

# O-Methyl-O-(1,1-difluoro-2-chloroethyl)-N-(3-meth

**Inchi:** InChI=1S/C8H17CIF2NO2PS/c1-7(2)4-5-12-15(16,13-3)14-8(10,11)6-9/h7H,4-6H2,1-3H3  
**InchiKey:** GFVLXLVTFSCVOY-UHFFFAOYSA-N  
**Formula:** C8H17CIF2NO2PS  
**SMILES:** COP(=S)(NCCC(C)C)OC(F)(F)CCI  
**Mol. weight [g/mol]:** 295.71

## Physical Properties

Property code	Value	Unit	Source
log10ws	0.49		Crippen Method
logp	3.341		Crippen Method
mcvol	197.890	ml/mol	McGowan Method
rinpol	1536.00		NIST Webbook
rinpol	1536.00		NIST Webbook

## Sources

**Crippen Method:** <http://pubs.acs.org/doi/abs/10.1021/ci9903071>  
**Crippen Method:** [https://www.cheméo.com/doc/models/crippen\\_log10ws](https://www.cheméo.com/doc/models/crippen_log10ws)  
**McGowan Method:** <http://link.springer.com/article/10.1007/BF02311772>  
**NIST Webbook:** <http://webbook.nist.gov/cgi/cbook.cgi?ID=R544175&Units=SI>

## Legend

**log10ws:** Log10 of Water solubility in mol/l  
**logp:** Octanol/Water partition coefficient  
**mcvol:** McGowan's characteristic volume  
**rinpol:** Non-polar retention indices

Latest version available from:

<https://www.cheméo.com/cid/114-454-1/O-Methyl-O-1-1-difluoro-2-chloroethyl-N-3-methylbutyl-phosphorothioamidat>

Generated by Cheméo on 2024-05-11 07:28:42.657913835 +0000 UTC m=+17701771.578491148.

Cheméo (<https://www.chemeo.com>) is the biggest free database of chemical and physical data for the process industry.