOXICITY DATA

Acute Toxicity
LD₅₀

Dosage
570 mg/kg
355 mg/kg
1390 mg/kg

Animal
rat
rat
rat

Route
oral
intraperitoneal
subcutaneous

Chronic Toxicity

U.S. Occupational Standard:

Carcinogenicity:

Mutagenicity:

Teratogenicity:

TLV:

Other Chronic Effects:
Irritant

o-Chlorophenol

CHEMICAL NAME

CAS Number: 95-57-8

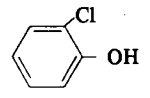
Chemical Name: o-Chlorophenol

Synonyms: 2-Chlorophenol

Molecular

Formula: C_6H_5ClO

Structure:



Chemical Properties

Molecular Weight: 128.6

Vapor Pressure: 2.97 at 25°C

Boiling Point: 174.5°C

Density: 1.256 at 25°C/25°C

Octanol/Water Partition Coefficient: 2.19

Physical State: Liquid

Vapor Density:

Melting Point: 7°C

Solubility: Slightly (H_2O)

Environmental Persistence

BOD: 100% of theoretical after 36 days

Atmospheric Reactivity: Reacts with oxidizing materials

Safety Hazard: Fire—moderate

o-Chlorophenol

PRODUCTION DATA

**Annual U.S.
Production:**

Consumption:

**Fraction of
Dispersion:**

**Fraction of Pro-
duction Lost:**

Release Rate (million lbs/yr):

TOXICITY DATA

Acute Toxicity

LD₅₀

Dosage

670 mg/kg

230 mg/kg

950 mg/kg

670 mg/kg

440 mg/kg

LD_{Lo}

950 mg/kg

800 mg/kg

Animal

rat

rat

rat

mouse

mammal

rabbit

guinea pig

Route

oral

intraperitoneal

subcutaneous

oral

oral

subcutaneous

subcutaneous

Chronic Toxicity

U.S. Occupational Standard:

Carcinogenicity: Neoplastic effect TD_{Lo} 38 gm/kg (12 weeks) mouse skin

Mutagenicity:

Teratogenicity:

TLV:

Other Chronic Effects:

p-Chlorophenol

CHEMICAL NAME

CAS Number: 106-48-9

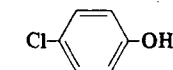
Chemical Name: p-Chlorophenol

Synonyms: 4-Chlorophenol

Molecular

Formula: C_6H_5ClO

Structure:



Chemical Properties

Molecular Weight: 128.56

Vapor Pressure: 1 mm at 49.8°C

Boiling Point: 220°C

Density: 1.246 at 60°C/25°C

Octanol/Water Partition Coefficient: 2.44

Physical State: Solid

Vapor Density:

Melting Point: 42 to 43°C

Solubility: Insoluble (H_2O)

Environmental Persistence

BOD: 100% of theoretical after 36 days

Atmospheric Reactivity:

Safety Hazard: Fire—moderate

p-Chlorophenol

PRODUCTION DATA

**Annual U.S.
Production:**

Consumption:

**Fraction of
Dispersion:**

**Fraction of Pro-
duction Lost:**

Release Rate (million lbs/yr):

TOXICITY DATA

Acute Toxicity
LD₅₀

Dosage
500 mg/kg
281 mg/kg
1030 mg/kg

Animal
rat
rat
rat

Route
oral
intraperitoneal
subcutaneous

Chronic Toxicity

U.S. Occupational Standard:

Carcinogenicity:

Mutagenicity:

Teratogenicity:

TLV:

Other Chronic Effects:

Chloroprene

CHEMICAL NAME

CAS Number: 126-99-8

Chemical Name: 2-Chloro-1,3-butadiene

Synonyms: 2-Chlorobutadiene-1,3
 β -Chloroprene
Chloroprene

**Molecular
Formula:** C_4H_5Cl

Structure:
$$\begin{array}{c} \text{Cl} \\ | \\ \text{CH}_2 = \text{C} - \text{CH} = \text{CH}_2 \end{array}$$

Chemical Properties

Molecular Weight: 88.54

Physical State: Liquid

Vapor Pressure: 215.4 mm at 25°C

Vapor Density:

Boiling Point: 59.4°C

Melting Point:

Density: 0.9583 at 20°C/4°C

Solubility: Slightly <10.0 gm/l (H_2O)

**Octanol/Water Parti-
tion Coefficient:**

Environmental Persistence

BOD:

Atmospheric Reactivity:

Activity toward RO_2 $t_{1/2} = 2.2$ yrs.

O_3 $t_{1/2} = 2$ hrs. producing 2-chloroacrolein

OH $t_{1/2} = 21$ hrs.

Photochemical degradation: reacts with reducing materials

Safety Hazard: Disaster hazard—dangerous chlorides

Chloroprene

PRODUCTION DATA

Annual U.S.**Production:** 396 x 10⁶ lbs.**Consumption:****Fraction of****Dispersion:** 0.01**Fraction of Pro-****duction Lost:** 0.015**Release Rate (million lbs/yr):** 6.02

TOXICITY DATA

Acute Toxicity**LD_{Lo}****Dosage**

1600 mg/kg

500 mg/kg

1000 mg/kg

300 mg/kg

96 mg/kg

LC_{Lo}600 mg/m³ (8 hrs)2500 mg/m³ (8 hrs)**Animal**

rat

rat

mouse

cat

rabbit

mouse

cat

Route

oral

subcutaneous

subcutaneous

subcutaneous

intravenous

inhalation

inhalation

Non-lethal Acute Effects: 80 ppm

Central nervous system depression

TC_{Lo} 80 ppm

human

inhalation

Degeneration of liver and kidneys

Dermatitis

Conjunctivitis—corneal necrosis

Anemia

Embryotoxic

Chronic Toxicity**U.S. Occupational Standard:****Carcinogenicity:****Mutagenicity:****TD_{Lo}** 39 mg/m³ (4 hrs per day for 48 days)

rat

inhalation

Teratogenicity:**TLV:****Other Chronic Effects:**

Chloropropham

CHEMICAL NAME

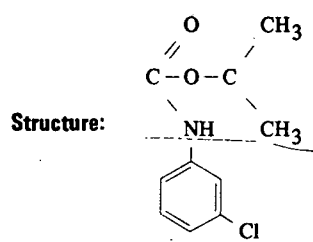
CAS Number: 101-21-3

Chemical Name: m-Chlorocarbanilic acid, isopropyl ester

Synonyms: Isopropyl-N-(3-chlorophenyl) carbanate
Isopropyl-m-chlorocarbanilate
Chloropropham

