

NEARLY FREE INTERNAL ROTATION IN PARATOLUALDEHYDE

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The rotational spectrum of paratolualdehyde ($\text{CH}_3\text{-C}_6\text{H}_4\text{-CHO}$) has been observed using a pulsed-molecular-beam Fourier-transform microwave spectrometer. The nearly free internal rotation of the methyl group splits each rotational transition into two components, A and E , separated by up to 2.5 GHz. The > 75 A -state a and b -type transitions fit an asymmetric-rotor Hamiltonian to better than 1 kHz to give $A=5128.6429(4)$ MHz $B=985.22385(3)$ MHz $C=827.28476(3)$. The inertial defect of $-0.61 \text{ u}\text{\AA}^2$ is significantly smaller than the expected rigid-molecule value of $-3.4 \text{ u}\text{\AA}^2$ and is consistent with nearly free internal rotation of the methyl top. The E state lines are strongly perturbed from a rigid-rotor pattern due to the low barrier to internal rotation of the methyl top. The potential barrier inhibiting the free rotation is approximated by $V(\alpha) = (1/2)V_3(1 - \cos 3\alpha) + (1/2)V_6(1 - \cos 6\alpha)$, where α is the internal rotation angle. For toluene ($\text{CH}_3\text{-C}_6\text{H}_5$) the V_3 term is zero by symmetry, while the V_6 term is approximately 5 cm^{-1} . In paratolualdehyde, the presence of the aldehyde group leads to a nonzero V_3 term due to a combination of long-range forces and the effect of the aldehyde substituent on the electronic structure of the benzene. Preliminary combined fits of the A and E state lines suggest that the V_3 and V_6 terms are of comparable magnitude. Searches are underway to observe the $|m| > 1$ internal rotor states to aid in the determination of the internal-rotation potential. Also, A -state transitions from the eight different ^{13}C isotopomers have been assigned to provide additional structural information.