

SPECTROSCOPY AND STRUCTURE OF CHAINS TYPE C_n , C_nH and C_nSi

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Structures, vertical excitation energies to the first electronic states and spectroscopic parameters for the ground electronic state are determined for several small C_n , C_nH and C_nSi astrophysically important molecular species ^a. The main aim of these investigations is to help the interpretation of several previous astrophysical observations ^b. State-of-the art ab initio calculations are performed for this purpose. All species display isomerism although the inter-transformation processes, that involve various electronic states, are not well known. Anharmonic spectroscopic parameters are computed from full-dimensional Potential Energy Surfaces which are mapped close to their respective local minima. In light of these calculated properties (band positions, spectroscopic parameters, predictions of ro-vibronic effects, spin-spin constants,...) assignments for FIR astrophysical observed bands, are discussed. vspace1em

^aH. Massó, M.L. Senent, P. Rosmus and M. Hochlaf, *J.Chem.Phys.* **124**, 234304 (2006); M.L. Senent H. Massó and M. Hochlaf, *ApJ*, **670**, 1510 (2007); H. Massó, V. Veryazov, P.A. Malmqvist, B.O. Roos and M.L. Senent, *J.Chem.Phys.* **127**, 154318 (2007); M. Hochlaf, C. Nicolas and L. Poisson, *J.Chem.Phys.* **127**, 014310 (2007) N. Inostroza, M. Hochlaf, M.L. Senent and J.R. Letelier (submitted 2008)

^bJ. Goicoechea, J. Cernicharo, H. Massó and M.L. Senent, *ApJ*, **609**, 225 (2004); J. Cernicharo, J.R. Goicoechea and Y. Benilan, *ApJ* **580**, L157, 2002.