Molecular

Formula: $C_{10}H_{12}ClNO_2$



Chemical Properties

Molecular Weight: 213.7

Physical State: Solid

Vapor Pressure:

Vapor Density:

Boiling Point: 247°C (decomposes)

Melting Point: 36°C

Density:

Solubility:

Octanol/Water Partition Coefficient:

Environmental Persistence

BOD:

Atmospheric Reactivity:

Safety Hazard: Disaster hazard—dangerous toxic fumes of phosgene

Chloropropham

PRODUCTION DATA

Annual U.S.
Production: 2×10^6 lbs.

Consumption:

Fraction of
Dispersion: 1:0

Fraction of Pro-
duction Lost: 0.015

Release Rate (million lbs/yr): 2.0

TOXICITY DATA

Acute Toxicity
 LD_{50}

Dosage
1200 mg/kg
5000 mg/kg

Animal
rat
rabbit

Route
oral
oral

Chronic Toxicity

U.S. Occupational Standard:

Carcinogenicity:

Neoplasm TD_{Lo}

60 mg/kg

mouse

oral

Mutagenicity:

Teratogenicity:

TLV:

Other Chronic Effects:

m-Chlorotoluene

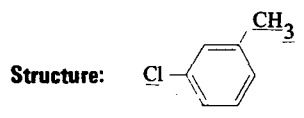
CHEMICAL NAME

CAS Number: 108-41-8

Chemical Name: m-Chlorotoluene
3-Chloro-1-methylbenzene

Synonyms:

**Molecular
Formula:** C₇H₇Cl



Chemical Properties

Molecular Weight: 126.6

Vapor Pressure: 4.17 mm at 25°C

Boiling Point: 162.3°C

Density: 1.0797 at 13.90°C/4°C

**Octanol/Water Parti-
tion Coefficient:** 3.28

Physical State: Liquid

Vapor Density:

Melting Point: -48.0°C

Solubility: Insoluble (H₂O)

Environmental Persistence

BOD:

Atmospheric Reactivity:

Safety Hazard: Fire—slight

m-Chlorotoluene

PRODUCTION DATA

**Annual U.S.
Production:**

Consumption:

**Fraction of
Dispersion:**

**Fraction of Pro-
duction Lost:**

Release Rate (million lbs/yr):

TOXICITY DATA

Acute Toxicity

Dosage

Animal

Route

Chronic Toxicity

U.S. Occupational Standard:

Carcinogenicity: Not tested

Mutagenicity:

Teratogenicity:

TLV:

Other Chronic Effects:

o-Chlorotoluene

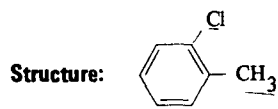
CHEMICAL NAME

CAS Number: 95-49-8

Chemical Name: o-Chlorotoluene

Synonyms: 2-Chloro-1-methylbenzene
 α -Chlorotoluene

**Molecular
Formula:** C_7H_7Cl



Chemical Properties

Molecular Weight: 126.6

Vapor Pressure: 4.11 mm at 25°C

Boiling Point: 159.3°C

Density: 1.1018 at 0°C/4°C

**Octanol/Water Parti-
tion Coefficient:** 3.42

Physical State: Liquid

Vapor Density:

Melting Point: -35.1°C

Solubility: Insoluble (H_2O)

Environmental Persistence

BOD:

Atmospheric Reactivity:

Safety Hazard: Fire—slight

o-Chlorotoluene

PRODUCTION DATA

**Annual U.S.
Production:**

Consumption:

**Fraction of
Dispersion:**

**Fraction of Pro-
duction Lost:**

Release Rate (million lbs/yr):

TOXICITY DATA

Acute Toxicity	Dosage	Animal	Route
LD ₅₀	1231 mg/kg	rat	oral
	1000 mg/kg	rat	subcutaneous
	1624 mg/kg	mouse	oral

Non-lethal Acute Effects:			
TC _{Lo}	16 ppm	human	inhalation

Chronic Toxicity

U.S. Occupational Standard: (air) TWA 1 ppm

Carcinogenicity: Neoplastic effects--TD_{Lo} 2100 mg/kg (51 weeks) rat subcutaneous

Mutagenicity:

Teratogenicity: TLV:

Other Chronic Effects:

p-Chlorotoluene

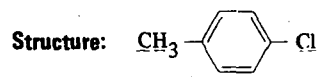
CHEMICAL NAME

CAS Number: 106-43-4

Chemical Name: p-Chlorotoluene

Synonyms: 4-Chloro-l-methyltoluene
p-Tolylchloride

**Molecular
Formula:** C₇H₇Cl



Chemical Properties

Molecular Weight: 126.6

Vapor Pressure: 406 mm at 25°C

Boiling Point: 162.3°C at 760 mm

Density: 1.0697 at 20°C/4°C

**Octanol/Water Parti-
tion Coefficient:** 3.33

Physical State: Liquid

Vapor Density:

Melting Point: 7.5°C

Solubility: Insoluble (H₂O)

Environmental Persistence

BOD:

Atmospheric Reactivity:

Safety Hazard: Fire—slight

p-Chlorotoluene

PRODUCTION DATA

Annual U.S.**Production:** 95 x 10⁶ lbs. (1976 capacity)**Consumption:****Fraction of****Dispersion:** 0.01**Fraction of Pro-****duction Lost:** 0.015**Release Rate (million lbs/yr):**

TOXICITY DATA

Acute Toxicity**Dosage****Animal****Route****Chronic Toxicity****U.S. Occupational Standard:****Carcinogenicity:** Not tested**Mutagenicity:****Teratogenicity:****TLV:****Other Chronic Effects:**

Citric acid

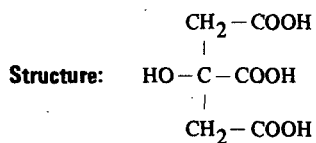
CHEMICAL NAME

CAS Number: 77-92-9

Chemical Name: Citric acid

Synonyms: β -Hydroxytricarballic acid
2-Hydroxy-1,2,3-propane tricarboxylic acid

**Molecular
Formula:** $C_6H_8O_7$



Chemical Properties

Molecular Weight: 210.2

Physical State: Solid

Vapor Pressure:

Vapor Density:

Boiling Point: Decomposes

Melting Point: 153°C

Density: 1.542

Solubility: 1,618 gm/l (H_2O)

**Octanol/Water Parti-
tion Coefficient:** 1.80

Environmental Persistence

BOD: 53% of theoretical
Total theoretical oxygen demand gm/gm = 0.75

Atmospheric Reactivity:

Safety Hazard:

Citric acid

PRODUCTION DATA

Annual U.S.

