

# Benzoic acid, 3-methyl-

|                      |                                                                                                      |
|----------------------|------------------------------------------------------------------------------------------------------|
| Other names:         | 3-Methylbenzoic acid<br>m-Methylbenzoic acid<br>m-Toluic acid<br>m-Toluylic acid<br>meta-Toluic acid |
| Inchi:               | InChI=1S/C8H8O2/c1-6-3-2-4-7(5-6)8(9)10/h2-5H,1H3,(H,9,10)                                           |
| InchiKey:            | GPSDUZXPYCFOSQ-UHFFFAOYSA-N                                                                          |
| Formula:             | C8H8O2                                                                                               |
| SMILES:              | Cc1cccc(C(=O)O)c1                                                                                    |
| Mol. weight [g/mol]: | 136.15                                                                                               |
| CAS:                 | 99-04-7                                                                                              |

## Physical Properties

| Property code | Value           | Unit   | Source                               |
|---------------|-----------------|--------|--------------------------------------|
| affp          | 829.80          | kJ/mol | NIST Webbook                         |
| basg          | 798.80          | kJ/mol | NIST Webbook                         |
| chs           | -3866.50 ± 0.90 | kJ/mol | NIST Webbook                         |
| chs           | -3865.30 ± 0.79 | kJ/mol | NIST Webbook                         |
| chs           | -3862.00        | kJ/mol | NIST Webbook                         |
| gf            | -146.48         | kJ/mol | Joback Method                        |
| hf            | -329.10         | kJ/mol | NIST Webbook                         |
| hf            | -327.90 ± 1.40  | kJ/mol | NIST Webbook                         |
| hfs           | -426.10 ± 0.96  | kJ/mol | NIST Webbook                         |
| hfs           | -424.90 ± 1.40  | kJ/mol | NIST Webbook                         |
| hfus          | 15.81           | kJ/mol | Joback Method                        |
| hsub          | 97.00           | kJ/mol | NIST Webbook                         |
| hsub          | 97.00 ± 0.30    | kJ/mol | NIST Webbook                         |
| hsub          | 97.00 ± 0.30    | kJ/mol | NIST Webbook                         |
| hvap          | 59.77           | kJ/mol | Joback Method                        |
| ie            | 9.40 ± 0.20     | eV     | NIST Webbook                         |
| log10ws       | -2.14           |        | Aqueous Solubility Prediction Method |
| logp          | 1.693           |        | Crippen Method                       |
| mcvol         | 107.260         | ml/mol | McGowan Method                       |
| pc            | 4356.87         | kPa    | Joback Method                        |
| rinpol        | 1325.00         |        | NIST Webbook                         |
| rinpol        | 1249.00         |        | NIST Webbook                         |

|    |               |         |                                                                                                                                                                              |
|----|---------------|---------|------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| tb | 536.20        | K       | NIST Webbook                                                                                                                                                                 |
| tc | 771.00        | K       | Vapor-liquid critical point measurements of fifteen compounds by the pulse-heating method                                                                                    |
| tf | 382.40        | K       | Abraham model correlations for describing the thermodynamic properties of solute transfer into pentyl acetate based on headspace chromatographic and solubility measurements |
| tf | 383.07        | K       | Aqueous Solubility Prediction Method                                                                                                                                         |
| tf | 384.40 ± 2.00 | K       | NIST Webbook                                                                                                                                                                 |
| tf | 384.00 ± 1.50 | K       | NIST Webbook                                                                                                                                                                 |
| tf | 381.90 ± 0.20 | K       | NIST Webbook                                                                                                                                                                 |
| vc | 0.401         | m3/kmol | Joback Method                                                                                                                                                                |

## Temperature Dependent Properties

| Property code | Value     | Unit    | Temperature [K] | Source        |
|---------------|-----------|---------|-----------------|---------------|
| cpg           | 262.13    | J/mol×K | 663.82          | Joback Method |
| cpg           | 253.91    | J/mol×K | 629.27          | Joback Method |
| cpg           | 283.75    | J/mol×K | 767.50          | Joback Method |
| cpg           | 277.03    | J/mol×K | 732.94          | Joback Method |
| cpg           | 269.83    | J/mol×K | 698.38          | Joback Method |
| cpg           | 235.84    | J/mol×K | 560.15          | Joback Method |
| cpg           | 245.16    | J/mol×K | 594.71          | Joback Method |
| cps           | 163.60    | J/mol×K | 298.00          | NIST Webbook  |
| dvisc         | 0.0019990 | Pa×s    | 368.03          | Joback Method |
| dvisc         | 0.0009222 | Pa×s    | 406.46          | Joback Method |
| dvisc         | 0.0004863 | Pa×s    | 444.88          | Joback Method |
| dvisc         | 0.0002839 | Pa×s    | 483.30          | Joback Method |
| dvisc         | 0.0001794 | Pa×s    | 521.73          | Joback Method |
| dvisc         | 0.0051891 | Pa×s    | 329.61          | Joback Method |
| dvisc         | 0.0001208 | Pa×s    | 560.15          | Joback Method |
| hfust         | 15.73     | kJ/mol  | 381.90          | NIST Webbook  |
| hfust         | 15.73     | kJ/mol  | 381.90          | NIST Webbook  |
| hfust         | 15.73     | kJ/mol  | 381.90          | NIST Webbook  |
| hvapt         | 62.80     | kJ/mol  | 503.00          | NIST Webbook  |
| sfust         | 41.20     | J/mol×K | 381.90          | NIST Webbook  |

