## Butanoic acid, 3-methylbutyl ester

Other names: 3-Methyl-1-butyl butanoate

3-Methylbutyl butanoate3-Methylbutyl butyrate3-Methylbutyl n-butyrateButyric acid, isopentyl ester

Isoamyl butanoate Isoamyl butylate Isoamyl butyrate Isoamyl-n-butyrate

Isopentyl alcohol, butyrate

Isopentyl butanoate Isopentyl butyrate Isopentyl-n-butyrate

UN 2620

Inchi: InChl=1S/C9H18O2/c1-4-5-9(10)11-7-6-8(2)3/h8H,4-7H2,1-3H3

InchiKey: PQLMXFQTAMDXIZ-UHFFFAOYSA-N

Formula: C9H18O2

SMILES: CCCC(=O)OCCC(C)C

Mol. weight [g/mol]: 158.24 CAS: 106-27-4

### **Physical Properties**

Property code	Value	Unit	Source
chl	-5491.10	kJ/mol	NIST Webbook
gf	-211.46	kJ/mol	Joback Method
hf	-479.17	kJ/mol	Joback Method
hfus	18.33	kJ/mol	Joback Method
hvap	44.40	kJ/mol	Joback Method
log10ws	-2.21		Crippen Method
logp	2.376		Crippen Method
mcvol	145.110	ml/mol	McGowan Method
рс	2431.44	kPa	Joback Method
rinpol	1002.00	NIST Webbook	
rinpol	1007.00	NIST Webbook	
rinpol	1039.00	NIST Webbook	
rinpol	1002.00	NIST Webbook	
rinpol	1041.00		NIST Webbook

	1002.22	\!! <b>?</b>
rinpol	1039.00	NIST Webbook
rinpol	1045.00	NIST Webbook
rinpol	1020.00	NIST Webbook
rinpol	1029.00	NIST Webbook
rinpol	1054.00	NIST Webbook
rinpol	1064.00	NIST Webbook
rinpol	1060.00	NIST Webbook
rinpol	1068.00	NIST Webbook
rinpol	1086.00	NIST Webbook
rinpol	1041.00	NIST Webbook
rinpol	1041.00	NIST Webbook
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rinpol	1008.00	NIST Webbook
rinpol	1060.00	NIST Webbook
rinpol	1013.00	NIST Webbook
rinpol	1042.00	NIST Webbook
rinpol	1038.00	NIST Webbook
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rinpol	1046.00	NIST Webbook

rinpol	1042.00	NIST Webbook
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rinpol	1016.00	NIST Webbook
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ripol	1256.00	NIST Webbook
ripol	1266.00	NIST Webbook
ripol	1266.00	NIST Webbook
ripol	1283.00	NIST Webbook

ripol	1255.00		NIST Webbook
ripol	1267.00		NIST Webbook
tb	451.70 ± 0.50	K	NIST Webbook
tb	452.20	K	NIST Webbook
tb	454.20 ± 0.30	K	NIST Webbook
tb	451.80 ± 0.50	K	NIST Webbook
tb	454.20 ± 0.25	K	NIST Webbook
tb	451.70 ± 0.50	K	NIST Webbook
tb	442.55 ± 1.00	K	NIST Webbook
tc	618.83 ± 6.00	K	NIST Webbook
tf	248.35	K	Joback Method
VC	0.557	m3/kmol	Joback Method

# **Temperature Dependent Properties**

Property code	Value	Unit	Temperature [K]	Source
cpg	323.06	J/mol×K	481.17	Joback Method
cpg	397.15	J/mol×K	658.49	Joback Method
cpg	386.04	J/mol×K	628.93	Joback Method
cpg	374.44	J/mol×K	599.38	Joback Method
cpg	362.35	J/mol×K	569.83	Joback Method
cpg	349.76	J/mol×K	540.28	Joback Method
cpg	336.66	J/mol×K	510.72	Joback Method
dvisc	0.0006860	Pa×s	323.15	Density and Viscosity Correlation for Several Common Fragrance and Flavor Esters
dvisc	0.0006460	Paxs	328.15	Density and Viscosity Correlation for Several Common Fragrance and Flavor Esters
dvisc	0.0006090	Paxs	333.15	Density and Viscosity Correlation for Several Common Fragrance and Flavor Esters
dvisc	0.0005750	Paxs	338.15	Density and Viscosity Correlation for Several Common Fragrance and Flavor Esters

dvisc	0.0005450	Pa×s	343.15	Density and Viscosity Correlation for Several Common Fragrance and Flavor Esters	
dvisc	0.0013170	Paxs	288.15	Densities and Viscosities of Binary Mixtures of Isoamyl Acetate, Ethyl Caproate, Ethyl Benzoate, Isoamyl Butyrate, Ethyl Phenylacetate, and Ethyl Caprylate with Ethanol at T = (288.15, 298.15, 308.15, and 318.15) K	
dvisc	0.0011110	Paxs	298.15	Densities and Viscosities of Binary Mixtures of Isoamyl Acetate, Ethyl Caproate, Ethyl Benzoate, Isoamyl Butyrate, Ethyl Phenylacetate, and Ethyl Caprylate with Ethanol at T = (288.15, 298.15, 308.15, and 318.15) K	
dvisc	0.0009520	Paxs	308.15	Densities and Viscosities of Binary Mixtures of Isoamyl Acetate, Ethyl Caproate, Ethyl Benzoate, Isoamyl Butyrate, Ethyl Phenylacetate, and Ethyl Caprylate with Ethanol at T = (288.15, 298.15, 308.15, and 318.15) K	