Production: 163.0×10^6 lbs.

Consumption: 152.4×10^6 lbs.

**Fraction of
Dispersion:** 0.75

**Fraction of Pro-
duction Lost:** 0.03

Release Rate (million lbs/yr): 119.2

TOXICITY DATA

Acute Toxicity
LD₅₀

Dosage
884 mg/kg
961 mg/kg
42 mg/kg
330 mg/kg

Animal
rat
mouse
mouse
rabbit

Route
intraperitoneal
intraperitoneal
intravenous
intravenous

Non-lethal Acute Effects:

Irritant with allergenic effects

Chronic Toxicity

U.S. Occupational Standard:

Carcinogenicity:

Mutagenicity:

Teratogenicity:

TLV:

Other Chronic Effects:

Crag herbicide 1

CHEMICAL NAME

CAS Number: 136-78-7

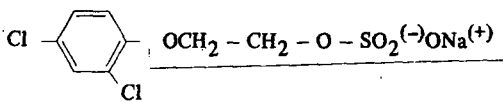
Chemical Name: 2-(2,4-Dichlorophenoxy)ethanol, hydrogensulfate sodium salt

Synonyms: Crag herbicide Sodium 2,4-dichlorophenyl cellosolve sulfate
Sesone
Sodium 2-(2,4-dichlorophenoxy)ethyl sulfate

Molecular

Formula: $C_8H_7Cl_2O_5S \cdot Na$

Structure:



Chemical Properties

Molecular Weight: 310.12

Physical State: Solid

Vapor Pressure:

Vapor Density:

Boiling Point:

Melting Point: 245°C

Density:

Solubility: Soluble (H₂O)

Octanol/Water Partition Coefficient:

Environmental Persistence

BOD:

Atmospheric Reactivity:

Safety Hazard:

Crag herbicide 1

PRODUCTION DATA

**Annual U.S.
Production:**

Consumption:

**Fraction of
Dispersion:**

**Fraction of Pro-
duction Lost:**

Release Rate (million lbs/yr):

TOXICITY DATA

**Acute Toxicity
LD₅₀**

**Dosage
730 mg/kg**

**Animal
rat**

**Route
oral**

Chronic Toxicity

U.S. Occupational Standard: (air) TWA 15 mg/m³

Carcinogenicity:

Mutagenicity:

Teratogenicity:

TLV:

Other Chronic Effects:

m-Cresol

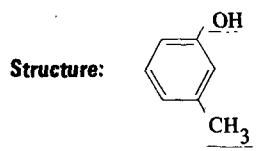
CHEMICAL NAME

CAS Number: 108-39-4

Chemical Name: m-Cresol

Synonyms: m-Methylphenol
m-Hydroxytoluene

Molecular Formula: $\text{CH}_3 \text{C}_6 \text{H}_4 \text{OH}$



Chemical Properties

Molecular Weight: 108.1

Physical State: Liquid

Vapor Pressure: 1 mm at 52.0°C

Vapor Density: 3.72

Boiling Point: 202.8°C

Melting Point: 12°C

Density: 1.034 at 20°C/4°C

Solubility: Slightly soluble (H₂O)

Octanol/Water Partition Coefficient: 2.37

Environmental Persistence

BOD:

Atmospheric Reactivity: Reacts with oxidizing materials

Safety Hazard: Fire—moderate
Explosion—moderate
Disaster hazard—dangerous toxic fumes

PRODUCTION DATA**Annual U.S.****Production:** 31.4×10^6 lbs. (1973) (m and p cresol) **Consumption:****Fraction of
Dispersion:****Fraction of Pro-
duction Lost:** 0.015**Release Rate (million lbs/yr):****TOXICITY DATA**

Acute Toxicity	Dosage	Animal	Route
LD ₅₀	242 mg/kg	rat	oral
	2050 mg/kg	rabbit	skin
LD _{Lo}	180 mg/kg	cat	subcutaneous
	1400 mg/kg	rabbit	oral
	280 mg/kg	rabbit	intravenous

Non-lethal Acute Effects:

CNS disorders
Kidney and liver damage
Irritation of skin and eyes

Chronic Toxicity**U.S. Occupational Standard:** (air) TWA: 5 ppm**Carcinogenicity:****Mutagenicity:****Teratogenicity:****TLV:** 5 ppm**Other Chronic Effects:**

o-Cresol

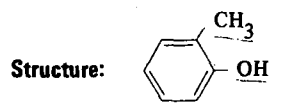
CHEMICAL NAME

CAS Number: 95-48-7

Chemical Name: o-Cresol

Synonyms: o-Cresylic Acid
o-Hydroxytoluene

**Molecular
Formula:** C₇H₈O



Chemical Properties

Molecular Weight: 108.1

Physical State: Solid

Vapor Pressure: 1 mm at 38.2°C

Vapor Density: 3.72

Boiling Point: 190.8°C

Melting Point: 30.9°C

Density: 1.047 at 20°C/4°C

Solubility: Soluble (H₂O) (31 gm/l)

**Octanol/Water Parti-
tion Coefficient:** 3.40

Environmental Persistence

BOD: 95% of theoretical at 6 days
Toxic to sewage organisms

Atmospheric Reactivity: Reacts with oxidizing materials
Less than a day in air } Yields quinones and
Less than 10 days in water } benzenes

