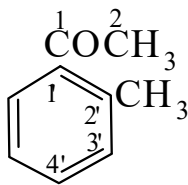




>>Entry Name: 2'-Methylacetophenone, 8Cl

Synonym(s): 1-(2-Methylphenyl)ethanone, 9Cl. Methyl *o*-tolyl ketone. *o*-Acetyloluene



CAS Registry Number: 577-16-2

Molecular Formula: C₉H₁₀O

Molecular Weight: 134.177

Accurate Mass: 134.073165

Percentage Composition: C 80.56%; H 7.51%; O 11.92%

Physical Description: Liq.

Boiling Point: Bp 214°. Bp₅ 79°

Density: d₄¹³1.02

Aldrich: M2659-3

Fluka: 65460

Sigma: M0145

>>Derivative: Semicarbazone

CAS Registry Number: 16882-28-3

Physical Description: Cryst. (EtOH)

Melting Point: Mp 218° (203-205°)

>>Derivative: 2,4-Dinitrophenylhydrazone

CAS Registry Number: 7287-83-4

Physical Description: Yellow cryst. (EtOH)

Melting Point: Mp 161°

References:

Aldrich Library of 13C and 1H FT NMR Spectra, 1992, **2**, 816B, (*nmr*)

Aldrich Library of FT-IR Spectra, 1st edn., 1985, **2**, 12D, (*ir*)

Aldrich Library of FT-IR Spectra: Vapor Phase, 1989, **3**, 1234B, (*ir*)

v. Auwers, K., *Annalen*, 1915, **408**, 242

Org. Synth., Coll. Vol., 1, 1932, 111

Jones, G., *J.C.S.*, 1960, 1918

Ghersetti, S. *et al.*, *Spectrosc. Lett.*, 1975, **8**, 391, (*ir*)

Shapiro, B.L. *et al.*, *J. Phys. Chem. Ref. Data*, 1977, **6**, 919, (*nmr*)