



CAS Registry Number: 1128-56-9
Molecular Formula: C₉H₉N₃
Molecular Weight: 159.19
Accurate Mass: 159.079647
Percentage Composition: C 67.91%; H 5.70%; N 26.40%
Melting Point: Mp 162-164° (as hydrochloride)
pKa Value: pK_a 2.66

>>Variant: 2H-form

Molecular Formula: C₃H₅N₃
Molecular Weight: 83.093
Accurate Mass: 83.048347
Percentage Composition: C 43.36%; H 6.06%; N 50.57%

>>Derivative: 2-Me
Synonym(s): 5-Amino-1-methylpyrazole

CAS Registry Number: 1192-21-8
Molecular Formula: C₄H₇N₃
Molecular Weight: 97.119
Accurate Mass: 97.063997
Percentage Composition: C 49.47%; H 7.26%; N 43.27%
pKa Value: pK_a 4.23 (25°, H₂O)

>>Derivative: 2-Me; hydrochloride

CAS Registry Number: 32016-34-5
Melting Point: Mp 141-143°

>>Derivative: 2-Me, picrate

CAS Registry Number: 1904-34-3
Physical Description: Large yellow prisms (MeOH)
Melting Point: Mp 221-224°

>>Derivative: 2-Ph
Synonym(s): 5-Amino-1-phenylpyrazole

CAS Registry Number: 826-85-7
Molecular Formula: C₉H₉N₃
Molecular Weight: 159.19
Accurate Mass: 159.079647
Percentage Composition: C 67.91%; H 5.70%; N 26.40%
Melting Point: Mp 182-185 dec.° (as hydrochloride)
pKa Value: pK_a 3.23

Rare Chemicals Library: S43807-3

References:

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