Safety Hazard: Fire—moderate
Explosion—slight
Disaster hazard—dangerous—toxic fumes

PRODUCTION DATA**Annual U.S.****Production:** 49.7 x 10⁶ lbs.**Consumption:** 49.7**Fraction of****Dispersion:** 0.3**Fraction of Pro-****duction Lost:** 0.015**Release Rate (million lbs/yr):** 15.6**TOXICITY DATA****Acute Toxicity**LD₅₀**Dosage**

121 mg/kg

1100 mg/kg

344 mg/kg

1380 mg/kg

LD_{Lo}

410 mg/kg

55 mg/kg

940 mg/kg

450 mg/kg

180 mg/kg

360 mg/kg

200 mg/kg

Animal

rat

rat

mouse

rabbit

mouse

cat

rabbit

rabbit

rabbit

guinea pig

frog

Route

oral

skin

oral

skin

subcutaneous

subcutaneous

oral

subcutaneous

intravenous

intraperitoneal

subcutaneous

Chronic Toxicity**U.S. Occupational Standard:** (air) TWA 5 ppm**Carcinogenicity:** 20% benzene solution—papilloma in mouse**Mutagenicity:****Teratogenicity:****TLV:****Other Chronic Effects:** Abnormal mitotic effects.

p-Cresol

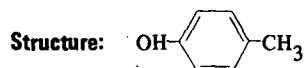
CHEMICAL NAME

CAS Number: 106-44-5

Chemical Name: p-Cresol

Synonyms: 4-Cresol
4-Methylphenol
p-Hydroxytoluene

**Molecular
Formula:** C₇H₈O



Chemical Properties

Molecular Weight: 108.1

Physical State: Solid

Vapor Pressure: 1 mm at 53.0°C

Vapor Density: 3.72

Boiling Point: 201.8°C

Melting Point: 35.26°C

Density: 1.0341 at 20°C/4°C

Solubility: Slightly (H₂O)

**Octanol/Water Parti-
tion Coefficient:** 2.35

Environmental Persistence

BOD:

Atmospheric Reactivity: Reacts with oxidizing materials

Safety Hazard: Fire—moderate
Explosion—moderate
Disaster hazard—dangerous—toxic fumes

PRODUCTION DATA**Annual U.S.****Production:** 31.4×10^6 lbs. (1973) (p and m cresol) **Consumption:****Fraction of
Dispersion:****Fraction of Pro-
duction Lost:** 0.015**Release Rate (million lbs/yr):****TOXICITY DATA**

Acute Toxicity	Dosage	Animal	Route
LD ₅₀	207 mg/kg	rat	oral
	750 mg/kg	rat	skin
	344 mg/kg	mouse	oral
	301 mg/kg	rabbit	skin
LD _{Lo}	150 mg/kg	mouse	subcutaneous
	80 mg/kg	cat	subcutaneous
	620 mg/kg	rabbit	oral
	300 mg/kg	rabbit	subcutaneous
	180 mg/kg	rabbit	intravenous
	100 mg/kg	guinea pig	intraperitoneal
	150 mg/kg	frog	subcutaneous

Non-lethal Acute Effects:

Corrosive to skin and mucous membranes
Damage to kidney, liver, nervous system
Dermatitis

Chronic Toxicity**U.S. Occupational Standard:** (air) TWA 5 ppm**Carcinogenicity:****Mutagenicity:****Teratogenicity:** TD_{Lo} 5500 mg/kg pregnant rat oral TLV: 5 ppm**Other Chronic Effects:**

Crotonaldehyde

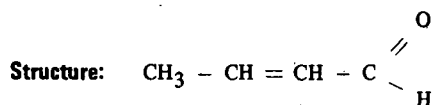
CHEMICAL NAME

CAS Number: 123-73-9

Chemical Name: (E)-Crotonaldehyde

Synonyms: 2-Butenal
Crotonic aldehyde
 β -Methyl acrolein

Molecular
Formula: C_4H_6O



Chemical Properties

Molecular Weight: 70.09

Physical State: Liquid

Vapor Pressure: 10 mm at 25°C

Vapor Density: 2.41

Boiling Point: 104°C

Melting Point: -69°C

Density: 0.853 at 20°C/20°C

Solubility: Soluble (H_2O)

Octanol/Water Parti-
tion Coefficient:

Environmental Persistence

BOD: 27% of theoretical at 10 days

Atmospheric Reactivity: Reacts with oxidizing materials

Safety Hazard:

Crotonaldehyde

PRODUCTION DATA

Annual U.S.

Production:

Consumption:

Fraction of

Dispersion:

Fraction of Pro-

duction Lost:

Release Rate (million lbs/yr):

TOXICITY DATA

Acute Toxicity

LD₅₀

Dosage

300 mg/kg

140 mg/kg

6 mg/kg

160 mg/kg

30

LC₅₀

4000 mg/m³ (30 minutes)

1510 mg/m³ (2 hours)

Animal

rat

rat

mouse

mouse

guinea pig

rat

mouse

Route

oral

subcutaneous

oral

subcutaneous

skin

inhalation

inhalation

Non-lethal Acute Effects:

Irritant- TC_{Lo}

4 ppm

human

inhalation

Dangerous to eyes

Causes corneal burns, irritant to skin

Chronic Toxicity

U.S. Occupational Standard: (air) TWA 2 ppm

Carcinogenicity: Not tested

Mutagenicity:

Teratogenicity:

TLV: 2 ppm

Other Chronic Effects:

Crotonic acid

CHEMICAL NAME

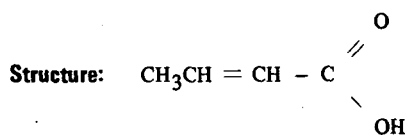
CAS Number: 324-65-0

Chemical Name: Crotonic acid

Synonyms: β -Methyl acrylic acid
Trans-2-butenic acid
Methyl-propenoic acid

Molecular

Formula: $C_4H_6O_2$



Chemical Properties

Molecular Weight: 86.09

Physical State: Solid

Vapor Pressure: 0.19 mm at 20°C

Vapor Density: 2.97

Boiling Point: 185°C

Melting Point: 72°C

Density: 1.018 at 15°C/4°C

Solubility: Soluble (H_2O)

