

Conformational analysis and vibrational assignments of 2,2,3,3-tetrafluoro-1-propanol CHF₂CF₂CH₂OH as three-rotors system. Badawi, Hassan M.; Foerner, Wolfgang

Abstract

The conformational stability of 2,2,3,3-tetrafluoro-1-propanol was investigated by DFT-B3LYP/6-311+G** and ab initio MP2/6-311+G** calcns. The calcd. potential energy curves of the mol. at DFT-B3LYP level were consistent with five distinct min. that correspond to gauche(-)-gauche-gauche (G1gg), trans-trans-gauche (Ttg), trans-gauche-gauche (Tgg), trans-gauche-gauche(-) (Tgg1) and gauche(-)-gauche-trans (G1gt) conformers in the order of decreasing relative stability. The equil. consts. for the conformational interconversion of 2,2,3,3-tetrafluoro-1-propanol were calcd. and found to correspond to an equil. mixt. of about 38% G1gg, 28% Ttg, 13% Tgg, 11% Tgg1 and 10% G1gt conformations at 298.15 K. The vibrational frequencies of 2,2,3,3-tetrafluoro-1-propanol in its five stable forms were computed at B3LYP level and complete vibrational assignments were made based on normal coordinate calcns. and comparison with exptl. data of the mol.