

# 2-Heptanone, 4-methyl-

|                      |                                                    |
|----------------------|----------------------------------------------------|
| Other names:         | 4-Methyl-2-heptanone                               |
| Inchi:               | InChI=1S/C8H16O/c1-4-5-7(2)6-8(3)9/h7H,4-6H2,1-3H3 |
| InchiKey:            | BIFPSSTVLFHHEI-UHFFFAOYSA-N                        |
| Formula:             | C8H16O                                             |
| SMILES:              | CCCC(C)CC(C)=O                                     |
| Mol. weight [g/mol]: | 128.21                                             |
| CAS:                 | 6137-06-0                                          |

## Physical Properties

| Property code | Value         | Unit    | Source         |
|---------------|---------------|---------|----------------|
| gf            | -114.88       | kJ/mol  | Joback Method  |
| hf            | -326.31       | kJ/mol  | Joback Method  |
| hfus          | 14.55         | kJ/mol  | Joback Method  |
| hvap          | 39.76         | kJ/mol  | Joback Method  |
| log10ws       | -2.21         |         | Crippen Method |
| logp          | 2.402         |         | Crippen Method |
| mcvol         | 125.150       | ml/mol  | McGowan Method |
| pc            | 2726.86       | kPa     | Joback Method  |
| rinpol        | 949.00        |         | NIST Webbook   |
| rinpol        | 943.00        |         | NIST Webbook   |
| rinpol        | 936.40        |         | NIST Webbook   |
| rinpol        | 943.00        |         | NIST Webbook   |
| rinpol        | 936.40        |         | NIST Webbook   |
| rinpol        | 949.00        |         | NIST Webbook   |
| ripol         | 1206.00       |         | NIST Webbook   |
| ripol         | 1213.00       |         | NIST Webbook   |
| ripol         | 1206.00       |         | NIST Webbook   |
| ripol         | 1206.00       |         | NIST Webbook   |
| ripol         | 1224.00       |         | NIST Webbook   |
| ripol         | 1206.00       |         | NIST Webbook   |
| ripol         | 1224.00       |         | NIST Webbook   |
| tb            | 429.00 ± 5.00 | K       | NIST Webbook   |
| tc            | 615.20        | K       | Joback Method  |
| tf            | 214.85        | K       | Joback Method  |
| vc            | 0.483         | m3/kmol | Joback Method  |

# Temperature Dependent Properties

| Property code | Value     | Unit    | Temperature [K] | Source        |
|---------------|-----------|---------|-----------------|---------------|
| cpg           | 255.56    | J/mol×K | 435.87          | Joback Method |
| cpg           | 268.41    | J/mol×K | 465.76          | Joback Method |
| cpg           | 280.75    | J/mol×K | 495.65          | Joback Method |
| cpg           | 292.58    | J/mol×K | 525.54          | Joback Method |
| cpg           | 303.91    | J/mol×K | 555.42          | Joback Method |
| cpg           | 314.75    | J/mol×K | 585.31          | Joback Method |
| cpg           | 325.12    | J/mol×K | 615.20          | Joback Method |
| dvisc         | 0.0067893 | Pa×s    | 214.85          | Joback Method |
| dvisc         | 0.0027108 | Pa×s    | 251.69          | Joback Method |
| dvisc         | 0.0013683 | Pa×s    | 288.52          | Joback Method |
| dvisc         | 0.0008063 | Pa×s    | 325.36          | Joback Method |
| dvisc         | 0.0005291 | Pa×s    | 362.20          | Joback Method |
| dvisc         | 0.0003753 | Pa×s    | 399.03          | Joback Method |
| dvisc         | 0.0002821 | Pa×s    | 435.87          | Joback Method |

## Correlations

| Information                 | Value                         |
|-----------------------------|-------------------------------|
| Property code               | pvap                          |
| Equation                    | $\ln(P_{vp}) = A + B/(T + C)$ |
| Coeff. A                    | 1.48402e+01                   |
| Coeff. B                    | -3.82591e+03                  |
| Coeff. C                    | -6.15380e+01                  |
| Temperature range (K), min. | 324.44                        |
| Temperature range (K), max. | 463.05                        |

## Sources

McGowan Method:

<http://link.springer.com/article/10.1007/BF02311772>

NIST Webbook:

<http://webbook.nist.gov/cgi/cbook.cgi?ID=C6137060&Units=SI>

The Yaws Handbook of Vapor Pressure:

<https://www.sciencedirect.com/book/9780128029992/the-yaws-handbook-of-vapor-pressure>

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)

## Legend

|                 |                                                 |
|-----------------|-------------------------------------------------|
| <b>cpg:</b>     | Ideal gas heat capacity                         |
| <b>dvisc:</b>   | Dynamic viscosity                               |
| <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>hvap:</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>pvap:</b>    | Vapor pressure                                  |
| <b>rinpol:</b>  | Non-polar retention indices                     |
| <b>ripol:</b>   | 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                                 |

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