Octanol/Water Partition Coefficient: 0.76

Environmental Persistence

BOD:

Atmospheric Reactivity: Reacts with oxidizing materials

Safety Hazard: Fire-slight

Crotonic acid

PRODUCTION DATA

**Annual U.S.
Production:**

Consumption:

**Fraction of
Dispersion:**

**Fraction of Pro-
duction Lost:**

Release Rate (million lbs/yr):

TOXICITY DATA

Acute Toxicity
LD₅₀

Dosage
1000 mg/kg
600 mg/kg

Animal
rat
guinea pig

Route
oral
skin

Chronic Toxicity

U.S. Occupational Standard:

Carcinogenicity:

Mutagenicity:

Teratogenicity:

TLV:

Other Chronic Effects:

Cumene

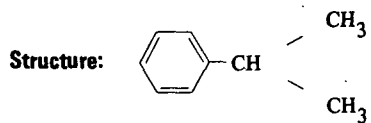
CHEMICAL NAME

CAS Number: 98-82-8

Chemical Name: Cumene

Synonyms: Isopropyl benzene
2-Phenyl propane
Cumol

**Molecular
Formula:** C_9H_{11}



Chemical Properties

Molecular Weight: 120.21

Physical State: Liquid

Vapor Pressure: 6.56 at 25°C

Vapor Density: 4.1

Boiling Point: 152°C

Melting Point: -96°C

Density: 0.864 at 20°C/4°C

Solubility: Insoluble (H₂O)

**Octanol/Water Parti-
tion Coefficient:**

Environmental Persistence

BOD: 70% of theoretical for 20 days at 20°C

Total theoretical oxygen demand gm/gm = 3.5

Atmospheric Reactivity: Reacts with oxidizing materials

Safety Hazard: Fire-moderate

Cumene

PRODUCTION DATA

Annual U.S.**Production:** 2292.9×10^6 lbs.**Consumption:** 2737.2×10^6 lbs.**Fraction of****Dispersion:** 0.01**Fraction of Pro-****duction Lost:** 0.01**Release Rate (million lbs/yr):** 50.3

TOXICITY DATA

Acute ToxicityLD₅₀LC₅₀LC_{Lo}**Dosage**

1400 mg/kg

8000 ppm

2000 ppm

Animal

rat

rat

mouse

Route

oral

inhalation

inhalation

Non-lethal Acute Effects:

Potent narcotic effects

Depressant to CNS

Dermatitis**Chronic Toxicity****U.S. Occupational Standard:** (air): TWA 50 ppm**Carcinogenicity:****Mutagenicity:****Teratogenicity:****TLV:****Other Chronic Effects:**

Cumene hydroperoxide

CHEMICAL NAME

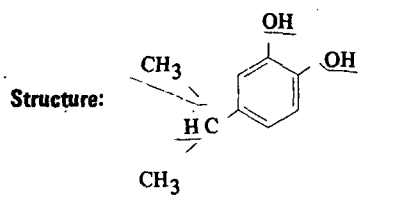
CAS Number: 80-15-9

Chemical Name: α,α -Dimethyl benzyl hydroperoxide

Synonyms: Caruacrol
o-Cresyl-5-isopropyl

2-Hydro-p-cumene

**Molecular
Formula:** $C_9H_{12}O_2$



Chemical Properties

Molecular Weight: 152.21

Physical State: Liquid

Vapor Pressure: 1 mm at 70°C

Vapor Density:

Boiling Point: 153°C

Melting Point:

Density: 1.05

Solubility: Slightly soluble (H₂O)

Octanol/Water Partition Coefficient:

Environmental Persistence

BOD:

Atmospheric Reactivity: Reacts with oxidizing materials

Safety Hazard: Explosion—dangerous

Cumene hydroperoxide

PRODUCTION DATA

Annual U.S.
Production:

Consumption:

Fraction of
Dispersion:

Fraction of Pro-
duction Lost:

Release Rate (million lbs/yr):

TOXICITY DATA

Acute Toxicity	Dosage	Animal	Route
LD ₅₀	382 mg/kg	rat	oral
	95 mg/kg	rat	intraperitoneal
	400 mg/kg	rat	subcutaneous
LD _{Lo}	5000 mg/kg	mouse	oral
	90 mg/kg	mouse	intraperitoneal
LC ₅₀	220 ppm (4 hours)	rat	inhalation
	200 ppm (4 hours)	mouse	inhalation

Chronic Toxicity

U.S. Occupational Standard:

Carcinogenicity:

Neoplastic effects:

TD_{Lo}

304 mg/kg
10 gm/kg (76 weeks)

mouse
mouse

inhalation
subcutaneous

Mutagenicity:

Teratogenicity:

TLV:

Other Chronic Effects:

Skin sensitizer

Cyanoacetic acid

CHEMICAL NAME

CAS Number: 372-09-8

Chemical Name: Cyanoacetic acid

Synonyms: Malonic acid, mononitrile

Molecular

Formula: $C_3H_3NO_2$

Structure: $N \equiv C - CH_2 - COOH$

Chemical Properties

Molecular Weight: 85.06

Physical State: Solid

Vapor Pressure:

Vapor Density:

Boiling Point: 108°C at 15 mm

Melting Point: 66.1 to 66.4°C

Density:

Solubility: Soluble (H_2O)

Octanol/Water Partition Coefficient:

Environmental Persistence

BOD:

Atmospheric Reactivity:

Safety Hazard:

Cyanoacetic acid

PRODUCTION DATA

Annual U.S.

