

ROTATIONAL SPECTROSCOPY OF THIOFORMALDEHYDE, H₂CS, IN ITS FOUR LOWEST EXCITED VIBRATIONAL, CORIOLIS-COUPLED STATES

HOLGER S. P. MÜLLER, CHRISTIAN P. ENDRES, STEPHAN SCHLEMMER, *I. Physikalisches Institut, Universität zu Köln, 50937 Köln, Germany*; ATSUKO MAEDA, IVAN MEDVEDEV, ERIC HERBST, MANFRED WINNEWISSER, FRANK C. De LUCIA, *Department of Physics, The Ohio State University, Columbus, OH 43210, USA*.

Rotational spectra of vibrationally excited states of H₂CS below 1500 cm⁻¹ have been obtained by long-path absorption spectroscopy between 120 and currently 670 GHz. The highest K_a accessed range from 9 to 12 and correspond to combined rotational and vibrational energies of almost 2400 cm⁻¹. The $v_4 = 1$ (990.185 cm⁻¹) and $v_6 = 1$ (991.019 cm⁻¹) states are almost degenerate and undergo very strong Coriolis interaction ($|\zeta_{46}^a| = 0.5148$) which causes $K_a > 0$ to be mixed completely. The $v_3 = 1$ state is at 1059.204 cm⁻¹ and the first order Coriolis terms are smaller ($|\zeta_{36}^c| = 0.2743$ and $|\zeta_{34}^b| = 0.0574$) so that perturbations are more local. Finally, the $v_2 = 1$ state (1455.496 cm⁻¹) is comparatively far away from the three lower states, but there is a large coupling term with $v_4 = 1$: $|\zeta_{24}^b| = 0.8555$. Because of the large energy difference, only higher order effects are pronounced in this case. The initial assignments were greatly facilitated by previous infrared studies on ν_4 , ν_6 , and ν_3^{ab} as well as on ν_2 .^c The results from these as well as previous and present pure rotational studies on the ground vibrational state were considered in the global fit. The very satisfactory fit required a modest number of vibrational corrections for the four vibrationally excited states and a comparatively large number of Coriolis interaction terms.

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