

THE TORSIONAL SPECTRUM OF MONO-DEUTERATED METHANOL CH₂DOH

A. EL HILALI, L. H. COUDERT, *LISA, CNRS/Universités Paris Est et Paris Diderot, 61 Avenue du Général de Gaulle, 94010 Créteil, France*; and S. KLEE, *Physikalisch-Chemisches Institut, Justus-Liebig-Universität Gießen, 35392 Gießen, Germany*.

Contrary to the isotopic specie of methanol with a symmetrical CH₃ or CD₃ methyl group, in the mono-deuterated species CH₂DOH the hindering potential no longer displays 3-fold symmetry and all moments of inertia depend on the angle of internal rotation.^a For this reason, this mono-deuterated species displays a fairly dense torsional spectrum difficult to compute and to assign.

In this paper an analysis of the torsional spectrum of mono-deuterated methanol is presented. More than 40 torsional subbands have been assigned in the 20 to 800 cm⁻¹ region. The body of data available for CH₂DOH consists of these newly assigned subbands and of the already available ones.^b The observed torsional subbands are characterized by $K \leq 10$ and $v_t \leq 10$. For 23 subbands, the rotational structure could be analyzed and this provides us with a mean to check their torsional assignment. The positions of the subband centers were analyzed with a theoretical approach accounting for the dependence of the inertia tensor on the angle of internal rotation. This allowed us to reproduce 58 experimental wavenumbers with an RMS value of 0.12 cm⁻¹. The spectroscopic parameters corresponding to the hindering potential and to the inertia tensor were also retrieved.

^aQuade and Lin, *J. Chem. Phys.* **38** (1963) 54.

^bLauvergnat, Coudert, Klee, and Smimov, *J. Mol. Spec.* **239** (2009) 204.