Improved Monte Carlo estimators for the one-body density

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An alternative Monte Carlo estimator for the one-body density \( \rho(\mathbf{r}) \) is presented. This estimator has a simple form and can be readily used in any type of Monte Carlo simulation. Comparisons with the usual regularization of the delta-function on a grid show that the statistical errors are greatly reduced. Furthermore, our expression allows accurate calculations of the density at any point in space, even in the regions never visited during the Monte Carlo simulation. The method is illustrated with the computation of accurate variational Monte Carlo electronic densities for the Helium atom (one-dimensional curve) and for the water dimer (three-dimensional grid containing up to 51 × 51 × 51 = 132 651 points).

The Monte Carlo approach is probably one of the most widely employed numerical approaches in the scientific and engineering community. In computational physics, it has been extensively used in the last 50 years for studying a great variety of many-body systems under many different conditions. To date, the most popular application of the method is probably the calculation of classical thermodynamical properties [1]. However, the Monte Carlo approach is also employed for evaluating quantum properties by using the path-integral formulation of quantum averages as classical ones (quantum Monte Carlo or path integral Monte Carlo approaches [2]). In the recent years, these later approaches have emerged as a unique and powerful tool for studying quantitatively the interplay between quantum and thermal effects in many-body systems (e.g., to understand the very rich physics of strongly correlated materials).

At the heart of all these applications lies the calculation of a number of high-dimensional integrals (or sums, for lattice problems) written under the following general form:

\[
I(\mathbf{r}) = \int d\mathbf{r}_1 \cdots d\mathbf{r}_N \Pi(\mathbf{r}_1, \ldots, \mathbf{r}_N) F(\mathbf{r}_1, \ldots, \mathbf{r}_N),
\]

where \( \Pi \) is some arbitrary \( N \)-body probability distribution (\( \Pi \) positive and normalized) and \( F \) some arbitrary real-valued function. The integration is performed over all accessible configurations for the \( N \)-particle system. The general idea of Monte Carlo approaches is to evaluate the integral by sampling the configuration space according to the probability distribution, \( \Pi \), and by averaging \( F \) over the various configurations generated by the sampling procedure, \( I(\mathbf{r}) = \langle F \rangle_{\Pi} \). Here and in what follows, the symbol \( \langle \cdots \rangle_{\Pi} \) indicates the statistical average over the density \( \Pi \). Various Monte Carlo algorithms (sampling procedures) can be found in the literature, the most celebrated one being, of course, the Metropolis algorithm [3]. The efficiency of a Monte Carlo approach is directly related to the magnitude of the fluctuations of the integrand in the regions where the probability distribution, \( \Pi \), is large. More precisely, for a given number of Monte Carlo steps, the statistical error \( \delta F \) is proportional to the square root of the variance of the integrand \( \tilde{F} \) defined as \( \sigma^2(\tilde{F}) = \langle \tilde{F}^2 \rangle - \langle \tilde{F} \rangle^2 \). Accordingly, a very attractive way of enhancing the convergence of a Monte Carlo simulation consists in introducing alternative “improved” estimators defined as new integrands \( \tilde{F} \) having the same average as \( F \) but a lower variance

\[
\langle \tilde{F} \rangle_{\Pi} = \langle F \rangle_{\Pi} \quad \text{and} \quad \sigma^2(\tilde{F}) < \sigma^2(F).
\]

In previous works [4,5] it has been shown how improved estimators can be designed for any type of integrand \( F \) and Monte Carlo algorithm, and some applications to the computation of forces have been presented [5].

In this Rapid Communication we present an efficient improved Monte Carlo estimator for calculating the one-body (or one-particle) density, \( \rho(\mathbf{r}) \).

\[
\rho(\mathbf{r}) = \left\langle \sum_{i=1}^{N} \delta(\mathbf{r}_i - \mathbf{r}) \right\rangle_{\Pi}
\]

and, more generally, any one-body average of the form \( \int d\mathbf{r} \delta(\mathbf{r}) F(\mathbf{r}) \). As we shall see, our estimator allows very important reductions in variance. In the example of the charge density of the water dimer presented below, a reduction of up to two orders of magnitude in CPU time is possible for some regions of space. In addition, and in sharp contrast with the usual estimator based on the regularization of the delta-function on a grid, our expression leads to accurate estimates of the density at any point in space, even in the regions never visited during the Monte Carlo simulation (e.g., in the large-distance regime). This property is particularly interesting when a global knowledge of the density map is searched for. For the water dimer case, we were able to accurately compute the charge density for 51 × 51 × 51 = 132 651 grid points. Note that such a calculation is vastly more difficult to perform when using the standard approach.

