

# Integrated Risk Information System

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# Methyl parathion (CASRN 298-00-0)

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Reference Dose for Chronic Oral Exposure (RfD)



#### 0174

Methyl parathion; CASRN 298-00-0

Health assessment information on a chemical substance is included in IRIS only after a comprehensive review of chronic toxicity data by U.S. EPA health scientists from several Program Offices and the Office of Research and Development. The summaries presented in Sections I and II represent a consensus reached in the review process. Background information and explanations of the methods used to derive the values given in IRIS are provided in the Background Documents.

STATUS OF DATA FOR Methyl parathion

#### File First On-Line 03/31/1987

Category (section)	Status	Last Revised
Oral RfD Assessment (I.A.)	on-line	03/01/1991
Inhalation RfC Assessment (I.B.)	no data	
Carcinogenicity Assessment (II.)	no data	

## \_I. Chronic Health Hazard Assessments for Noncarcinogenic Effects

### \_I.A. Reference Dose for Chronic Oral Exposure (RfD)

Substance Name — Methyl parathion CASRN — 298-00-0 Last Revised — 03/01/1991

The oral Reference Dose (RfD) is based on the assumption that thresholds exist for certain toxic effects such as cellular necrosis. It is expressed in units of mg/kg-day. In general, the RfD is an estimate (with uncertainty spanning perhaps an order of magnitude) of a daily exposure to the human population (including sensitive subgroups) that is likely to be without an appreciable risk of deleterious effects during a lifetime. Please refer to the Background

Document for an elaboration of these concepts. RfDs can also be derived for the noncarcinogenic health effects of substances that are also carcinogens. Therefore, it is essential to refer to other sources of information concerning the carcinogenicity of this substance. If the U.S. EPA has evaluated this substance for potential human carcinogenicity, a summary of that evaluation will be contained in Section II of this file.

### \_\_I.A.1. Oral RfD Summary

Critical Effect	Experimental Doses*	UF	MF	RfD
RBC, ChE inhibition; reduced hemoglobin, hematocrit and RBCs	NOEL: 0.5 ppm (0.025 mg/kg/day)	100	1	2.5E-4 mg/kg/day
2-Year Rat Feeding Study	LEL: 5.0 ppm (0.25 mg/kg/day)			
Monsanto Co., 1984				

<sup>\*</sup>Conversion Factors -- 1 ppm = 0.05 mg/kg/day (assumed rat food consumption)

### \_\_I.A.2. Principal and Supporting Studies (Oral RfD)

Monsanto Company. 1984. MRID No. 000139023, 00143965, 00145507. Available from EPA. Write to FOI, EPA, Washington, DC 20460.

Sixty rats/sex/group were fed diet containing methyl parathion at concentrations of 0.5, 5 or 50 ppm for 2 years. This study was classified as supplementary because a NOEL for neurologic changes was not adequately defined. Sciatic nerve preparations from 1 of 5 males in the low-dose group and 1 of 5 in the mid-dose group reportedly showed moderate degenerative changes. However, based on effects observed in hematological parameters, a NOEL of 0.5 ppm can be established. Hemoglobin, hematocrit, and RBCs were slightly reduced in mid- and high-dose males, and moderately reduced in high- dose females.

#### \_\_I.A.3. Uncertainty and Modifying Factors (Oral RfD)

UF — Based on a chronic exposure study, an uncertainty factor of 100 was used to account for inter- and intraspecies differences in the extrapolation of toxicity to humans.

MF — None

#### \_\_I.A.4. Additional Studies/Comments (Oral RfD)

In a subchronic study with methyl parathion in humans (Rider et al., 1971), RBC cholinesterase depression was reported, with a NOEL of approximately 0.3 mg/kg/day. Using a UF of 100 to adjust for chronic exposure and intraspecies sensitivity, an RfD based on this study would be 0.003 mg/kg/day. Adequate supporting data for human studies are not available. Nevertheless, even anecdotal data directly relating to human exposure should not be dismissed. Therefore, an RfD based on animal studies should not exceed 0.003 mg/kg/day unless additional data for humans can be found to support such a determination.

Data Considered for Establishing the RfD:

- 1) 2-Year Feeding rat: Principal study see previous description; core grade supplementary
- 2) Teratology rat: Embryo/fetotoxicity, developmental NOEL=I0 mg/kg/day (i.p., single dose on day 12 of gestation); LEL=I5 mg/kg/day (mortality, growth retardation retardation, delayed ossification); core grade supplementary (Stauffer Chemical Co., 1967a)
- 3) Teratology mouse: Teratogenic NOEL=20 mg/kg/day; LEL=60\mg/kg/day (cleft palate); (i.p., single dose on day IO of gestation) (Stauffer Chemical Co., 1967b)
- 4) 3-Month Feeding rat: NOEL=2.5 ppm; LEL=25 ppm [reduced hematocrit and ChE inhibition (brain, plasma and RBC), increased serum alkaline phosphatase urine specific gravity]; core grade guideline (Monsanto, 1980a)
- 5) 3-Month Feeding mouse: NOEL=none; LEL=10 ppm (LDT) [decreased testicular weight (no abnormal histopathology), ChE not determined]; core grade minimum (Monsanto, 1980b)
- 6) 3-Month Feeding dog: NOEL=0.3 mg/kg/day; LEL=I mg/kg/day (plasma, RBC ChE inhibition); core grade guideline (Monsanto, 1978)
- 7) Teratology rabbit: NOEL=3 mg/kg/day (HDT); LEL=none; core grade supplementary (A/S Cheminova, 1984)
- 8) Teratology rat: NOEL=0.3 mg/kg/day (gavage); LEL=I mg/kg/day; core grade supplementary (A/S Cheminova, 1977)
- 9) 3-Generation Reproduction rat: NOEL=I0 ppm; LEL=30 ppm (stillbirths, weanling mortality, reduced weanling body weight); core grade minimum (Natural Agricultural Chem. Assoc., 1964)
- 10) 2-Generation Reproduction rat: Reproductive NOEL=25 ppm (HDT); Maternal NOEL=5 ppm; LEL=25 ppm (reduced body weight); core grade minimum (Monsanto, 1982)

