

Malathion

Other names:

(dl) malathion
1,2-Di(ethoxycarbonyl)ethyl O,O-dimethyl phosphordithioate
1,2-Di(ethoxycarbonyl)ethyl O,O-dimethyl phosphorodithioate
AC 26691
American Cyanamid 4,049
Butanedioic acid, [(dimethoxyphosphinothioyl)thio]-, diethyl ester
Calmathion
Carbethoxy malathion
Carbetovur
Carbetox
Carbofos
Carbophos
Celthion
Chemathion
Cimexan
Cleensheen
Compound 4049
Cythion
Derbac-M
Detmol MA
Dicarboethoxyethyl O,O-dimethyl phosphorodithioate
Diethyl (dimethoxyphosphinothioylthio) butanedioate
Diethyl (dimethoxyphosphinothioylthio)succinate
Diethyl 2-(dimethoxyphosphinothioylthio)succinate
Diethyl [(dimethoxyphosphinothioyl)thio]butanedioate
Diethyl mercaptosuccinate S-ester with O,O-dimethyl phosphorodithioate
Diethyl mercaptosuccinate, O,O-dimethyl dithiophosphate, S-ester
Diethyl mercaptosuccinate, O,O-dimethyl phosphorodithioate
Diethyl mercaptosuccinate, O,O-dimethyl thiophosphate
Diethyl mercaptosuccinic acid O,O-dimethyl phosphorodithioate
Diethyl mercaptosuccinic acid, S-ester of O,O-dimethyl phosphorodithioate
Dithiophosphate de O,O-dimethyle et de S-(1,2-dicarboethoxyethyle)
EL 4049
ENT 17,034
Emmatos
Emmatos extra
Ethiolacar
Etiol
Experimental insecticide 4049
Extermathion

Formal
Forthion
Fosfotion
Fosfotion
Fosfotion 550
Four thousand forty-nine
Fyfanon
Gammazine
Hilthion
Hilthion 25WDP
Insecticide No. 4049
Karbofos
Kop-thion
Kypfos
Latka 4049
MLT
Malacide
Malafor
Malagran
Malakill
Malamar
Malamar 50
Malaphele
Malaphos
Malasol
Malaspray
Malataf
Malathion E50
Malathion LV Concentrate
Malathion ULV
Malathione
Malathiozoo
Malathon
Malathyl
Malathyne
Malation
Malatol
Malatox
Maldison
Malmed
Malphos
Maltox
Maltox MLT

Mercaptosuccinic acid diethyl ester
 Mercaptosuccinic acid diethyl ester, S-ester with O,O-dimethyl phosphorothioate
 Mercaptothion
 Mercaptotion
 Moscarda
 NA 2783
 NCI-C00215
 O,O-Dimethyl S-(1,2-dicarbethoxyethyl) dithiophosphate
 O,O-Dimethyl S-(1,2-dicarbethoxyethyl) phosphorodithioate
 O,O-Dimethyl S-(1,2-dicarbethoxyethyl) thiothionophosphate
 O,O-Dimethyl S-1,2-di(ethoxycarbonyl)ethyl phosphorodithioate
 O,O-Dimethyl S-1,2-di(ethoxycarbonyl)ethyl phosphorothioate
 O,O-Dimethyl dithiophosphate diethylmercaptosuccinate
 O,O-Dimethyl-S-(1,2-di(ethoxycarbonyl)ethyl) phosphorodithioate
 O,O-Dimethyl-S-1,2-dikarbetoxyethyliditiofosfat
 O,O-Dwumetylo-S-1,2-bis(karboetoksyetylo)-dwutiofosforan
 OMS 1
 Oleophosphothion
 Organoderm
 Ortho Malathion
 Ovide
 Phosphothion
 Prioderm
 Radotion
 S-(1,2-Bis(aethoxy-carbonyl)-aethyl)-O,O-dimethyl-dithiophosphat
 S-(1,2-Bis(ethoxy-carbonyl)-ethyl)-O,O-dimethyl-dithiofosfaat
 S-(1,2-Bis(etossi-carbonil)-etil)-O,O-dimetil-ditiofosfato
 S-(1,2-Di(ethoxycarbonyl)ethyl) dimethyl phosphorothiolothionate
 S-(1,2-Dicarbethoxyethyl) O,O-dimethyl phosphorodithioate
 S-(1,2-Dicarbethoxyethyl) O,O-dimethyldithiophosphate
 S-1,2-Bis(ethoxycarbonyl)ethyl-O,O-dimethyl thiophosphate
 S-[1,2-Bis(carbethoxy)ethyl] O,O-dimethyl dithiophosphate
 S-[1,2-Bis(ethoxycarbonyl)ethyl] O,O-dimethyl phosphorodithioate
 SF 60
 Sadofos
 Sadofos 30
 Sadophos
 Siptox I
 Staeubol-Puder
 Succinic acid, mercapto-, diethyl ester, S-ester with O,O-dimethyl phosphorodithioate
 Suleo M
 Sumitox
 TAK

