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Quantum Monte Carlo method for attractive Coulomb potentials

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Starting from an exact lower bound on the imaginary-time propagator, we present a path-integral quantum Monte Carlo method that can handle singular attractive potentials. We illustrate the basic ideas of this quantum Monte Carlo algorithm by simulating the ground state of hydrogen and helium.

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I. INTRODUCTION

Quantum Monte Carlo (QMC) simulation is a powerful method for computing the ground state and nonzero temperature properties of quantum many-body systems [1,2]. There are two fundamental problems that limit the application of these methods. The first and most important is the minus-sign problem, which we do not address in this paper (see, however, [3,4]). The second problem arises if one wants to simulate systems with attractive singular potentials, the Coulomb interaction being the prime example. The purpose of this paper is to present an approach that solves the latter problem in a form that fits rather naturally in the standard path integral QMC (PIQMC) approach and leaves a lot of room for further systematic improvements.

Let us first recapitulate the basic steps of the procedure to set up a PIQMC simulation. Writing \( K \) and \( V \) for the kinetic and potential energy, respectively, the first step is to approximate the imaginary-time propagator by a product of short-time imaginary-time propagators. The standard approach is to invoke the Trotter-Suzuki formula [5,6],

\[
e^{-\beta(K+V)} = \lim_{m \to \infty} (e^{-\beta K/m} e^{-\beta V/m})^m, \tag{1}
\]

to construct a sequence of systematic approximations \( Z_m \) to the partition function \( Z \) [6,7],

\[
Z = \text{Tr} \exp(-\beta H) = \lim_{m \to \infty} Z_m \tag{2}
\]

\[
Z_m = \int dr_1 \cdots dr_m \prod_{n=1}^m \langle r_n | e^{-\beta K/m} | r_{n+1} \rangle e^{-\beta V(r_{n+1}/m)}, \tag{3}
\]

where \( r_{m+1} = r_1 \) and use has been made of the fact that the potential energy is diagonal in the coordinate representation. Taking the limit \( m \to \infty \), Eq. (3) yields the Feynman path integral [8] for a system with Hamiltonian \( H = K + V \). Expression (3) is the starting point for the PIQMC simulation.

In the case of the attractive Coulomb interaction, it is easy to see why the standard PIQMC approach fails. Let us take the hydrogen atom as an example. The Hamiltonian reads

\[
H = -\frac{\hbar^2}{2M} \nabla^2 - \frac{q^2}{r}, \tag{4}
\]

where \( q \) denotes the charge of the electron and \( M = m_e/(1 + m_e/m_p) \), \( m_e(m_p) \) being the mass of the electron (proton). Replacing the imaginary-time free-particle propagator in Eq. (3) by its explicit, exact expression,

\[
\langle r | e^{-\beta K/m} | r' \rangle = \left( \frac{mM}{2 \pi \beta \hbar^2} \right)^{3/2} \exp \left( -\frac{mM|x-x'|^2}{2\beta \hbar^2} \right), \tag{5}
\]

we obtain

\[
Z_m = \left( \frac{mM}{2 \pi \beta \hbar^2} \right)^{3m/2} \int dr_1 \cdots dr_m \\
\times \exp \left[ -\frac{mM}{2\beta \hbar^2} \sum_{n=1}^m (r_n - r_{n+1})^2 \right] \\
\times \exp \left[ +\frac{\beta q^2}{m} \sum_{n=1}^m \frac{1}{r_n} \right]. \tag{6}
\]

PIQMC calculates the ratio of integrals such as Eq. (6) by using a Monte Carlo procedure to generate the coordinates \( \{r_1, \ldots, r_m\} \). The integrand in Eq. (6) serves as the weight for the importance sampling process. As the latter tends to maximize the integrand, it is clear that because of the factors \( \exp(+\beta q^2 m^{-1} r_n^{-1}) \), the points \( \{r_1, \ldots, r_m\} \) will, after a few steps, end up very close to the origin. In the case of a singular, attractive potential, importance sampling based on Eq. (6) fails. Using instead of the simplest Trotter-Suzuki formula (1) a more sophisticated one [9] only makes things worse because these hybrid product formulas contain derivatives of the potential with respect to the coordinates.

The problem encountered in setting up a PIQMC scheme for models with a singular, attractive potential is just a signature of the fundamental difficulties that arise when one tries to define the Feynman path integral for the hydrogen atom [10]. The formal solution to this problem is known [10,11]. It is rather complicated and not easy to incorporate in a PIQMC simulation.

