

1,2,3,8,9-pentabromo-dibenzo-dioxin

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|-----------------------------|---|
| Inchi: | InChI=1S/C12H3Br5O2/c13-4-1-2-6-11(9(4)16)19-12-7(18-6)3-5(14)8(15)10(12)17/h1-3H |
| InchiKey: | BASDYTCMFMVWAF-UHFFFAOYSA-N |
| Formula: | C12H3Br5O2 |
| SMILES: | BrC1cc2c(c(Br)c1Br)Oc1c(ccc(Br)c1Br)O2 |
| Mol. weight [g/mol]: | 578.67 |

Physical Properties

| Property code | Value | Unit | Source |
|---------------|---------|----------------------|----------------|
| gf | 187.49 | kJ/mol | Joback Method |
| hf | 68.71 | kJ/mol | Joback Method |
| hfus | 53.74 | kJ/mol | Joback Method |
| hvap | 92.74 | kJ/mol | Joback Method |
| log10ws | -8.80 | | Crippen Method |
| logp | 7.397 | | Crippen Method |
| mcvol | 220.800 | ml/mol | McGowan Method |
| pc | 5503.26 | kPa | Joback Method |
| rinpol | 3223.00 | | NIST Webbook |
| rinpol | 3223.00 | | NIST Webbook |
| tb | 954.02 | K | Joback Method |
| tc | 1259.09 | K | Joback Method |
| tf | 743.32 | K | Joback Method |
| vc | 0.809 | m ³ /kmol | Joback Method |

Temperature Dependent Properties

| Property code | Value | Unit | Temperature [K] | Source |
|---------------|-----------|---------|-----------------|---------------|
| cpg | 428.97 | J/molxK | 954.02 | Joback Method |
| cpg | 436.52 | J/molxK | 1004.86 | Joback Method |
| cpg | 444.52 | J/molxK | 1055.71 | Joback Method |
| cpg | 453.23 | J/molxK | 1106.55 | Joback Method |
| cpg | 462.89 | J/molxK | 1157.40 | Joback Method |
| cpg | 473.76 | J/molxK | 1208.24 | Joback Method |
| cpg | 486.09 | J/molxK | 1259.09 | Joback Method |
| dvisc | 0.0006039 | Paxs | 743.32 | Joback Method |

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|-------|-----------|------|--------|---------------|
| dvisc | 0.0005175 | Paxs | 778.44 | Joback Method |
| dvisc | 0.0004494 | Paxs | 813.55 | Joback Method |
| dvisc | 0.0003948 | Paxs | 848.67 | Joback Method |
| dvisc | 0.0003505 | Paxs | 883.79 | Joback Method |
| dvisc | 0.0003140 | Paxs | 918.90 | Joback Method |
| dvisc | 0.0002835 | Paxs | 954.02 | Joback Method |

Sources

| | |
|------------------------|---|
| McGowan Method: | http://link.springer.com/article/10.1007/BF02311772 |
| NIST Webbook: | http://webbook.nist.gov/cgi/cbook.cgi?ID=R172339&Units=SI |
| Crippen Method: | http://pubs.acs.org/doi/abs/10.1021/ci9903071 |
| Crippen Method: | https://www.chemeo.com/doc/models/crippen_log10ws |
| Joback Method: | https://en.wikipedia.org/wiki/Joback_method |

Legend

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

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<https://www.chemeo.com/cid/117-719-4/1-2-3-8-9-pentabromo-dibenzo-dioxin.pdf>

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