Production: 200 x 10⁶ lbs. (1967)

Consumption:

Fraction of

Dispersion: 0.01

Fraction of Pro-

duction Lost: 0.015

Release Rate (million lbs/yr):

TOXICITY DATA

Acute Toxicity

LD₅₀

Dosage

200 mg/kg

Animal

mouse

Route

intraperitoneal

Very toxic

Chronic Toxicity

U.S. Occupational Standard:

Carcinogenicity:

Mutagenicity:

Teratogenicity:

TLV:

Other Chronic Effects:

Cyanogen chloride

CHEMICAL NAME

CAS Number: 506-77-4

Chemical Name: Cyanogen chloride

Synonyms: Chlorine cyanide
CK

**Molecular
Formula:** CCIN

Structure: $\text{Cl}-\text{C}\equiv\text{N}$

Chemical Properties

Molecular Weight: 61.48

Physical State: Liquid

Vapor Pressure: 1010 mm at 20°C

Vapor Density: 2.1

Boiling Point: 13.1°C

Melting Point: -6.5°C

Density: 1.218 at 4°C/4°C

Solubility: Soluble (H₂O)

**Octanol/Water Parti-
tion Coefficient:**

Environmental Persistence

BOD:

Atmospheric Reactivity:

Safety Hazard:

Cyanogen chloride

PRODUCTION DATA

Annual U.S.
Production:

Consumption:

Fraction of
Dispersion:

Fraction of Pro-
duction Lost:

Release Rate (million lbs/yr):

TOXICITY DATA

Acute Toxicity	Dosage	Animal	Route
LD _{Lo}	39 mg/kg	mouse	subcutaneous
	5 mg/kg	dog	subcutaneous
	20 mg/kg	rabbit	subcutaneous
LD ₅₀	118 ppm (30 minutes)	rat	inhalation
	177 ppm (30 minutes)	mouse	inhalation
	207 ppm (30 minutes)	rabbit	inhalation
LD _{Lo}	207 ppm (30 minutes)	guinea pig	inhalation

Non-lethal Acute Effects:

Irritant to eyes and mucous membranes

Chronic Toxicity

U.S. Occupational Standard:

Carcinogenicity:

Mutagenicity:

Teratogenicity:

TLV:

Other Chronic Effects:

Cyanuric acid

CHEMICAL NAME

CAS Number: 108-80-5

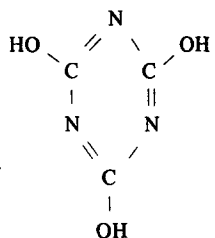
Chemical Name: s-Triazine -2, 4, 6-triol

Synonyms: sym-Triazinetriol Cyanuric acid, dihydrate
Tricyanic acid Tricarbimide

Molecular

Formula: $C_3H_3N_3O_3$

Structure:



Chemical Properties (dihydrate)

Molecular Weight: 165.11

Physical State: solid

Vapor Pressure:

Vapor Density:

Boiling Point: decomposes

Melting Point: 320 to 350°C (decomposes)

Density: 1.768 at 0°C

Solubility: Soluble (H₂O)

Octanol/Water Partition Coefficient:

Environmental Persistence

BOD:

Atmospheric Reactivity:

Safety Hazard: Disaster hazard—cyanides

Cyanuric acid

PRODUCTION DATA

Annual U.S.

Production:

Consumption:

Fraction of

Dispersion:

Fraction of Pro-

duction Lost:

Release Rate (million lbs/yr):

TOXICITY DATA

Acute Toxicity

Highly toxic

Dosage

Animal

Route

Chronic Toxicity

U.S. Occupational Standard:

Carcinogenicity:

Mutagenicity:

Teratogenicity:

TLV:

Other Chronic Effects:

Cyanuric chloride

CHEMICAL NAME

CAS Number: 108-77-0

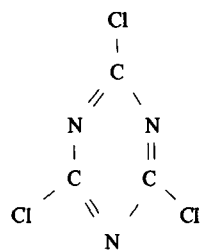
Chemical Name: 2,4,6 Trichloro-s-triazine

Synonyms: Cyanuric chloride

Molecular

Formula: $C_3Cl_3N_3$

Structure:



Chemical Properties

Molecular Weight: 184.42

Vapor Pressure: 2 mm at 70°C

Boiling Point: 190°C

Density: 1.32 at 20°C/4°C

Octanol/Water Partition Coefficient:

Physical State: Solid

Vapor Density: 6.36

Melting Point: 154°C

Solubility: Insoluble (H₂O)

Environmental Persistence

BOD:

Atmospheric Reactivity:

Safety Hazard: Disaster hazard—dangerous—chlorides and nitriles

Cyanuric chloride

PRODUCTION DATA

Annual U.S.
Production:

Consumption:

Fraction of
Dispersion:

Fraction of Pro-
duction Lost:

Release Rate (million lbs/yr):

TOXICITY DATA

Acute Toxicity
LD₅₀

Dosage
485 mg/kg

Animal
rat

Route
oral

Non-lethal Acute Effects:

Irritation of mucous membranes
Disturbed heart rhythm

Chronic Toxicity

U.S. Occupational Standard:

Carcinogenicity: Neoplastic effects

TD_{Lo}

22 gm/kg (82 weeks)

rat

oral

750 mg/kg (15 weeks)

rat

subcutaneous

Mutagenicity:

Teratogenicity:

TLV:

Other Chronic Effects:

Cyclohexane

CHEMICAL NAME

CAS Number: 110-82-7

Chemical Name: Cyclohexane

Synonyms: Hexahydrobenzene
Hexanaphene
Hexamethylene

**Molecular
Formula:** C_6H_{12}

Structure:



Chemical Properties

Molecular Weight: 84.16

Physical State: Liquid

Vapor Pressure: 98.14 at 25°C

Vapor Density: 2.90

Boiling Point: 80.7°C

Melting Point: 6.3°C

Density: 0.77855 at 20°C/4°C

Solubility: Insoluble (H₂O) (<1 gm/l)

**Octanol/Water Parti-
tion Coefficient:**

Environmental Persistence

BOD:

Atmospheric Reactivity: Reacts with oxidizing materials

Safety Hazard: Fire—dangerous
Explosion—moderate

Cyclohexane

PRODUCTION DATA

Annual U.S. Production: 416×10^6 lbs. (1976 capacity) **Consumption:**

Fraction of Dispersion: 0.10 **Fraction of Production Lost:** 0.015

Release Rate (million lbs/yr): 94.2 (1972)

TOXICITY DATA

Acute Toxicity	Dosage	Animal	Route
LD ₅₀	1297 mg/kg	mouse	oral
	5500 mg/kg	rabbit	oral
LD _{Lo}	77 mg/kg	rabbit	intravenous

