# Benzenamine, N,N-dimethyl-4-[(3-methylphenyl)azo]-

Other names: Aniline, N,N-dimethyl-p-(m-tolylazo)-

**MDAB** 

N,N-Dimethyl-p-(m-tolylazo)aniline

3-Methyl-4'-(dimethylamino)azobenzene

3'-Mdab

3',N,N-Trimethyl-4-aminoazobenzene 4-Dimethylamino-3'-methylazobenzene 3'-Methyl-4-dimethylaminoazobenzene m'-Methyl-p-dimethylaminoazobenzene

3'-Me-dab

3'-Methyl-4-dimethylaminoazobenzen

3'-Methylbuttergelb

3'-Methyldimethylaminoazobenzol

3'Methyl-dab

4-(N,N-Dimethylamino)-3'-methylazobenzene Aniline, N,N-dimethyl-p-(3'-methylphenylazo)-3'-Methyl-N,N-dimethyl-4-aminoazobenzene 3'-Methyl-4-(N,N-dimethylamino)azobenzene

NSC 59783

N,N-dimethyl-4-(m-tolylazo)aniline

InChl=1S/C15H17N3/c1-12-5-4-6-14(11-12)17-16-13-7-9-15(10-8-13)18(2)3/h4-11H,1-3I

InchiKey: LVTFSVIRYMXRSR-UHFFFAOYSA-N

Formula: C15H17N3

**SMILES:** Cc1cccc(N=Nc2ccc(N(C)C)cc2)c1

Mol. weight [g/mol]: 239.32 CAS: 55-80-1

## **Physical Properties**

Property code	Value	Unit	Source
hf	211.94	kJ/mol	Joback Method
hvap	63.57	kJ/mol	Joback Method
log10ws	-3.90		Crippen Method
logp	4.476		Crippen Method
mcvol	200.330	ml/mol	McGowan Method
pc	1906.90	kPa	Joback Method
tb	767.56	K	Joback Method
tc	1017.23	K	Joback Method

### Sources

McGowan Method: http://link.springer.com/article/10.1007/BF02311772

NIST Webbook: http://webbook.nist.gov/cgi/cbook.cgi?ID=C55801&Units=SI

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

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

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

## Legend

hf: Enthalpy of formation at standard conditions

**hvap:** Enthalpy of vaporization at standard conditions

log10ws: Log10 of Water solubility in mol/llogp: Octanol/Water partition coefficientmcvol: McGowan's characteristic volume

pc: Critical Pressure

**tb:** Normal Boiling Point Temperature

tc: Critical Temperature

#### Latest version available from:

https://www.chemeo.com/cid/49-644-3/Benzenamine-N-N-dimethyl-4-3-methylphenyl-azo.pdf

Generated by Cheméo on 2024-04-10 00:22:18.531341549 +0000 UTC m=+14997787.451918861.

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