

# Ipanguline C2

**Inchi:** InChI=1S/C20H33NO7/c1-6-12(2)18(23)28-16-8-10-21-9-7-15(17(16)21)11-26-19(24)20  
**InchiKey:** QSDVUZIHMWDHEK-SRMJLOKFSA-N  
**Formula:** C20H33NO7  
**SMILES:** CCC(C)C(=O)OC1CCN2CCC(COC(=O)C(C)(O)C(C)OC(C)=O)C12  
**Mol. weight [g/mol]:** 399.48

## Physical Properties

| Property code | Value   | Unit   | Source         |
|---------------|---------|--------|----------------|
| log10ws       | -2.37   |        | Crippen Method |
| logp          | 1.284   |        | Crippen Method |
| mcvol         | 309.110 | ml/mol | McGowan Method |
| rinpol        | 2426.00 |        | NIST Webbook   |
| rinpol        | 2426.00 |        | NIST Webbook   |

## Sources

**Crippen Method:** <http://pubs.acs.org/doi/abs/10.1021/ci990307l>  
**Crippen Method:** [https://www.chemeo.com/doc/models/crippen\\_log10ws](https://www.chemeo.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=R394915&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

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