Chronic Toxicity

U.S. Occupational Standard: (air) TWA 300 ppm

Carcinogenicity:

Mutagenicity:

Teratogenicity:

TLV: 300 ppm

Other Chronic Effects: Skin irritation
May be an asphyxiant

Cyclohexanol

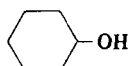
CHEMICAL NAME

CAS Number: 108-93-0

Chemical Name: Cyclohexanol

Synonyms: Hexahydrophenol
Hexalin

**Molecular
Formula:** $C_6H_{12}O$

Structure: 

Chemical Properties

Molecular Weight: 100.17

Physical State: Liquid

Vapor Pressure: 1.7 mm at 25°C

Vapor Density: 3.45

Boiling Point: 161.5°C

Melting Point: 23°C

Density: 0.9449 at 25°C/4°C

Solubility: Soluble (H_2O)

**Octanol/Water Parti-
tion Coefficient:**

Environmental Persistence

BOD:

Atmospheric Reactivity: Reacts with oxidizing materials

Safety Hazard: Fire-moderate

Cyclohexanol

PRODUCTION DATA

Annual U.S.**Production:** 760 x 10⁶ lbs.**Consumption:****Fraction of****Dispersion:** 0.01**Fraction of Pro-****duction Lost:** 0.007**Release Rate (million lbs/yr):**

TOXICITY DATA

Acute ToxicityLD₅₀LD_{Lo}**Dosage**

2060 mg/kg

1300 mg/kg

800 mg/kg

2200 mg/kg

1420 mg/kg

Animal

rat

dog

cat

rabbit

rabbit

Route

oral

oral

subcutaneous

oral

intraperitoneal

Non-lethal Acute Effects:Mucous membrane effect: TC_{Lo} 100 ppm

Conjunctivitis

Corneal necrosis

Irritation of skin and respiratory tract

Chronic Toxicity**U.S. Occupational Standard:** (air) TWA 50 ppm**Carcinogenicity:** Not tested**Mutagenicity:****Teratogenicity:****TLV:** 50 ppm**Other Chronic Effects:** Damage to kidneys, liver, blood vessels

Cyclohexanone

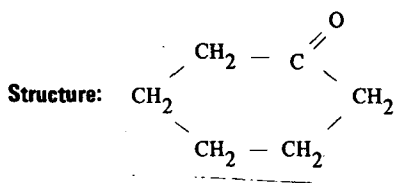
CHEMICAL NAME

CAS Number: 108-94-1

Chemical Name: Cyclohexanone

Synonyms: Ketohexamethylene Cyclohexylketone
Pimelic ketone
Hexanon

Molecular Formula: $C_6H_{10}O$



Chemical Properties

Molecular Weight: 98.14

Physical State: Liquid

Vapor Pressure: 4.77 mm at 25°C

Vapor Density: 3.4

Boiling Point: 155.6°C

Melting Point: -47°C

Density: 0.9478 at 20°C/4°C

Solubility: 50 gm/l (H₂O)

Octanol/Water Partition Coefficient: 0.01

Environmental Persistence

BOD:

Atmospheric Reactivity: Reacts with oxidizing materials

Safety Hazard: Fire—moderate
Explosion—slight

Cyclohexanone

PRODUCTION DATA

Annual U.S.
Production: 784.4 x 10⁶ lbs. Consumption: 784.4

Fraction of
Dispersion: 0.05 Fraction of Pro-
duction Lost: 0.015

Release Rate (million lbs/yr): 51.0

TOXICITY DATA

Acute Toxicity	Dosage	Animal	Route
LD ₅₀	1620 mg/kg	rat	oral
	1350 mg/kg	mouse	intraperitoneal
LD _{Lo}	1000 mg/kg	rabbit	skin
	1300 mg/kg	mouse	oral
	630 mg/kg	dog	intravenous
	1600 mg/kg	rabbit	oral
LC _{Lo}	2000 ppm (4 hours)	rat	inhalation

Non-lethal Acute Effects:

Irritant: eyes and throat

TC _{Lo}	50 ppm	human	inhalation
------------------	--------	-------	------------

Chronic Toxicity

U.S. Occupational Standard: (air) TWA 50 ppm

Carcinogenicity:

Mutagenicity:

Teratogenicity:

TLV: 50 ppm

Other Chronic Effects:

Cyclohexene

CHEMICAL NAME

CAS Number: 110-83-8

Chemical Name: Cyclohexène

Synonyms: 1,2,3,4-Tetrahydrobenzene

Molecular

Formula: C₆H₁₀

Structure:



Chemical Properties

Molecular Weight: 82.14

Physical State: Liquid

Vapor Pressure: 160 mm at 38°C

Vapor Density: 2.8

Boiling Point: 83°C at 760°C

Melting Point: -103.7

Density: 0.8102 at 20°C/4°C

Solubility: Insoluble (H₂O)

Octanol/Water Partition Coefficient:

Environmental Persistence

BOD:

Atmospheric Reactivity: Reacts with oxidizing materials

Safety Hazard: Fire-dangerous

Cyclohexene

PRODUCTION DATA

Annual U.S.
Production: 2298.4 x 10⁶ lbs. Consumption: 1779.4

Fraction of
Dispersion: 0.04 Fraction of Pro-
duction Lost: 0.01

Release Rate (million lbs/yr): 94.2

TOXICITY DATA

Acute Toxicity	Dosage	Animal	Route
----------------	--------	--------	-------

Chronic Toxicity

U.S. Occupational Standard: (air) TWA 300 ppm

Carcinogenicity:

Mutagenicity:

Teratogenicity:

TLV:

Other Chronic Effects:

Cyclohexylamine

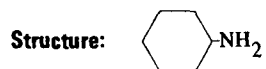
CHEMICAL NAME

CAS Number: 108-91-8

Chemical Name: Cyclohexylamine

Synonyms: Hexahydroaniline
Aminocyclohexane

**Molecular
Formula:** $C_6H_{13}N$



Chemical Properties

Molecular Weight: 99.2

Physical State: Liquid

Vapor Pressure:

