

Me-TMS

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InchiKey: DQKFOBXXAKZGIPX-AZROICAXSA-N

Formula: C₃₄H₆₆O₅Si₃

SMILES: COC(=O)CCC(C)C1CCC2C3C(O[Si](C)(C)C)CC4CC(O[Si](C)(C)C)CCC4(C)C3CC(O[Si](C)(C)C)C1

Mol. weight [g/mol]: 639.14

Physical Properties

Property code	Value	Unit	Source
log10ws	-2.31		Crippen Method
logp	9.115		Crippen Method
rinpol	3211.00		NIST Webbook
rinpol	3211.00		NIST Webbook
rinpol	3211.00		NIST Webbook
ripol	3365.00		NIST Webbook
ripol	3365.00		NIST Webbook

Sources

Crippen Method: <http://pubs.acs.org/doi/abs/10.1021/ci990307l>

Crippen Method: https://www.chemeo.com/doc/models/crippen_log10ws

NIST Webbook: <http://webbook.nist.gov/cgi/cbook.cgi?ID=R533398&Units=SI>

Legend

log10ws:	Log10 of Water solubility in mol/l
logp:	Octanol/Water partition coefficient
rinpol:	Non-polar retention indices
ripol:	Polar retention indices

Latest version available from:

<https://www.cheméo.com/cid/27-825-6/allo-Cholanic-acid-3-alpha-7-alpha-12-alpha-trihydroxy-Me-TMS.pdf>

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