JET-COOLED CAVITY RING-DOWN SPECTROSCOPY OF THE $\tilde{A}^2 E'' - \tilde{X}^2 A'_2$ VIBRONIC TRANSITION OF NO₃

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The three energetically lowest electronic states $(\tilde{X} \ ^2A'_2, \ \tilde{A} \ ^2E'')$, and $\tilde{B} \ ^2E')$ of NO₃ are strongly coupled by vibronic interactions and have been treated in considerable detail theoretically.^b Corresponding experimental characterization of the interaction is much less detailed. Previous experimental results primarily consist of IR measurements of vibrational transitions in the ground state.^{cd} In addition, the electronically forbidden $\tilde{A} \cdot \tilde{X}$ transition has been observed in ambient temperature CRDS studies.^{ef} A slit-jet nozzle with a high voltage pulsed discharge has been applied to produce the NO₃ radical by dissociating the N-O bond of N₂O₅, and the jet-cooled NO₃ CRDS absorption spectrum has been successfully observed with a high-resolution laser source ($\Delta \nu \approx 250$ MHz, intrinsic resolution considering the instrumental linewidth and the residual Doppler broadening in the jet). The 4¹₀ band (parallel band) shows complex rotational structure which is presently being analyzed. The 2¹₀ band has also been measured as an example of a perpendicular band. Besides the ν_2 and ν_4 vibronic bands, the vibronically forbidden origin band (0⁰₀ band) has been recorded under the same experimental conditions. The weakly observed $\tilde{A} \cdot \tilde{X}$ origin band is likely either a magnetic dipole or an electric quardrupole transition.

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