DEUTERATED PROPANAL FIR SPECTRUM

M. VILLA, Departamento de Química (CBI, UAM-I), Av. San Rafael Atlixco 186, Col. Vicentina, Iztapalapa 09340, Mexico D.F., MEXICO; M. L. SENENT, Instituto de Estructura de la Materia (C.S.I.C.), Serrano 113bis, Madrid, SPAIN.

In previous work, the zero point energy correction has been applied to methyamine and propanal in order to reproduce more properly their FIR spectrum. In this work it will be shown the zero point energy correction applied to different deuterated species of propanal, and the FIR spectrum obtained experimentally is going to be compared with experimental data reported in the literature. It is important to emphasize that the zero point energy correction method that has been used is a parametrized one, in order to improve the HF/gaussian 98 energy and frequency calculations that have been obtained.