In essence, the method proposed in this paper is similar to the one used to solve the hydrogen path integral, i.e., use the quantum fluctuations to smear out the singularity of the potential. Mathematically we implement this idea by applying Jensen’s inequality to the propagator [12]. Applications of the Feynman path-integral formalism are often based on a
combination of Jensen’s inequality and a variational approach [8,10], so it is not a surprise that similar tricks may work for PIQMC as well.

The paper is organized as follows. In Sec. II, we give a simple derivation of an exact lower bound on the imaginary-time propagator. This inequality naturally defines a sequence of systematic approximations \( \tilde{Z}_m \) to the partition function. Although each \( \tilde{Z}_m \) looks very similar to \( Z_m \), the former can be used for PIQMC with attractive, singular potentials. For pedagogical reasons, in Sec. III we illustrate the approach by presenting an analytical treatment of the harmonic oscillator. In Sec. IV, we give the explicit form of the approximate propagator for the attractive Coulomb potential and present PIQMC results for the ground state of the hydrogen and helium atom.

II. LOWER BOUND ON THE PROPAGATOR

Consider a system with Hamiltonian \( H = K + V \) and a complete set of states \( \{x\} \) that diagonalizes the Hermitian operator \( V \). In the case in which \( V \) contains a singular attractive part, we replace \( V = \lim_{\epsilon \to 0} V_\epsilon \) by a regular \( V_\epsilon(x) > -\infty \) and take the limit \( \epsilon \to 0 \) at the end of the calculation. Using the Trotter-Suzuki formula, we can write

\[
\langle x | e^{-\tau(K + V)} | x' \rangle = \lim_{m \to \infty} \langle x | (e^{-\tau K/m} e^{-\tau V_\epsilon/m})^m | x' \rangle,
\]

where \( m \) is the number of Trotter steps.

\[
\lim_{m \to \infty} \int dx_1 \cdots dx_n \prod_{i=1}^{m} \langle x_i | e^{-\tau K/m} | x_i+1 \rangle e^{-\tau V_\epsilon(x_i)/m} \]

\[
\times \int dx_1 \cdots dx_n \prod_{i=1}^{m} \langle x_i | e^{-\tau K/m} | x_i+1 \rangle.
\]

If \( \langle x | e^{-\tau K} | x' \rangle \geq 0 \) for all \( \tau \), \( x \), and \( x' \), the function

\[
\rho(\{x_i\}) = \prod_{i=1}^{m} \langle x_i | e^{-\tau K/m} | x_i+1 \rangle
\]

\[
\times \left( \int dx_1 \cdots dx_n \prod_{i=1}^{m} \langle x_i | e^{-\tau K/m} | x_i+1 \rangle \right)
\]

is a proper probability density. Clearly, Eq. (10) is of the form \( \int dx_1 \cdots dx_n \rho(\{x_i\})f(\{x_i\}) \) so that we can apply Jensen’s inequality

\[
\int dx_1 \cdots dx_n \rho(\{x_i\}) e^{\tau f(\{x_i\})}
\]

\[
\geq \exp \left( \int dx_1 \cdots dx_n \rho(\{x_i\}) f(\{x_i\}) \right)
\]

and obtain

\[
\langle x | e^{-\tau K + V_\epsilon} | x' \rangle
\]

\[
\geq \langle x | e^{-\tau K} | x' \rangle \lim_{m \to \infty} \exp \left( -\frac{\tau}{m} \sum_{i=1}^{m} \int dx_1 \cdots dx_m 
\times \frac{V_\epsilon(x_i) \prod_{n=1}^{m} \langle x_n | e^{\tau K/m} | x_{n+1} \rangle}{\langle x | e^{-\tau K} | x' \rangle} \right),
\]

\[
\geq \langle x | e^{-\tau K} | x' \rangle \exp \left( -\frac{\tau}{m} \sum_{i=1}^{m} \int dx_i 
\times \frac{\langle x | e^{\tau K/m} | x_i \rangle V_\epsilon(x_i) \langle x_i | e^{\tau K/m} | x' \rangle}{\langle x | e^{-\tau K} | x' \rangle} \right).
\]

