## A FIRST PRINCIPLE EFFECTIVE HAMILTONIAN FOR INCLUDING NON-ADIABATIC EFFECTS FOR $\mathrm{H_2^+}$ AND $\mathrm{HD^+}$

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We compute non-adiabatic corrections for all bound and long lived quasi-bound vibrational levels of  $H_2^+$  and  $HD^+$  for selected rotational levels. This is done using the Bunker and Moss formalism with the correction factors computed from *ab initio* wavefunctions. The electronic wave functions are expanded in terms of nuclear centered gaussian basis functions. The agreement with accurate calculations is very good: for  $H_2^+$ , most transition frequencies are predicted to within about 0.0001 cm<sup>-1</sup>. For HD<sup>+</sup>, the results are not quite as good due to the uncertainties in the adiabatic correction. This paves the way for using these techniques to accurately predict the non-adiabatic effects for more complicated molecules.