

FTS AND ICLAS RECORDS OF PREDISSOCIATED ROVIBRATIONAL STRUCTURE NEAR 10400 CM⁻¹ FOR SYMMETRIC ISOTOPOMERS ¹⁶O₃ AND ¹⁸O₃ OF OZONE AND CONTOUR ANALYSIS

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This work is devoted to the study of low-energy quasi-bound states of the O₃ molecule. Previous analyses^{a,b,c} allowed to characterize a few vibrational levels of the first excited state ³A₂.

The absorption spectrum of ³A₂ in the region of its band 1₀⁰ 2₀² 3₀⁰ presents unassigned features. The theoretical profiles related to the probable connected transitions make difficult an unambiguous analysis of the spectrum at room temperature. To decrease the superpositions, absorption spectra at low temperature appeared necessary. Special cooled cells allowed us to record spectra by Fourier Transform Spectroscopy (FTS) and Intra Cavity Laser Absorption Spectroscopy (ICLAS). The predissociation broadening effects are important for these transitions and the rotational structure cannot be observed.

A comparison between theoretical and experimental profiles shows two or three types of transitions from the ground state \tilde{X} ¹A₁. i) One centred at 10360 cm⁻¹ corresponds to ³A₂ (0,0,1) \leftarrow \tilde{X} ¹A₁ (0,0,0). ii) The others in the 10480 cm⁻¹ region could be due to ³B₂ \leftarrow \tilde{X} ¹A₁ and/or ³B₁ \leftarrow \tilde{X} ¹A₁ and, simulations indicate that the upper level of the most intense absorption is probably ³B₂.

^aA.J. BOUVIER, G. WANNOUS, S. CHURASSY, R. BACIS, J. BRION, J. MALICET and R.H. JUDGE, Spectrochimica Acta (in press).

^bA.J. BOUVIER et al. Spectrochimica Acta, 55A, 2811 (1999)

^cA.J. BOUVIER et al. J. Mol. Spectrosc., 190, 189 (1998)