

2-Amino-2-phenylethanol, ferrocenylboronate

Inchi: InChI=1S/C13H13BNO.C5H5.Fe/c1-2-6-11(7-3-1)13-10-16-14(15-13)12-8-4-5-9-12;1-2-4
InchiKey: PFRXMCIPDPKKOR-UHFFFAOYSA-N
Formula: C18H18BFeNO
SMILES: c1ccc(C2COB(C34C5C6C7C3[Fe]6754389%10C4C3C8C9C4%10)N2)cc1
Mol. weight [g/mol]: 331.00

Physical Properties

Property code	Value	Unit	Source
rinpol	2480.00		NIST Webbook

Sources

NIST Webbook: <http://webbook.nist.gov/cgi/cbook.cgi?ID=R163112&Units=SI>

Legend

rinpol: Non-polar retention indices

Latest version available from:

<https://www.cheméo.com/cid/23-789-1/2-Amino-2-phenylethanol-ferrocenylboronate.pdf>

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