TORSIONAL, VIBRATIONAL AND VIBRATION-TORSIONAL LEVELS IN THE \mathbf{S}_1 AND GROUND CATIONIC \mathbf{D}_0^+ STATES OF PARA-XYLENE

ADRIAN M. GARDNER, WILLIAM DUNCAN TUTTLE, School of Chemistry, University of Nottingham, Nottingham, United Kingdom; PETER GRONER, Department of Chemistry, University of Missouri - Kansas City, Kansas City, MO, USA; TIMOTHY G. WRIGHT, School of Chemistry, University of Nottingham, Nottingham, United Kingdom.

Insight gained from examining the "pure" torsional, vibrational and vibration-torsional (vibtor) levels of the single rotor molecules: toluene (methylbenzene)^a and para-fluorotoluene (pFT), b is applied to the double rotor para-xylene (p-dimethylbenzene) molecule . c Resonance-enhanced multiphoton ionization (REMPI) spectroscopy and zero-kinetic-energy (ZEKE) spectroscopy are employed in order to investigate the S_1 and ground cationic states of para-xylene. Observed transitions are assigned in the full molecular symmetry group (G_{72}) for the first time.

^aJ. R. Gascooke, E. A. Virgo, and W. D. Lawrance, J. Chem. Phys., 143, 044313 (2015).

^bA. M. Gardner, W. D. Tuttle, L. Whalley, A. Claydon, J. H. Carter and T. G. Wright, J. Chem. Phys., 145, 124307 (2016).

^cA. M. Gardner, W. D. Tuttle, P. Groner and T. G. Wright, *J. Chem. Phys.*, (2017, in press).