

# (Z)-1-Phenylpropene

<b>Other names:</b>	(Z)-1-Propenylbenzene 1-Phenyl-1-propene, cis- Benzene, (1Z)-1-propenyl- Benzene, 1-propenyl-, (Z)- Benzene, propen-1-yl-, (Z)- Benzene, propenyl-, (Z)- cis-1-Phenyl-1-propene cis-1-Phenylpropene cis-1-Propenylbenzene cis-Propenylbenzene cis-«beta»-Methylstyrene cis-Â«betaÂ»-Methylstyrene
<b>Inchi:</b>	InChI=1S/C9H10/c1-2-6-9-7-4-3-5-8-9/h2-8H,1H3/b6-2-
<b>InchiKey:</b>	QROGIFZRVHSFLM-KXFIGUGUSA-N
<b>Formula:</b>	C9H10
<b>SMILES:</b>	CC=Cc1ccccc1
<b>Mol. weight [g/mol]:</b>	118.18
<b>CAS:</b>	766-90-5

## Physical Properties

Property code	Value	Unit	Source
affp	836.40	kJ/mol	NIST Webbook
basg	807.50	kJ/mol	NIST Webbook
chg	-5092.14	kJ/mol	NIST Webbook
gf	217.53	kJ/mol	Joback Method
hf	124.66	kJ/mol	Joback Method
hfus	13.31	kJ/mol	Joback Method
hvap	37.86	kJ/mol	Joback Method
ie	8.15	eV	NIST Webbook
ie	8.15	eV	NIST Webbook
ie	8.45	eV	NIST Webbook
ie	8.48	eV	NIST Webbook
ie	8.28	eV	NIST Webbook
log10ws	-2.71		Crippen Method
logp	2.720		Crippen Method
mcvol	109.610	ml/mol	McGowan Method
pc	3464.28	kPa	Joback Method

rinpol	975.30		NIST Webbook
rinpol	975.00		NIST Webbook
rinpol	155.00		NIST Webbook
rinpol	975.00		NIST Webbook
rinpol	155.00		NIST Webbook
rinpol	973.00		NIST Webbook
rinpol	973.00		NIST Webbook
rinpol	984.00		NIST Webbook
ripol	1324.30		NIST Webbook
ripol	1324.00		NIST Webbook
tb	440.58 ± 0.30	K	NIST Webbook
tb	441.65 ± 3.00	K	NIST Webbook
tc	654.10	K	Joback Method
tf	211.23 ± 0.20	K	NIST Webbook
tf	203.40 ± 1.00	K	NIST Webbook
tf	211.47 ± 0.15	K	NIST Webbook
vc	0.411	m <sup>3</sup> /kmol	Joback Method

## Temperature Dependent Properties

Property code	Value	Unit	Temperature [K]	Source
cpg	268.59	J/mol×K	654.10	Joback Method
cpg	199.03	J/mol×K	436.16	Joback Method
cpg	212.72	J/mol×K	472.48	Joback Method
cpg	225.50	J/mol×K	508.81	Joback Method
cpg	237.43	J/mol×K	545.13	Joback Method
cpg	248.56	J/mol×K	581.46	Joback Method
cpg	258.93	J/mol×K	617.78	Joback Method
dvisc	0.0001928	Paxs	436.16	Joback Method
dvisc	0.0032236	Paxs	212.53	Joback Method
dvisc	0.0014202	Paxs	249.80	Joback Method
dvisc	0.0007741	Paxs	287.07	Joback Method
dvisc	0.0004851	Paxs	324.34	Joback Method
dvisc	0.0003347	Paxs	361.62	Joback Method
dvisc	0.0002476	Paxs	398.89	Joback Method
hvapt	44.80	kJ/mol	423.00	NIST Webbook

# Correlations

Information	Value
Property code	pvap
Equation	$\ln(P_{vp}) = A + B/(T + C)$
Coeff. A	1.56145e+01
Coeff. B	-4.32144e+03
Coeff. C	-4.75840e+01
Temperature range (K), min.	329.54
Temperature range (K), max.	467.02

## Sources

Joback Method:	<a href="https://en.wikipedia.org/wiki/Joback_method">https://en.wikipedia.org/wiki/Joback_method</a>
McGowan Method:	<a href="http://link.springer.com/article/10.1007/BF02311772">http://link.springer.com/article/10.1007/BF02311772</a>
NIST Webbook:	<a href="http://webbook.nist.gov/cgi/cbook.cgi?ID=C766905&amp;Units=SI">http://webbook.nist.gov/cgi/cbook.cgi?ID=C766905&amp;Units=SI</a>
The Yaws Handbook of Vapor Pressure:	<a href="https://www.sciencedirect.com/book/9780128029992/the-yaws-handbook-of-vapor-pressure">https://www.sciencedirect.com/book/9780128029992/the-yaws-handbook-of-vapor-pressure</a>
Crippen Method:	<a href="http://pubs.acs.org/doi/abs/10.1021/ci9903071">http://pubs.acs.org/doi/abs/10.1021/ci9903071</a>
Crippen Method:	<a href="https://www.chemeo.com/doc/models/crippen_log10ws">https://www.chemeo.com/doc/models/crippen_log10ws</a>

## Legend

<b>affp:</b>	Proton affinity
<b>basg:</b>	Gas basicity
<b>chg:</b>	Standard gas enthalpy of combustion
<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>hvapt:</b>	Enthalpy of vaporization at a given temperature
<b>ie:</b>	Ionization energy
<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

Latest version available from:

<https://www.cheméo.com/cid/48-884-8/Z-1-Phenylpropene.pdf>

Generated by Cheméo on 2024-04-19 18:43:42.412588429 +0000 UTC m=+15841471.333165773.

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