Vapor Density: 3.42

Boiling Point: 134.5°C

Melting Point: -18°C

Density: 0.865 at 25°C/25°C

Solubility: Soluble (H_2O)

**Octanol/Water Parti-
tion Coefficient:**

Environmental Persistence

BOD:

Atmospheric Reactivity: Reacts with oxidizing materials

Safety Hazard: Fire—moderate
Disaster hazard—dangerous—toxic fumes

Cyclohexylamine

PRODUCTION DATA

Annual U.S.
Production: 12×10^6 lbs. (1967)

Consumption:

Fraction of
Dispersion: 0.01

Fraction of Pro-
duction Lost: 0.015

Release Rate (million lbs/yr):

TOXICITY DATA

Acute Toxicity

LD₅₀

Dosage

710 mg/kg

200 mg/kg

300 mg/kg

320 mg/kg

500 mg/kg

8000 ppm (4 hours)

Animal

rat

rat

mouse

rabbit

rabbit

rat

Route

oral

intraperitoneal

intraperitoneal

skin

parenteral

inhalation

LD_{Lo}

LC_{Lo}

Chronic Toxicity

U.S. Occupational Standard:

Carcinogenicity:

Mutagenicity:

Teratogenicity:

TLV: 10 ppm

Other Chronic Effects:

Dermatitis

Convulsions

CNS depression

Cyclopentadiene

CHEMICAL NAME

CAS Number: 542-92-7

Chemical Name: 1,3-Cyclopentadiene

Synonyms:

Molecular

Formula: C₅H₆

Structure:



Chemical Properties

Molecular Weight: 66.11

Vapor Pressure: 10 mm at 25°C

Boiling Point: 40.0°C at 760 mm

Density: 0.80475 at 19°C/4°C

Octanol/Water Partition Coefficient:

Physical State:

Vapor Density:

Melting Point: -97.2°C

Solubility: Insoluble (H₂O)

Environmental Persistence

BOD:

Atmospheric Reactivity: Reacts with oxidizing materials

Safety Hazard: Fire—moderate
Explosion—moderate

Cyclopentadiene

PRODUCTION DATA

Annual U.S. Production: 100×10^6 lbs. (Total capacity) **Consumption:**

Fraction of Dispersion: 0.01 **Fraction of Production Lost:** 0.015

Release Rate (million lbs/yr):

TOXICITY DATA

Acute Toxicity	Dosage	Animal	Route
TD _{Lo}	250 ppm	human	inhalation

Chronic Toxicity

U.S. Occupational Standard: (air) TWA 75 ppm

Carcinogenicity: Not tested

Mutagenicity:

Teratogenicity:

TLV:

Other Chronic Effects: Abdominal pain
Jaundice
Anemia

Cyprex

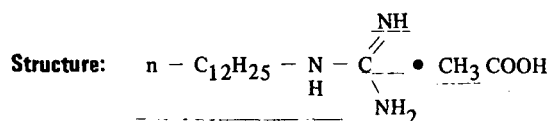
CHEMICAL NAME

CAS Number: 2439-10-3

Chemical Name: Dodecylguanidine, monoacetate

Synonyms: Cyprex
Dodine

Molecular Formula: $C_{13}H_{29}N_3.C_2H_4O_2$



Chemical Properties

Molecular Weight: 287.36

Physical State: Solid

Vapor Pressure:

Vapor Density:

Boiling Point:

Melting Point: 136°C

Density:

Solubility: Soluble (hot H₂O)

Octanol/Water Partition Coefficient:

Environmental Persistence

BOD:

Atmospheric Reactivity:

Safety Hazard:

PRODUCTION DATA

Annual U.S. Production:	2.0 x 10 ⁶ lbs.	Consumption:	2.0 x 10 ⁶ lbs.
Fraction of Dispersion:	1.0	Fraction of Production Lost:	0.01
Release Rate (million lbs/yr): 2.0			

TOXICITY DATA

Acute Toxicity	Dosage	Animal	Route
LD ₅₀	566 mg/kg	rat	oral
	1000 mg/kg	rat	unknown

Chronic Toxicity

U.S. Occupational Standard:

Carcinogenicity:

Mutagenicity:

Teratogenicity:

TLV:

Other Chronic Effects:

Strong solutions cause severe irritation of skin and eyes

TECHNICAL REPORT DATA <i>(Please read Instructions on the reverse before completing)</i>		
1. REPORT NO. EPA-450/3-77-008b	2.	3. RECIPIENT'S ACCESSION NO.
4. TITLE AND SUMMARY Preliminary Scoring of Organic Air Pollutants: Chemistry, Production and Toxicity of Selected Synthetic Organic Chemicals (Chemicals A-C) Appendix I	5. REPORT DATE October 1976	
	6. PERFORMING ORGANIZATION CODE	
7. AUTHOR(S) J. Dorigan, B. Fuller, R. Duffy	8. PERFORMING ORGANIZATION REPORT NO.	
9. PERFORMING ORGANIZATION NAME AND ADDRESS The Mitre Corporation Metrek Division Westgate Research Park McLean, Virginia 22101	10. PROGRAM ELEMENT NO.	
	11. CONTRACT/GRANT NO. 68-02-1495	
12. SPONSORING AGENCY NAME AND ADDRESS EPA, Office of Air Quality Planning and Standards Strategies and Air Standards Division Research Triangle Park, N.C. 27711	13. TYPE OF REPORT AND PERIOD COVERED	
	14. SPONSORING AGENCY CODE	
15. SUPPLEMENTARY NOTES		
16. ABSTRACT This is the first of a series of four appendices to the report "Scoring of Organic Air Pollutants." The entire appendix contains a compilation of available data on chemical structure and properties, environmental persistence, production, and toxicity for 637 synthetic organic chemicals.		
17. KEY WORDS AND DOCUMENT ANALYSIS		
a. DESCRIPTORS	b. IDENTIFIERS/OPEN ENDED TERMS	c. COSATI Field/Group
organic chemical structure organic chemical properties organic chemical production organic chemical toxicity		
18. DISTRIBUTION STATEMENT unlimited	19. SECURITY CLASS (This Report) unclassified	21. NO. OF PAGES 338
	20. SECURITY CLASS (This page) unclassified	22. PRICE