# Sources

|                                                                                                                                                                                                                                                               |                                                                                                                                                                                                                                                                                                                                                                                                                                                                        |
|---------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| McGowan Method:                                                                                                                                                                                                                                               | <a href="http://link.springer.com/article/10.1007/BF02311772">http://link.springer.com/article/10.1007/BF02311772</a>                                                                                                                                                                                                                                                                                                                                                  |
| Aqueous Solubility Prediction Method:                                                                                                                                                                                                                         | <a href="http://onschallenge.wikispaces.com/file/view/AqueousDataset002.xlsx/351826032/AqueousDataset002.xlsx">http://onschallenge.wikispaces.com/file/view/AqueousDataset002.xlsx/351826032/AqueousDataset002.xlsx</a>                                                                                                                                                                                                                                                |
| Abraham model correlations for describing the thermodynamic properties of solute transfer into pentyl acetate based on headspace chromatographic and solubility measurements:                                                                                 | <a href="https://www.doi.org/10.1016/j.jct.2018.05.003">https://www.doi.org/10.1016/j.jct.2018.05.003</a>                                                                                                                                                                                                                                                                                                                                                              |
| Joback Method:                                                                                                                                                                                                                                                | <a href="https://en.wikipedia.org/wiki/Joback_method">https://en.wikipedia.org/wiki/Joback_method</a>                                                                                                                                                                                                                                                                                                                                                                  |
| Crippen Method:                                                                                                                                                                                                                                               | <a href="http://pubs.acs.org/doi/abs/10.1021/ci990307l">http://pubs.acs.org/doi/abs/10.1021/ci990307l</a>                                                                                                                                                                                                                                                                                                                                                              |
| Solubilities of Benzoic Acid, p-Methylbenzoic Acid, m-Methylbenzoic Acid, o-Methylbenzoic Acid, p-Methoxybenzoic Acid, and p-tert-butylbenzoic acid derivatives in water-liquid critical point measurements of fifteen compounds by the pulse-heating method: | <a href="https://www.doi.org/10.1021/je700677d">https://www.doi.org/10.1021/je700677d</a><br><a href="http://webbook.nist.gov/cgi/cbook.cgi?ID=C99047&amp;Units=SI">http://webbook.nist.gov/cgi/cbook.cgi?ID=C99047&amp;Units=SI</a><br><a href="https://www.doi.org/10.1016/j.fluid.2006.10.014">https://www.doi.org/10.1016/j.fluid.2006.10.014</a><br><a href="https://www.doi.org/10.1016/j.fluid.2014.07.038">https://www.doi.org/10.1016/j.fluid.2014.07.038</a> |

## Legend

|          |                                                          |
|----------|----------------------------------------------------------|
| affp:    | Proton affinity                                          |
| basg:    | Gas basicity                                             |
| chs:     | Standard solid enthalpy of combustion                    |
| cpg:     | Ideal gas heat capacity                                  |
| cps:     | Solid phase heat capacity                                |
| dvisc:   | Dynamic viscosity                                        |
| gf:      | Standard Gibbs free energy of formation                  |
| hf:      | Enthalpy of formation at standard conditions             |
| hfs:     | Solid phase enthalpy of formation at standard conditions |
| hfus:    | Enthalpy of fusion at standard conditions                |
| hfust:   | Enthalpy of fusion at a given temperature                |
| hsub:    | Enthalpy of sublimation at standard conditions           |
| hvap:    | Enthalpy of vaporization at standard conditions          |
| hvapt:   | Enthalpy of vaporization at a given temperature          |
| ie:      | Ionization energy                                        |
| 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                              |
| sfust:   | Entropy of fusion at a given temperature                 |
| tb:      | Normal Boiling Point Temperature                         |
| tc:      | Critical Temperature                                     |
| tf:      | Normal melting (fusion) point                            |
| vc:      | Critical Volume                                          |

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