## Computer predicted ionization energy of carbon accurate to within $\pm 1~{\rm cm}^{-1}$

Nike Dattani\*
Harvard Smithsonian Center for Astrophysics, 02138, Cambridge, MA, USA.

## Giovanni LiManni<sup>†</sup>

Department of Electronic Structure Theory, Max Planck Institute for Solid State Systems, Suttgart, Germany.

## David Feller<sup>‡</sup>

Department of Chemistry, Washington State University, Pullman, Washington 99164-4630, USA,

## Jacek Koput§

Department of Chemistry, Adam Mickiewicz University, 60-780 Poznan, Poland. (Dated: 24th November 2018)

We show that we can predict the first ionization energy in carbon to within  $1.076~\rm cm^{-1}$  of the experimental value. This is an improvement of more than a factor of 5 over the preceding best prediction in [Phys. Rev. A 81, 022503] and comparable to the accuracies achieved for the smaller atoms up to boron. The sum of all error contributions in our calculations up to the fifth power of the fine structure constant is estimated to be at most  $0.806~\rm cm^{-1}$ , which yields our estimate of contributions in the sixth and higher powers to be:  $1.076 \pm 0.806~\rm cm^{-1}$ .

In the last seven years, ionization energies (IEs) have been calculated with unprecedented precision for the Li atom [?], Be atom [?] and B atom [?]. Tight variational bounds for non-relativistic ground state energies assuming a clamped, point-sized nucleus have reached 49 digits in units of Hartree for He, 19 digits for Li, 12 digits for Be, and 11 digits for B (see Table ??). Calculated IEs have been made in agreement with experiment to within  $10^{-3}$  cm<sup>-1</sup> for <sup>7</sup>Li,  $10^{-1}$  cm<sup>-1</sup> for <sup>9</sup>Be and 1 cm<sup>-1</sup> for <sup>11</sup>B (see Table ??).

For the C atom, before this present study, no high-precision calculation had been reported to predict an IE to  $\sim 1~{\rm cm^{-1}}$  agreement with experiment. Table ?? shows that the method used for the smaller atoms up to boron has not had success for carbon. With twice as many variationally optimizable parameters, 1 fewer digit was obtained for the B atom than for the Be atom, which also suggests that it would be very difficult to variationally optimize a fully explicitly correlated wavefunction ansatz for atoms and molecules coming from most of the periodic table.

The best known variational bound for the NR,CPN ground state energy for C was calculated in 2015 using fixed-node diffusion Monte Carlo (FN-DMC) with the nodes of the electronic wavefunction fixed at the locations of a CISD/cc-pV5Z wavefunction, and the statistical uncertainty on the Monte Carlo was  $\pm 20~\mu E_{\rm Hartree}$  [?]. However, the IE for C predicted by FN-DMC was in discrepancy with experiment by more than  $40~{\rm cm}^{-1}$ . In this paper, the approach we use to calculate the NR,CPN

energy of the ground state of C is FCIQMC (full configuration interaction quantum Monte Carlo) in an augcc-pCV8Z basis set. Table ?? shows that our NR,CPN energy is at least 76  $\mu E_h$  higher than the variational upper bound obtained from FN-DMC; but since in our approach, imperfections in the description of the wavefunction for the neutral atom are almost the same as in the cation, the errors are almost completely eliminated when taking the energy difference, therefore bringing us to agreement with experiment that is comparable to what has been seen with the explicitly correlated approach for atoms as big as (but not exceeding) boron.

After adding relativistic and quantum electrodynamics (QED) corrections, and corrections to the clamped nucleus approximation, we obtained an IE for the ground sate of C which is within 1.076 cm<sup>-1</sup> disagreement with the best known experimental estimate. While this is not as impressive as the method of variationally optimizing parameters in an explicitly correlated wavefunction ansatz has proven to be for Li and Be, the disagreement has the same order of magnitude as the latter approach for B (see Table ??). We finally note that the approach used in this paper, of calculating FCIQMC on a basis set of non-explicitly correlated orbitals has successfully treated systems with far more electrons (transition metal atoms [?], diatomics [?], larger molecules such as butadiene [?], and even solid state systems [?]), so it is conceivable that the approach used in this paper may in the near future be able to determine (with fair accuracy) the IEs which at present remain experimentally elusive or poorly known. These include arsenic (whose experimental IE has an uncertainty of  $\pm 2 \,\mathrm{cm}^{-1}$ ), Pm, Pa, Fm, Md, No, Sg, Bh and Hs (whose IEs are only known based on extrapolations of other experimental data and have uncertainties between  $\pm 140\,\mathrm{cm}^{-1}$  and  $\pm 4000\,\mathrm{cm}^{-1}$ ), Rf and Db (whose IEs are only known from theoretical calculations), and Mt, Ds, Rg, Cn, Nh, Fl, Mc, Lv, Ts,

<sup>\*</sup> ndattani@cfa.harvard.edu

<sup>†</sup> g.limanni@fkf.mpg.de

<sup>‡</sup> dfeller@owt.com

<sup>§</sup> koput@amu.edu.pl