

N,N'-di-2-Naphthyl-p-phenylenediamine

Other names:

Diafen NN
1,4-Benzenediamine, N,N'-di-2-naphthalenyl-
p-Phenylenediamine, N,N'-di-2-naphthyl-
s-Di(«beta»-naphthyl)-p-phenylenediamine
Aceto DIPP
AgeRite W
AgeRite White
Antigene F
Antioxidant DNP
Antioxidant 123
ASM-DNT
Di(«beta»-naphthyl)-p-phenyldiamine
Di(«beta»-naphthyl)-p-phenylenediamine
DNPD
DNPDA
N,N'-p-Phenylenebis(2-naphthylamine)
N,N'-Bis(«beta»-naphthyl)-p-phenylenediamine
N,N'-Di(«beta»-naphthyl)-p-phenylenediamine
Nonox CL
Santowhite CL
Tisperse MB-2X
1,4-Bis(2-naphthylamino)benzene
2-Naphthyl-p-phenylenediamine
N,N'-Bis-(2-naftyl)-p-fenylendiamin
N,N'-Bis(2-naphthyl)-p-phenylenediamine
sym-Di-«beta»-naphthyl-p-phenylenediamine
Dwu-«beta»-naftylo-p-fenylodwuamina
1,4-Benzenediamine, N,N'-bis-2-naphthylenyl-
DBNPD
N,N'-Di-beta-naphthyl-p-phenylenediamine
Benzene-1,4-diamine, N,N'-di(2-naphthyl)-
N,N'-Di-(2-naphthyl)-1,4-diaminobenzene
NSC 3410

Inchi:

InChI=1S/C26H20N2/c1-3-7-21-17-25(11-9-19(21)5-1)27-23-13-15-24(16-14-23)28-26-1

InchiKey:

VETPHHXZEJAYOB-UHFFFAOYSA-N

Formula:

C₂₆H₂₀N₂

SMILES:

c1ccc2cc(Nc3ccc(Nc4ccc5ccccc5c4)cc3)ccc2c1

Mol. weight [g/mol]:

360.45

CAS:

93-46-9

Physical Properties

Property code	Value	Unit	Source
gf	868.46	kJ/mol	Joback Method
hf	584.29	kJ/mol	Joback Method
h _{fus}	48.29	kJ/mol	Joback Method
h _{vap}	98.44	kJ/mol	Joback Method
log ₁₀ w _s	-8.69		Crippen Method
log _p	7.480		Crippen Method
m _{cvol}	286.960	ml/mol	McGowan Method
pc	1975.31	kPa	Joback Method
tb	1027.56	K	Joback Method
tc	1299.52	K	Joback Method
tf	670.32	K	Joback Method
vc	1.081	m ³ /kmol	Joback Method

Temperature Dependent Properties

Property code	Value	Unit	Temperature [K]	Source
c _{pg}	897.59	J/mol×K	1027.56	Joback Method
c _{pg}	912.51	J/mol×K	1072.89	Joback Method
c _{pg}	927.03	J/mol×K	1118.21	Joback Method
c _{pg}	941.41	J/mol×K	1163.54	Joback Method
c _{pg}	955.93	J/mol×K	1208.87	Joback Method
c _{pg}	970.85	J/mol×K	1254.19	Joback Method
c _{pg}	986.44	J/mol×K	1299.52	Joback Method

Sources

- 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
- McGowan Method:** <http://link.springer.com/article/10.1007/BF02311772>
- NIST Webbook:** <http://webbook.nist.gov/cgi/cbook.cgi?ID=C93469&Units=SI>

Legend

cpg:	Ideal gas heat capacity
gf:	Standard Gibbs free energy of formation
hf:	Enthalpy of formation at standard conditions
hfus:	Enthalpy of fusion at standard conditions
hvac:	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
tb:	Normal Boiling Point Temperature
tc:	Critical Temperature
tf:	Normal melting (fusion) point
vc:	Critical Volume

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