dvisc	0.0008270	Paxs	318.15	Densities and Viscosities of Binary Mixtures of Isoamyl Acetate, Ethyl Caproate, Ethyl Benzoate, Isoamyl Butyrate, Ethyl Phenylacetate, and Ethyl Caprylate with Ethanol at T = (288.15, 298.15, 308.15, and 318.15) K	
dvisc	0.0010440	Paxs	293.15	Density and Viscosity Correlation for Several Common Fragrance and Flavor Esters	
dvisc	0.0009670	Paxs	298.15	Density and Viscosity Correlation for Several Common Fragrance and Flavor Esters	
dvisc	0.0008970	Paxs	303.15	Density and Viscosity Correlation for Several Common Fragrance and Flavor Esters	
dvisc	0.0008360	Paxs	308.15	Density and Viscosity Correlation for Several Common Fragrance and Flavor Esters	
dvisc	0.0007810	Paxs	313.15	Density and Viscosity Correlation for Several Common Fragrance and Flavor Esters	
dvisc	0.0007310	Paxs	318.15	Density and Viscosity Correlation for Several Common Fragrance and Flavor Esters	
hvapt	47.40	kJ/mol	373.00	NIST Webbook	

rhol	858.69	kg/m3	298.15	Refractive Indices and Surface Tensions of Binary Mixtures of Isoamyl Acetate, Ethyl Caproate, Ethyl Benzoate, Isoamyl Butyrate, Ethyl Phenylacetate, and Ethyl Caprylate with Ethanol at (288.15, 298.15, 308.15, and 318.15) K	
srf	0.02	N/m	358.15	Densities, Viscosities, Refractive Indices, and Surface Tensions for 12 Flavor Esters from T) 288.15 K to T) 358.15 K	
srf	0.02	N/m	348.15	Densities, Viscosities, Refractive Indices, and Surface Tensions for 12 Flavor Esters from T) 288.15 K to T) 358.15 K	
srf	0.02	N/m	328.15	Densities, Viscosities, Refractive Indices, and Surface Tensions for 12 Flavor Esters from T ) 288.15 K to T ) 358.15 K	
srf	0.02	N/m	318.15	Densities, Viscosities, Refractive Indices, and Surface Tensions for 12 Flavor Esters from T) 288.15 K to T) 358.15 K	
srf	0.02	N/m	308.15	Densities, Viscosities, Refractive Indices, and Surface Tensions for 12 Flavor Esters from T ) 288.15 K to T ) 358.15 K	

srf	0.02	N/m	298.15	Densities, Viscosities, Refractive Indices, and Surface Tensions for 12 Flavor Esters from T) 288.15 K to T) 358.15 K	
srf	0.03	N/m	288.15	Densities, Viscosities, Refractive Indices, and Surface Tensions for 12 Flavor Esters from T) 288.15 K to T) 358.15 K	
srf	0.02	N/m	338.15	Densities, Viscosities, Refractive Indices, and Surface Tensions for 12 Flavor Esters from T) 288.15 K to T) 358.15 K	

### **Correlations**

Information Value

Property code	pvap
Equation	ln(Pvp) = A + B/(T + C)
Coeff. A	1.51047e+01
Coeff. B	-4.02835e+03
Coeff. C	-6.68480e+01
Temperature range (K), min.	338.72
Temperature range (K), max.	478.19

#### **Sources**

Densities, Viscosities, Refractive Indices, and Surface Tensions for 12 Flav งักษรเปลาพิธิการ p2ชั่วเคียง to T) ชีวธรรมเล:
Density and Viscosity Correlation for Several Common Fragrance and Flavor ประจาชายงาน

**Crippen Method:** 

https://www.doi.org/10.1021/je050170x

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

https://www.doi.org/10.1021/je050001c

http://webbook.nist.gov/cgi/cbook.cgi?ID=C106274&Units=SI

http://pubs.acs.org/doi/abs/10.1021/ci990307l

**Refractive Indices and Surface** Tensions of Binary Mixtures of Isoamyl **Aceaale,M⊇thy**PCaproate, Ethyl Benzoate, Isoamyl Butyrate, Ethyl Hoffylacetale, and Ethyl Caprylate with http://link.springer.com/article/10.1007/BF02311772 Ethanol at (288 15, 298 15, 308 15 and belistings and viscosities of Binary Mixtures of Isoamyl Acetate, Ethyl Ceippeand Ethyl Phenylacetate, and Ethyl Copyrigate with Ethanol at T Ethyl Caprylate with Ethanol at T = (288.15, 298.15, 308.15, and 318.15) K:

https://www.doi.org/10.1021/je0601208

https://en.wikipedia.org/wiki/Joback\_method

https://www.doi.org/10.1021/je050389b

https://www.chemeo.com/doc/models/crippen\_log10ws

#### Legend

chl: Standard liquid enthalpy of combustion

Ideal gas heat capacity cpg:

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 Enthalpy of vaporization at a given temperature hvapt:

log10ws: Log10 of Water solubility in mol/l Octanol/Water partition coefficient logp: mcvol: McGowan's characteristic volume

Critical Pressure pc: Vapor pressure pvap: rhol: Liquid Density

rinpol: Non-polar retention indices

ripol: Polar retention indices

Surface Tension srf:

Normal Boiling Point Temperature tb:

Critical Temperature tc:

tf: Normal melting (fusion) point

Critical Volume VC:

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