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Erratum: Determination of Structure Parameters in Strong-Field Tunneling Ionization Theory of Molecules (Physical Review a (2010) 81 (033423))

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There are several errors in Tables V and VI of our article. In Table V, the C_{2m} of the HOMO-1 (1π) of CO molecule should be 0.014. In Table VI, the binding energies of $2p\pi_g$ and $2p\pi_u$ of H_2^+ should be 0.2267 and 0.4288, respectively.