Let us recall that accurate one-particle properties are of central interest for the understanding of the physics of many
complex many-body systems. Such systems include all those which are not translationally invariant (typically, all finite systems: atoms, molecules, clusters, nuclei, etc.) and all those whose translational symmetry has been explicitly broken, e.g., by the application of an inhomogeneous external field. In addition to this, many physical modelizations and/or effective theories rely explicitly on the knowledge of the one-body density. Many examples could be cited but let us mention, for example, the various modelings of the electro-static field of molecules from the one-electron density [6], the studies of the structure and reactivity of molecular systems based on the topological analysis of the electron density and/or its Laplacian [7], and, also, the very important case of density functional theories (DFT) which could greatly benefit from the possibility of computing accurate three-dimensional (3D) charge/spin density maps for large molecular systems (e.g., via accurate fits of the exchange-correlation Kohn-Sham potential, see [8]).

Finally, let us note that the use of alternative forms for evaluating the density is not new. For example, in the works of Hiller et al. [9], Sucher and Drachman [10], Hariman [11], and Rassolov and Chipman [12] new classes of global operators built for computing the density have been introduced. However, in these works, the general idea is to design operators whose expectation values give an accurate estimate of the unknown exact one-body density and not the exact one-body density associated with a known N-body density. Actually, our strategy is more closely related to what has been presented by Vrbik et al. [13], Langfelder et al. [14], and Alexander and Coldwell [15]. In these works, alternative Monte Carlo estimators with lower variances are also introduced. However, the emphasis is only put on the case of evaluating the charge and/or spin density at the nuclei. Here, such ideas are extended to any point in space and a general formula allowing one to control all possible sources of statistical fluctuations in all possible regimes is presented.

**General improved density estimator.** Due to the presence of Dirac functions in the Monte Carlo estimator of the density, Eq. (3), some sort of regularization has to be introduced. It is usually done by partitioning the physically relevant part of the one-particle space (usually, the 3D ordinary space) into small domains of finite volume and by evaluating the corresponding locally averaged densities. In practice, such a procedure is particularly simple to implement by counting the number of particles present in each elementary domain at each step of the simulation. However, the statistical fluctuations can be rather large. This is particularly true for the low-density regions which are rarely visited by the particles. Even worse, there is no way of evaluating the density in regions which are never visited during the finite Monte Carlo simulation. One way to escape from these difficulties is to introduce some global estimators defined in the whole space. To do that, we regularize the Dirac-function by using the following equality:

\[
\delta(r_i - r) = \frac{-f(r_i; r)}{4\pi} \nabla_i^2 \frac{1}{|r_i - r|},
\]

where \(f(r_i; r)\) is a smooth function of \(r_i\) verifying \(f(r_i = r; r) = 1\). Here, \(\nabla_i^2\) denotes the Laplacian acting on the \(r_i\) variable and \(r\) the fixed position where the density is computed. Note that this formula is just a slightly generalized form of the well-known equality corresponding to \(f = 1\). Now, injecting this expression into Eq. (3) and integrating by parts, the density can be rewritten as

\[
\rho(r) = -\frac{1}{4\pi} \sum_{i=1}^{N} \left\langle \frac{1}{|r_i - r|} - f(r_i; r) \nabla_i^2 \frac{1}{|r_i - r|} \right\rangle_{\Pi}, \tag{5}
\]

Next, we introduce some additional function \(g(r)\) (independent on the particle coordinates) and write \(\rho(r)\) as

\[
\rho(r) = -\frac{1}{4\pi} \sum_{i=1}^{N} \left\langle \frac{1}{|r_i - r|} - g(r) \nabla_i^2 \frac{1}{|r_i - r|} \right\rangle_{\Pi} \tag{6}
\]

the last step being allowed since \(\left\langle \nabla_i^2 f(r_i; r) \right\rangle_{\Pi} = 0\).

Expression (6) is our general form for the improved estimator of the one-body density. The two functions \(f(r_i; r)\) and \(g(r)\) play the role of auxiliary quantities. They are introduced to decrease the variance of the density estimator. As with any optimization problem, there is no universal strategy for choosing \(f\) and \(g\). However, the guiding principle is to identify the leading sources of fluctuations and, then, to adjust the auxiliary functions to remove most of them.