	TM-4049
	Taskil
	Vegfru malatox
	Vetiol
	Zithiol
	[(Dimethoxyphosphinothioyl)thio]butanedioic acid, diethyl ester
	diethyl 2-dimethoxyphosphinothioylsulfanylbutanedioate
Inchi:	InChI=1S/C10H19O6PS2/c1-5-15-9(11)7-8(10(12)16-6-2)19-17(18,13-3)14-4/h8H,5-7H2
InchiKey:	JXSJBGJIGXNWCI-UHFFFAOYSA-N
Formula:	C10H19O6PS2
SMILES:	CCOC(=O)CC(SP(=S)(OC)OC)C(=O)OCC
Mol. weight [g/mol]:	330.36
CAS:	121-75-5

Physical Properties

Property code	Value	Unit	Source
hvap	71.10	kJ/mol	NIST Webbook
log10ws	-3.36		Aqueous Solubility Prediction Method
log10ws	-3.37		Estimated Solubility Method
logp	2.122		Crippen Method
mcvol	231.540	ml/mol	McGowan Method
rinpol	1938.00		NIST Webbook
rinpol	1922.00		NIST Webbook
rinpol	1920.00		NIST Webbook
rinpol	1962.00		NIST Webbook
rinpol	1917.00		NIST Webbook
rinpol	1959.00		NIST Webbook
rinpol	1965.00		NIST Webbook
rinpol	1919.00		NIST Webbook
rinpol	1946.00		NIST Webbook
rinpol	1919.00		NIST Webbook
rinpol	1930.00		NIST Webbook
rinpol	1930.00		NIST Webbook
rinpol	1920.00		NIST Webbook
rinpol	1963.00		NIST Webbook
rinpol	1965.00		NIST Webbook
rinpol	1962.00		NIST Webbook
rinpol	1930.00		NIST Webbook
rinpol	1965.00		NIST Webbook

rinpol	1970.00		NIST Webbook
rinpol	1975.00		NIST Webbook
rinpol	1986.00		NIST Webbook
rinpol	1986.00		NIST Webbook
rinpol	1969.00		NIST Webbook
rinpol	1913.00		NIST Webbook
rinpol	1917.00		NIST Webbook
rinpol	1917.00		NIST Webbook
rinpol	1926.00		NIST Webbook
rinpol	1930.00		NIST Webbook
rinpol	1934.00		NIST Webbook
rinpol	1931.00		NIST Webbook
rinpol	1928.00		NIST Webbook
rinpol	1919.00		NIST Webbook
ripol	2872.00		NIST Webbook
tf	276.25 ± 0.25	K	NIST Webbook
tf	275.95	K	Aqueous Solubility Prediction Method
tf	275.95 ± 0.35	K	NIST Webbook
tf	273.95 ± 0.50	K	NIST Webbook

Correlations

Information	Value
Property code	pvap
Equation	$\ln(P_{vp}) = A + B/(T + C)$
Coeff. A	1.77036e+01
Coeff. B	-8.55779e+03
Temperature range (K), min.	491.37
Temperature range (K), max.	690.58

Sources

McGowan Method:	http://link.springer.com/article/10.1007/BF02311772
NIST Webbook:	http://webbook.nist.gov/cgi/cbook.cgi?ID=C121755&Units=SI
The Yaws Handbook of Vapor Pressure:	https://www.sciencedirect.com/book/9780128029992/the-yaws-handbook-of-vapor-pressure
Crippen Method:	http://pubs.acs.org/doi/abs/10.1021/ci990307I
Aqueous Solubility Prediction Method:	http://onschallenge.wikispaces.com/file/view/AqueousDataset002.xlsx/351826032/AqueousDataset002.xlsx

Legend

hvap:	Enthalpy of vaporization at standard conditions
log10ws:	Log10 of Water solubility in mol/l
logp:	Octanol/Water partition coefficient
mcvol:	McGowan's characteristic volume
pvap:	Vapor pressure
rinpola:	Non-polar retention indices
ripola:	Polar retention indices
tf:	Normal melting (fusion) point

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