For \( m \to \infty \), the sum over \( n \) can be replaced by an integral over imaginary time. Finally, we let \( \epsilon \to 0 \) and obtain [12]

\[
\langle x | e^{-\tau(K + V)} | x' \rangle
\]

\[
\geq \langle x | e^{-\tau K} | x' \rangle \times \exp \left( -\int_0^\tau d\tau \frac{\langle x | e^{-\tau K} | x' \rangle}{\langle x | e^{-\tau K} | x' \rangle} \right).
\]

Note that the left-hand side of Eq. (14) reduces to the standard, symmetrized Trotter-Suzuki formula approximation [13,14] if we replace the integral over \( u \) by a two-point trapezium-rule approximation. This replacement also changes the direction of inequality as can been seen directly from the upper bound [12],

\[
\langle x | e^{-\tau K + V_\epsilon} | x' \rangle
\]

\[
\leq \langle x | e^{-\tau K} | x' \rangle \times \exp \left( -\int_0^\tau d\tau \ln \left( \frac{\langle x | e^{-\tau K} | x' \rangle}{\langle x | e^{-\tau K} | x' \rangle} \right) \right)
\]

\[
\leq \langle x | e^{-\tau K} | x' \rangle e^{-\tau V(x)}.
\]
\[
Z_m = \left( \frac{M}{2\pi\hbar^2} \right)^{3m/2} \int dr_1 \cdots dr_m \times \prod_{n=1}^{m} \exp \left[ -\frac{M}{2\pi\hbar^2} (r_n - r_{n+1})^2 \right. \\
- \left. \int_0^{1} du \langle r_n e^{-\nu K} V e^{-(\tau - \nu) K} | r_{n+1} \rangle \right], 
\]

where \( \tau = \beta/m \). The simplest approximant \( \hat{Z}_1 \) corresponds to Feynman’s variational approximation to the full Feynman path integral [8,10]. The main difference between Eqs. (3) and (16) is that the bare potential \( e^{-V(x)} \) is replaced by an effective potential that is obtained by convoluting the bare potential and free-particle propagators \( e^{-u K} \) and \( e^{-(\tau - u) K} \). Convolution smears out singularities. As we show below, in the case of the attractive Coulomb interaction, expression (14) is finite for any choice of \( x \) and \( x' \). For the approximants \( \hat{Z}_m \) to be useful in PIQMC, it is necessary that the integral over \( u \) be done efficiently. In the next two sections, we show how this can be done.

### III. ILLUSTRATIVE EXAMPLE

It is instructive to have at least one example for which the details can be worked out analytically, without actually using PIQMC. Not surprisingly, this program can be carried out for the harmonic oscillator. For notational convenience, we will consider the one-dimensional model Hamiltonian \( H = K + V \), with \( K = -(\hbar^2/2m) d^2/dx^2 \) and \( V = M/2x^2/2 \). Calculating the matrix element \( \langle x | e^{-u K} V e^{-(\tau - u) K} | x' \rangle \) in Eq. (16) is a straightforward exercise in performing Gaussian integrals [15]. We obtain

\[
\hat{Z}_m = \left( \frac{m M}{2\pi\hbar^2} \right)^{m/2} \int dx_1 \cdots dx_m \times \prod_{n=1}^{m} \exp \left[ -\frac{m M}{2\beta\hbar^2} (x_n - x_{n+1})^2 - \frac{\beta M\omega^2}{6m} x_n^2 + x_{n+1}^2 \right. \\
\left. + x_n x_{n+1} + \frac{\beta\hbar^2}{2mM} \right].
\]

For comparison, if we use the standard Trotter-Suzuki formula, we obtain [7]

\[
E_m = \beta\hbar^2 \omega^2 \sum_{n=0}^{m-1} \frac{1}{1 - \cos(2\pi n/m) + \beta^2\hbar^2 \omega^2/2m^2}.
\]

In Table I, we present numerical results obtained from Eqs. (19) and (20) and compare with the exact value of the energy \( E = (\hbar \omega/2) \coth(\beta\hbar \omega/2) \). Note that the average of the two approximations, i.e., \( (\hat{E}_m + E_m)/2 \), is remarkably close to the exact value \( E \), an observation for which we have no mathematical justification at this time.
IV. ATTRACTIVE COULOMB POTENTIAL

As a second example, we will consider a neutral system consisting of two electrons with opposite spin and a nucleus. The Hamiltonian reads [16,17]

\[ H = -\frac{\hbar^2}{2M_1} \nabla_1^2 - \frac{\hbar^2}{2M_2} \nabla_2^2 - \frac{q^2}{|r_1|} - \frac{2q^2}{|r_1 - r_2|}, \]

