

PSEUDO JAHN-TELLER ORIGIN OF THE PROTON-TRANSFER ENERGY BARRIER IN THE HYDROGEN-BONDED [FHF]⁻ SYSTEM

Natalia Gorinchoy^{a*}, Iolanta Balan^a, Victor Polinger^b, Isaak Bersuker^{a,c}

^aInstitute of Chemistry, 3, Academiei str., Chisinau MD-2028, Republic of Moldova

^bUniversity of Washington, 109, Bagley Hall Box 351700 Seattle, WA 98195-1700, USA

^cUniversity of Texas at Austin, Austin, Texas 78712, USA

*email: ngorinchoy@yahoo.com; natalia.gorincioi@ichem.md

Abstract. The results of *ab initio* calculations of the adiabatic potential energy surfaces for the proton-bound [FHF]⁻ system at different F-F distances have been rationalized in the framework of the vibronic theory. It is shown that the instability of the symmetric D_{σ_h} structure at increased F...F distances and the proton displacement to one of the fluorine atoms are due to the pseudo Jahn–Teller mixing of the ground electronic state $1^1\Sigma_g$ with the lowest excited state of $1^1\Sigma_u$ symmetry through the asymmetric σ_u vibrational mode.

Keywords: proton transfer, hydrogen bond, pseudo Jahn–Teller effect, potential energy surface, bifluoride anion.

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