**Improved electronic density estimator for molecules.** In what follows, we consider the one-electron density of molecular systems obtained from the N-body quantum probability distribution written as

\[
\Pi(r_1, \ldots, r_N) = \frac{\psi_f^2(r_1, \ldots, r_N)}{\int dr_1 \cdots dr_N \psi_f^2(r_1, \ldots, r_N)}, \tag{7}
\]

where \(\psi_f\) is some electronic trial wave function.

1. **Short electron-nucleus distance regime.** For a system of electrons in Coulombic interaction with a set of fixed nuclei the exact wave function is known to obey the following electron-nuclear cusp condition:

\[
\psi_{r \rightarrow r_A} \rightarrow 1 = Z_A |r_i - R_A|, \tag{8}
\]

where \(R_A\) denotes the position of a given nucleus \(A\) of charge \(Z_A\). Most of the accurate trial wave functions employed in the literature fulfill this important condition. Now, as a consequence of the cusp condition we have

\[
\frac{\nabla_i^2 \Pi}{\Pi} \rightarrow _{r \rightarrow R_A} \frac{4Z_A}{|r_i - R_A|}. \tag{9}
\]

In the neighborhood of nucleus \(A\), this term is an important source of fluctuations. This is easily seen by noting that, in the regime \(r \sim R_A\), the estimator of \(\rho(r)\), Eq. (6), behaves as \(\frac{1}{|r - R_A|}\), a quantity which has an infinite variance \(\int [\frac{1}{|r - R_A|}]^2 dr = + \infty\). To remove this source of wild fluctuations we adjust the function \(f\) so that \(\nabla_i f\) exactly cancels the divergence of \(\frac{\nabla_i^2 \Pi}{\Pi}\). A simple suitable form for \(f\) obeying such a condition plus the constraint, \(f(r_i = r; r) = 1\), is given by
2. Large-distance regime. In the large-distance regime, \(|r| \rightarrow +\infty\), the exact one-electron density is known to decay exponentially. In contrast, our basic estimator decays algebraically as a function of \(r\). Here, we propose to force the latter estimator to decay also exponentially. In practice, a simple choice for the function \(f\) is

\[
f(r_{1}; r) = (1 + \lambda |r_{1} - r|) \exp[-\lambda |r_{1} - r|],
\]

where \(\lambda\) is some real parameter. Note that the coefficients of the linear and exponential terms have been taken identical \((=\lambda)\) to avoid the divergence of the Laplacian of \(f\) at \(r_{1} = r\). The parameter \(\lambda\) can be adjusted either by minimizing the fluctuations of the average density or fixed at some value close to the theoretical value of \(\lambda_{sl} = 2\sqrt{2}\) \((f\) first ionization potential, the exact density is known to decay as \(\rho \sim \exp(-\lambda_{sl} r)\) see, e.g., Ref. [16]).

Another useful remark is that, in the regime \(|r| \rightarrow +\infty\), the electrons of the molecule are, in first approximation, at the same distance from the point where \(\rho\) is evaluated. As a consequence, during the simulation the quantity \(1/|r_{1} - r|\) fluctuates very little around the approximate average \(1/\langle |r_{1} - r| \rangle\) Therefore to reduce the fluctuations it is valuable to remove a quantity close to the latter average from the former one. Here, this idea is implemented by introducing the following function \(g\):

\[
g(r) = \frac{1}{M} \sum_{A=1}^{M} \frac{1}{|R_{A} - r|}.
\]

