

AN INVESTIGATION OF EXCITED-STATE STRUCTURE AND DYNAMICS IN ACETYLACETONE THROUGH USE OF RESONANCE RAMAN SPECTROSCOPY

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Acetylacetone (AA), one of the simplest β -diketones, exhibits a strong intramolecular hydrogen bond that stabilizes the enol tautomer of the isolated (gas-phase) species and mediates the attendant proton-transfer process. Both experiment^a and theory^b have demonstrated conclusively that the \tilde{X}^1A_1 ground electronic state exhibits an asymmetrical equilibrium geometry with a potential barrier of finite height separating two equivalent conformers of C_s symmetry. In contrast, *ab initio* calculations^c have suggested that the electronically excited \tilde{B}^1B_2 ($\pi^*\pi$) manifold supports a symmetric (C_{2v}) minimum energy configuration which has the shuttling hydron located midway between the oxygen atom centers. This assertion, with its prediction of a low-barrier hydrogen-bonding motif, has been investigated experimentally by means of Resonance Raman Spectroscopy. Excitation at 266 nm, essentially coincident with the peak of the $\pi^* \leftarrow \pi$ transition, results in Raman profiles dominated by intense spectral features that stem from vibrational modes involving substantial distortion of the chelate ring, including marked displacement of the O \cdots O distance. Of special note is the 1620 – 2800 cm^{-1} region, which is not expected to contain any fundamental transitions, yet exhibits rich structure that has been assigned to overtone and combination bands. All of these data are consistent with a large change in molecular geometry upon electronic excitation. Resonance Raman spectra of deuterated derivatives and structural analogues of AA afford an additional means for unraveling the observed excited-state behavior. Ongoing extensions of these studies will be discussed, as well as efforts toward theoretical analysis based on the time-dependent formalism for Raman scattering.

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