

«alpha»-Altropyranose, TMS

Inchi: InChI=1S/C21H52O6Si5/c1-28(2,3)22-16-17-18(24-29(4,5)6)19(25-30(7,8)9)20(26-31(10)11)23-15-12-13-14-27-21-20
InchiKey: PPFHNIVPOLWPCF-FCMAGTKHSA-N
Formula: C₂₁H₅₂O₆Si₅
SMILES: C[Si](C)(C)OCC1OC(O[Si](C)(C)C)C(O[Si](C)(C)C)C(O[Si](C)(C)C)C1O[Si](C)(C)C
Mol. weight [g/mol]: 541.06

Physical Properties

Property code	Value	Unit	Source
logp	6.075		Crippen Method
rinpol	1830.00		NIST Webbook
rinpol	1830.00		NIST Webbook
ripol	1703.00		NIST Webbook
ripol	1703.00		NIST Webbook

Sources

Cheméo Relay (1.0, beta):

NIST Webbook: <http://webbook.nist.gov/cgi/cbook.cgi?ID=R52498&Units=SI>

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

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

Cheméo Relay (1.0):

Legend

logp: Octanol/Water partition coefficient

rinpol: Non-polar retention indices

ripol: Polar retention indices

Latest version available from:

<https://www.chemeo.com/cid/49-732-5/alpha-Altropyranose-TMS.pdf>

Generated by Cheméo on 2026-07-22 07:42:18.737027198 +0000 UTC m=+217234.578912668.

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