ANALYSIS AND FIT OF THE HIGH-RESOLUTION VISIBLE SPECTRUM OF THE TWO-EQUIVALENT-TOP MOLECULE BIACETYL

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The $A^1A_u(S_1) - X^1A_g(S_0)$ LIF spectrum of biacetyl (CH₃C(=O)C(=O)CH₃) shows a long progression in the torsional vibrations of the two methyl tops. The S_1 torsional level pattern can be qualitatively understood^{*a*} using local mode ideas applied to the two equivalent methyl rotors and symmetry ideas from the PI group G_{36} . For the present rotational analysis, we assigned a G_{36} symmetry species, two local-mode torsional quantum numbers, and rotational quantum numbers $J_{Ka,Kc}$ to each observed torsion-rotation level. Transitions were globally fit with a two-equivalent-top computer program, which uses the molecular principal axis system, a free-rotor basis set for each top, a symmetric-top basis set for the rotational functions, and a single-step diagonalization procedure. The program is similar to our two-inequivalent-top program for N-methylacetamide.^b Last year, we refit 179 previously published^c rotational lines involving 7 torsional tunneling sublevels with zero or one quantum of torsional excitation in the S_1 state of biacetyl, using 14 parameters (5 for S_0 , 9 for S_1) to obtain a standard deviation of 0.0041 cm⁻¹. We have now extended that fit by adding 131 unpublished lines involving 5 tunneling sublevels with two quanta of torsional excitation, using 19 parameters to obtain a standard deviation of 0.0040 cm⁻¹. Including transitions with still higher torsional excitation leads to a degraded standard deviation. We thus suspect that a number of higher-order terms describing top-top-overall-rotation interactions are missing from the present model, but the possibility of misassignments in the higher-energy regions must also be considered.

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