(21)

where the vectors \( r_1 \) and \( r_2 \) describe the position of the two electrons, with the nucleus placed in the origin. It is convenient to introduce the notation \( K_i = -D_i \nabla_i^2, \) \( D_i = \hbar^2/2M_i, \) \( V_i = V(r_i), \) \( V_{12} = V(r_1 - r_2), \) and \( V(r) = q^2/|r|, \) for \( i = 1, 2. \)

Application of inequality (14) requires the evaluation of

\[ I(r_1, r_2, r_1', r_2') = -\int_0^\tau du(r_1 r_2 | e^{-u(K_1 + K_2)} | r_1 r_2') \frac{\langle r_1 r_2 | e^{-\beta(K_1 + K_2)} | r_1' r_2' \rangle}{\langle r_1 | e^{-\beta K_1} | r_1' \rangle \langle r_2 | e^{-\beta K_2} | r_2' \rangle} \]

\[ = -\int_0^\tau du(r_1 | e^{-uK_1} V_1 e^{-(\tau-u)K_1} | r_1') \frac{\langle r_1 | e^{-\beta K_1} | r_1' \rangle}{\langle r_1 r_2 | e^{-\beta K_1} V_1 e^{-(\tau-u)K_1} | r_1' r_2' \rangle} \]

\[ + 2 \frac{\langle r_1 r_2 | e^{-\beta K_1} V_1 e^{-(\tau-u)K_1} | r_1' r_2' \rangle}{\langle r_1 r_2 | e^{-\beta K_1} | r_1' r_2' \rangle}, \]

(22)

where we made use of the fact that \([K_1, V_2] = [K_2, V_1] = 0. \) It is sufficient to consider the last term of Eq. (22). Inserting a complete set of states for both particles, we obtain

\[ I_{12}(r_1, r_2, r_1', r_2') = \int_0^\tau du \int dr_1^2 \int dr_2^2 \int dr_1' dr_2' \frac{\langle r_1 r_2 | e^{-u(K_1 + K_2)} | r_1' r_2' \rangle V(r_1 - r_2) V(r_1' - r_2') \langle r_1' r_2' | e^{-(\tau-u)(K_1 + K_2)} | r_1 r_2 \rangle}{\langle r_1 r_2 | e^{-\beta(K_1 + K_2)} | r_1' r_2' \rangle}. \]

(23)

Inserting the explicit expression for the free-particle propagator (5), a straightforward manipulation of the Gaussian integrals in Eq. (23) gives

\[ I_{12}(r_1, r_2, r_1', r_2', D) = \int_0^\tau du \int dr_1 \int dr_2 \frac{\tau}{4\pi u(\tau-u)D} \left( \frac{\tau}{4\pi u(\tau-u)D} \right)^{3/2} V(r) \exp \left\{ -\frac{[\tau -(\tau-u)(r_1 - r_2) - u(r_1' - r_2')]^2}{4u(\tau-u)D} \right\}, \]

(24)

where \( D = D_1 + D_2. \)

In the case of the Coulomb potential, the integral over \( r \) can be evaluated analytically by changing to spherical coordinates. The remaining integral over \( u \) is calculated numerically. In practice, it is expedient to replace the integration over \( u \) by an integration over an angle. An expression that is adequate for numerical purposes is

\[ I_{12}(r_1, r_2, r_1', r_2', D) = \frac{2\pi q^2}{D} \int_0^{\pi/2} d\phi \]

\[ \times \frac{\text{erf}[(4\pi D)^{-1/2}((r_1 - r_2)\tan \phi + (r_1' - r_2')\cot \phi)]}{((r_1 - r_2)\tan \phi + (r_1' - r_2')\cot \phi)} \]

(25)

It is easy to check that \( I_{12}(r_1, r_2, r_1', r_2', D) \) is finite.
whereas in the case of the hydrogen atom we have
\[
Z_m^H = \left( \frac{M}{2\pi\tau h^2} \right)^{3m/2} \int dr_1 \cdots dr_m 
\times \exp \left\{ -\frac{M}{2\tau h^2} \sum_{n=1}^{m} (r_n - r_{n+1})^2 
+ \tau \sum_{n=1}^{m} I_1(r_n, r_{n+1}, D_1) \right\},
\]
with \( \tau = \beta/m \). As the integrands in Eqs. (26) and (27) are always finite, expressions (26) and (27) can be used to perform PIQMC simulations.