### Other Data Reviewed:

- 1) 2-Year Feeding rat (Wistar): NOEL=2 ppm; LEL=10 ppm; (ChE inhibition); body weight gain depression at 50 ppm; (incomplete toxicity data); core grade supplementary (A/S Cheminova, 1981)
- 2) 30-Day Studies human
- a) Rider, J., J. Swader, E. Puletti. 1971. Anticholinesterase toxicity studies with methyl parathion, guthion and phosdrin in human subjects. Fed. Proc. 30(2): 443 [Abstract]. RBC ChE inhibition at 28 and 30 mg/kg/day. Summary data only. No basis for validation (no core grade).
- b) Rider, J., J. Swader, E. Puletti. 1970. Methyl parathion and guthion anticholinesterase effects in human subjects. Fed. Proc. 29(2): 347 [Abstract]. Summary data only. No basis for validation (no core grade).

Data Gap(s): Chronic Dog Feeding Study; Chronic Rat Feeding Study I.A.5. Confidence in the Oral RfD Study — Medium Database — Medium RfD — Medium The principal study was well conducted with a good number of animals and doses, but confidence is considered medium because it is incomplete in regard to neurological evaluation. Confidence in the database is medium because although it is extensive it fails to confirm the possible neurological problems. Medium confidence in the RfD follows. \_\_I.A.6. EPA Documentation and Review of the Oral RfD Pesticide Registration Standard, May 1986 (Draft) Agency Work Group Review — 05/14/1986, 12/09/1986 Verification Date — 12/09/1986 \_\_I.A.7. EPA Contacts (Oral RfD) Please contact the IRIS Hotline for all questions concerning this assessment or IRIS, in general, at (202)566-1676 (phone), (202)566-1749 (FAX) or hotline.iris@epa.gov (internet address). \_I.B. Reference Concentration for Chronic Inhalation Exposure (RfC) Substance Name — Methyl parathion CASRN — 298-00-0 Not available at this time. \_II. Carcinogenicity Assessment for Lifetime Exposure Substance Name — Methyl parathion CASRN — 298-00-0

This substance/agent has not undergone a complete evaluation and determination under US EPA's IRIS program for evidence of human carcinogenic potential.

\_III. [reserved] \_IV. [reserved] \_V. [reserved]

### \_VI. Bibliography

Substance Name — Methyl parathion CASRN — 298-00-0
Last Revised — 03/01/1991

#### \_VI.A. Oral RfD References

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Stauffer Chemical Company. 1967b. MRID No. 00127241. Available from EPA. Write to FOI, EPA, Washington, DC 20460.

#### \_VI.B. Inhalation RfC References

None

### \_VI.C. Carcinogenicity Assessment References

None

# \_VII. Revision History

Substance Name — Methyl parathion CASRN — 298-00-0

Date	Section	Description
03/01/1991	I.A.4.	Citations added
03/01/1991	III.A.	Health Advisory on-line
03/01/1991	VI.	Bibliography on-line
01/01/1992	IV.	Regulatory actions updated
04/01/1997	III., IV., V.	Drinking Water Health Advisories, EPA Regulatory Actions, and Supplementary Data were removed from IRIS on or before April 1997. IRIS users were directed to the appropriate EPA Program Offices for this information.
02/22/2001	I.A., II.	This chemical is being reassessed under the IRIS Program.
02/09/2004	I.A., II.	This chemical is no longer being reassessed under the IRIS Program. See Federal Register February 9, 2004 (Volume 69, Number 26).

# \_VIII. Synonyms

Substance Name — Methyl parathion CASRN — 298-00-0 Last Revised — 03/31/1987

> 298-00-0 8056 HC

Azofos

Azophos

BAY 11405

Bladan-M

Dalf

Dimethylfenitrothion

Dimethyl 4-Nitrophenyl Phosphorothionate

Dimethyl p-Nitrophenyl Phosphorothionate

Dimethyl p-Nitrophenyl Thiophosphate

Dimethyl Parathion

E 601

ENT 17,292

Folidol M

Folidol M-40

Gearphos

Meptox

Metacid 50

Metacide

Metafos http://www.epa.gov/iris/subst/0174.htm Metaphos Last updated on Thursday, January 19, 2012 Methyl-E 605 Methyl Parathion Methylthiophos Metron M-Parathion NCI-C02971 Nitrox Nitrox 80 Oleovofotox O,O-Dimethyl O-(p-Nitrophenyl) Phosphorothioate O,O-Dimethyl O-(p-Nitrophenyl) Thionophosphate O,O-Dimethyl O-(p-Nitrophenyl) Thiophosphate Partron M Penncap M Penncap MLS Phosphorothioic Acid, O,O-Dimethyl O-(4-Nitrophenyl) Ester Phosphorothioic Acid, O,O-Dimethyl O-(p-Nitrophenyl) Ester Quinophos Sinafid M-48 Thiophenit Vofatox Wofatox

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