

(20R/S)-3.beta.,29-bis-(Trimethylsilyoxy)lupan-29-

#1
InChI: InChI=1S/C36H66O3Si2/c1-24(31(37)39-41(11,12)13)25-16-19-33(4)22-23-35(6)26(30(2
InChIKey: DKFXWBJRSYECOY-HSWMNOIGSA-N
Formula: C36H66O3Si2
SMILES: CC(C(=O)O[Si](C)(C)C)[C@@H]1CC[C@]2(C)CC[C@]3(C)C(CCC4[C@@]5(C)CC[C@H
Mol. weight [g/mol]: 603.09

Physical Properties

Property code	Value	Unit	Source
logp	10.322		Crippen Method

Sources

Crippen Method: <http://pubs.acs.org/doi/abs/10.1021/ci990307l>
Crippen Method: https://www.cheméo.com/doc/models/crippen_log10ws
Cheméo Relay (1.0):
Cheméo Relay (1.0, beta):
NIST Webbook: <http://webbook.nist.gov/cgi/cbook.cgi?ID=R583269&Units=SI>

Legend

logp: Octanol/Water partition coefficient

Latest version available from:

<https://www.cheméo.com/cid/70-047-2/20R-S-3-beta-29-bis-Trimethylsilyoxy-lupan-29-oate-1.pdf>

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