

THE $\tilde{X}^2A_1 - \tilde{A}^2B_2$ CONICAL INTERSECTION IN NO_2 , OR HOW EXPERIMENTS AND THEORY INTERPLAY TOGETHER

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We present a comparison between experimental results (vibronic energies, absorption and LIDFS intensity ratios, rotational constants) and ab-initio calculations (diabatic and adiabatic energies, vibronic matrix elements). The three main inputs required in the analysis of the $\tilde{X}^2A_1 - \tilde{A}^2B_2$ conical intersection are: i) the diabatic levels of the \tilde{X}^2A_1 state (*i.e.*, the complete set of approximately 200 low lying levels of the \tilde{X}^2A_1 observed by LIDFS), ii) the diabatic levels of the \tilde{A}^2B_2 state (they are approximately predicted by the ab-initio calculations), iii) the matrix elements of the vibronic interaction, V_{12} , between the \tilde{X}^2A_1 and \tilde{A}^2B_2 electronic states. The validity of the simplified form proposed for V_{12} , namely “ λQ_3 ”, will be discussed. The comparison allows to assign some observed vibronic levels, which in return can be used to improve some parameters of the initial ab-initio PESurfaces. Up to now, our analysis is limited to the four lowest polyads of the \tilde{A}^2B_2 state ranging from 9700 cm^{-1} to 12300 cm^{-1} . At higher energy the interactions are stronger, leading to vibronic chaos above $\sim 17000\text{ cm}^{-1}$.