

# 4-Hydroxybenzyl alcohol, bis(heptafluorobutyrate)

|                             |   |
|-----------------------------|---|
| <b>Inchi:</b>               | InChI=1S/C15H6F14O4/c16-10(17,12(20,21)14(24,25)26)8(30)32-5-6-1-3-7(4-2-6)33-9(3 |
| <b>InchiKey:</b>            | FGALMUVNHSMOSX-UHFFFAOYSA-N   |
| <b>Formula:</b>             | C15H6F14O4  |
| <b>SMILES:</b>              | O=C(OCc1ccc(OC(=O)C(F)(F)C(F)(F)C(F)(F)F)cc1)C(F)(F)C(F)(F)C(F)(F)F               |
| <b>Mol. weight [g/mol]:</b> | 516.18  |

## Physical Properties

| Property code | Value    | Unit                 | Source         |
|---------------|----------|----------------------|----------------|
| gf            | -2999.94 | kJ/mol               | Joback Method  |
| hf            | -3415.51 | kJ/mol               | Joback Method  |
| hfus          | 32.47    | kJ/mol               | Joback Method  |
| hvap          | 51.02    | kJ/mol               | Joback Method  |
| log10ws       | -6.62    |                      | Crippen Method |
| logp          | 5.301    |                      | Crippen Method |
| mcvol         | 238.110  | ml/mol               | McGowan Method |
| pc            | 1307.06  | kPa                  | Joback Method  |
| rinpol        | 1287.00  |                      | NIST Webbook   |
| rinpol        | 1287.00  |                      | NIST Webbook   |
| tb            | 697.24   | K                    | Joback Method  |
| tc            | 863.56   | K                    | Joback Method  |
| tf            | 464.85   | K                    | Joback Method  |
| vc            | 1.002    | m <sup>3</sup> /kmol | Joback Method  |

## Temperature Dependent Properties

| Property code | Value  | Unit    | Temperature [K] | Source        |
|---------------|--------|---------|-----------------|---------------|
| cpg           | 719.91 | J/mol×K | 697.24          | Joback Method |
| cpg           | 729.98 | J/mol×K | 724.96          | Joback Method |
| cpg           | 739.18 | J/mol×K | 752.68          | Joback Method |
| cpg           | 747.59 | J/mol×K | 780.40          | Joback Method |
| cpg           | 755.28 | J/mol×K | 808.12          | Joback Method |
| cpg           | 762.32 | J/mol×K | 835.84          | Joback Method |
| cpg           | 768.78 | J/mol×K | 863.56          | Joback Method |

# Sources

|                        |   |
|------------------------|---|
| <b>Crippen Method:</b> | <a href="https://www.chemeo.com/doc/models/crippen_log10ws">https://www.chemeo.com/doc/models/crippen_log10ws</a>                         |
| <b>Joback Method:</b>  | <a href="https://en.wikipedia.org/wiki/Joback_method">https://en.wikipedia.org/wiki/Joback_method</a>                                     |
| <b>McGowan Method:</b> | <a href="http://link.springer.com/article/10.1007/BF02311772">http://link.springer.com/article/10.1007/BF02311772</a>                     |
| <b>NIST Webbook:</b>   | <a href="http://webbook.nist.gov/cgi/cbook.cgi?ID=U376194&amp;Units=SI">http://webbook.nist.gov/cgi/cbook.cgi?ID=U376194&amp;Units=SI</a> |
| <b>Crippen Method:</b> | <a href="http://pubs.acs.org/doi/abs/10.1021/ci990307I">http://pubs.acs.org/doi/abs/10.1021/ci990307I</a>                                 |

# Legend

|                 |   |
|-----------------|---|
| <b>cpg:</b>     | Ideal gas heat capacity                         |
| <b>gf:</b>      | Standard Gibbs free energy of formation         |
| <b>hf:</b>      | Enthalpy of formation at standard conditions    |
| <b>hfus:</b>    | Enthalpy of fusion at standard conditions       |
| <b>hvpap:</b>   | Enthalpy of vaporization at standard conditions |
| <b>log10ws:</b> | Log10 of Water solubility in mol/l              |
| <b>logp:</b>    | Octanol/Water partition coefficient             |
| <b>mcvol:</b>   | McGowan's characteristic volume                 |
| <b>pc:</b>      | Critical Pressure                               |
| <b>rinpola:</b> | Non-polar retention indices                     |
| <b>tb:</b>      | Normal Boiling Point Temperature                |
| <b>tc:</b>      | Critical Temperature                            |
| <b>tf:</b>      | Normal melting (fusion) point                   |
| <b>vc:</b>      | Critical Volume                                 |

Latest version available from:

<https://www.chemeo.com/cid/117-474-6/4-Hydroxybenzyl-alcohol-bis-heptafluorobutyrate.pdf>

Generated by Cheméo on 2024-04-28 02:33:11.978245782 +0000 UTC m=+16560840.898823094.

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