In our first application we consider the He atom described by a simple trial wave function written as \(\psi_{f} = \phi_{1}(r_{1})\phi_{1}(r_{2})\) with \(\phi = \exp(-\gamma r)\), and \(\gamma = 1.6875\) (Slater value). For this problem, the exact density is known and is given by \(\rho(r) = 2\gamma^{2}/\pi \exp(-2\gamma r)\). Figure 1 shows the results obtained for \(\rho(r)\) for a relatively short Monte Carlo run. The main curve displays the results obtained with (i) the usual estimator based on the delta representation, Eq. (3), and (ii) the simple improved estimator corresponding to Eq. (6) with \(f=1\) and \(g=0\), and our best improved estimator defined via Eqs. (6) and (10)–(12). In the latter case, the densities corresponding to each of the two possible choices for \(f\) [Eqs. (10) or (11)] have been computed for each distance, the final value corresponding to the value having the smallest statistical error. The usual estimator, Eq. (3), has been regularized by introducing small elementary cubes of length \(a=0.2\). For all distances the statistical error associated with the usual estimator is very large with respect to improved estimators. At intermediate distances, at least one order of magnitude in accuracy is lost. For example, at \(r=0.6\) the statistical error is about ten times larger than for the simple estimator case and a factor of about 20 is found with respect to the best improved estimator. At large distances, a region rarely visited by the electrons, the standard estimator is so noisy that it is useless in practice. Now, regarding improved estimators it is clear that the auxiliary functions \(f\) and \(g\) have a great impact on reducing the errors. At very small distances (first inset) a gain of about 5 in statistical error is obtained with the best improved estimator. At \(r=0\), the gain is even larger since the simple estimator has an infinite variance. At large distances where both improved estimators have a finite variance, it is seen that introducing some exponential decay into the estimator plus a proper shift \((g\)-contribution) improves considerably the convergence. In the range 2.5–3, the gain in error increases from 15 at \(r=2.5\) to 40 at \(r=3\).

In our second application we consider the water dimer in a nonsymmetric nuclear geometry (structure #2 of Ref. [17]) described by an electronic wave function consisting of a Hartree-Fock part (cc-pVTZ basis set) plus a standard explicitly correlated Jastrow term. Figure 2 shows the density plots obtained with our best improved estimator for a number of points equal to \(51 \times 51 \times 51\) (=132,651). As seen in the figure the density obtained displays a very smooth aspect. A closer look shows that this regularity is present at a rather small scale. In Fig. 3 we present a more quantitative comparison of the data along the O-O axis. The figure clearly shows that the best estimator outperforms the usual one. First, the curve corresponding to the new estimator (solid
line connecting the points) is very smooth, although it has been obtained by simple linear extrapolation of the data. In sharp contrast, this is absolutely not true for the usual estimator curve whose overall behavior is particularly chaotic. Second, the statistical error has been greatly reduced using the new estimator. Depending on the distance, a gain in accuracy ranging roughly from 5 to 10 (i.e., up to two orders of magnitude in CPU time) has been obtained. An interesting point to mention is the presence of some very wild fluctuations in the neighborhood of \( r_{\text{O-O}} \sim 1.5 \) for the standard estimator. These fluctuations are due to the presence of a hydrogen atom close to the O-O axis. We can verify that, in sharp contrast, our estimator, which has been built to correctly take into account the nuclear cusp, Eq. (10), performs well in that region. Remark that in the large-\( r \) regime (data not shown here) where the standard estimator is strictly zero (no sampling of this region), the improved estimator still continues to give accurate values of the very small density.

Finally, let us make some comments about the additional computational cost associated with the use of our improved estimator instead of the very simple standard one, Eq. (3). Remark first that the calculation of \( \Pi \) and its first and second derivatives is done only once at each Monte Carlo step for all of the grid. The cost for evaluating these quantities scales as \( \sim n_{\text{part}}^2 \), where \( n_{\text{part}} \) is the number of particles and \( \gamma \) is typically 2 for bosons (two-body interaction) and 3 for fermions (the computation of determinants is the dominant part). Note that this scaling is not problematic since it is the scaling of the Metropolis step itself. On the other hand, the evaluation of the \( r \)-dependent quantities is much more simple [see, expression of \( f \), Eqs. (10) and (11)], and the corresponding CPU cost scales as \( \sim n_{\text{grid}}^2 n_{\text{part}} \). This cost is negligible for a large number of particles \( n_{\text{part}} \gg n_{\text{grid}} \) regime but not for small and intermediate systems (such as the water dimer). In these latter cases, different ways are at our disposal to reduce the CPU cost. We shall not enter into the details here, let us just mention the use of grids adapted to the density profile (less points are needed), the possibility of avoiding the computation of the improved estimator at each Monte Carlo step (Monte Carlo configurations are highly correlated) and, also, exploiting the fact that the contribution to the density of a given particle is exponentially small for large grid point-particle distances, a property which can be useful for limiting the amount of calculations. In conclusion, the additional computational cost associated with our proposed estimator can be easily kept under control and, thus, is not a limitation of the approach. In the water dimer case presented above, a naive Cartesian grid of 132 651 points has been used and the improved estimator has been computed for each 15 Monte Carlo steps. We found a CPU-time increase with respect to the usual estimator of about 6.

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