

A NUMERICAL-TENSORIAL “HYBRID” NUCLEAR-MOTION HAMILTONIAN AND DIPOLE MOMENT FOR SPECTRA CALCULATION OF POLYATOMIC NONRIGID MOLECULES

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The analysis and modelling of high-resolution spectra of nonrigid molecules require a specific Hamiltonian and group-theoretical formulation which differs significantly from that of more familiar rigid systems. Within the framework of the Hougen-Bunker-Johns (HBJ) theory¹, this work is devoted to the construction of a nonrigid Hamiltonian based on a suitable combination of numerical calculations for the nonrigid part in conjunction with the irreducible tensor operator method for the rigid part. A so-called “hybrid” model is thus introduced². For the first time, a variational calculation from *ab initio* potential energy surfaces is performed using the hybrid formulation of the exact HBJ kinetic energy operator expressed in terms of vibrational, large-amplitude motion and rotational tensor operators through the use of both curvilinear and rectilinear coordinates. Group theory for nonrigid molecules plays a central role in the characterization of the overall tunnelling splittings and will be discussed. The construction of the dipole moment operator will be also examined.

Validation tests consisting in a careful convergence study of the energy levels as well as in a comparison of results obtained from independent computer codes are given for the nonrigid molecules CH₂, CH₃, NH₃ and H₂O₂. This work—combined with our recent method for constructing effective models³—will pave the way to the analysis of high-resolution spectra of larger nonrigid systems.

This work is supported by the french Agence Nationale de la Recherche (ANR TEMMEX, Grant No. 21-CE30-0053-01) and the Russian Scientific Foundation (RSF, No. 22-42-09022).

¹J. T. Hougen, P. R. Bunker, J. W. C. Johns, *J. Mol. Spectrosc.*, **34** (1970) 136

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³M. Rey, *J. Chem. Phys.* **156** (2022) 224103. doi.org/10.1063/5.0089097