In the path integral formalism, the ground-state energy is obtained by setting \( \beta \to \infty \) and \( \beta/m \to 0 \), i.e., \( E = \lim_{\beta \to \infty} \lim_{\beta/m \to 0} \hat{E}_m \). Of course, in numerical work, taking one or both of these limits is impossible. In Tables II and III, we present numerical results of PIQMC estimates of the ground-state energy \( E \) of the hydrogen and helium atom. These results have been obtained from five statistically independent simulations of 100,000 Monte Carlo steps per degree of freedom each. The systematic errors due to the discretization of the path integral are hidden in the statistical noise. The PIQMC procedure we have used is standard [1,7] except for a trick we have used to improve the efficiency of sampling the paths, details of which are given in the Appendix. Although a ground-state calculation pushes the PIQMC method to the point of becoming rather inefficient, the numerical results are in satisfactory agreement with the known values.

### V. DISCUSSION

We have shown that it is possible to perform PIQMC simulations for quantum systems with attractive Coulomb potentials. Instead of the conventional Trotter-Suzuki formulæ approach, one can use Eq. (16) to construct a path integral that is free of singularities. In practice, a numerical calculation of the latter requires only minor modifications of a standard PIQMC code.

The efficiency of the PIQMC method described above can be improved with relatively modest efforts. Instead of using the free-particle propagator \( \hat{K} \), we are free to pick any other model Hamiltonian \( \hat{H}_0 \) for which the matrix elements of \( e^{-\tau \hat{H}_0} \) are positive and integrals involving these matrix elements are known analytically. An obvious choice would be to take for \( \hat{H}_0 \) a set of harmonic oscillators. The matrix elements of \( e^{-\tau \hat{H}_0} \) are Gaussians and hence the conditions used to derive Eq. (14) are satisfied. If necessary, the approximant \( \hat{Z}_m \) can be improved further by optimization of the parameters of the oscillators. For \( m = 1 \), this approach is identical to the variational method proposed by Feynman and Kleinert [18–22] and independently by Giachetti and Tognetti [23,24].

Recently, a systematic approximation scheme was proposed, combining the smearing of the potential with the cumulant expansion [25–28]. In view of the good PIQMC results obtained with the primitive formalism adopted in this paper, the application of these more sophisticated schemes to PIQMC seems promising. Using a more accurate effective potential will yield results of the same quality with fewer time slices (\( m \)). However, the computational work required to numerically evaluate the effective potential will increase with its accuracy. A good compromise between improved accuracy on the one hand and increasing computational work on the other will have to be found. Extending the PIQMC method in this direction is left for future research.

### APPENDIX

In PIQMC, the simplest method for sampling paths is to change one degree of freedom at each Monte Carlo step. Usually, this is rather inefficient and one adds Monte Carlo moves that make global changes of the path, e.g., moves that resemble classical motion. In this appendix, we present a more sophisticated scheme, which we found performed very well at very low temperature. The basic idea is to change variables such that the kinetic-energy term in the path integral becomes a diagonal quadratic form, i.e.,
\[
\sum_{k=1}^{m} (x_k - x_{k+1})^2 = \sum_{k=2}^{m} y_k^2, \tag{A1}
\]
where \( x_{m+1} = x_1 \). After some straightforward algebra, one finds that the transformation from the \( \{x_i\} \) to the \( \{y_i\} \) is given by
\[
y_k^2 = \frac{m-k+2}{m-k+1} \left( x_k - \frac{m-k+1}{m-k+2} x_{k-1} + x_{m+1} \right)^2. \tag{A2}
\]
The expression for \( x_k \) in terms of the \( \{u_i\} \) reads
\[
x_k = y_1 + \sum_{j=2}^{k} \frac{m-j+1}{m-j+2} \frac{m-j+1}{m-j+2}^{1/2} y_j, \quad 1 < k \leqslant m. \tag{A3}
\]
with $x_1 = y_1$. From Eq. (A3), we conclude that the computational work for making a global change of the path (i.e., simultaneously changing all $y_i$) is linear in $m$, hence optimal. It is also clear that the variable $y_1$ plays the role of the ‘‘classical’’ position. The variables $y_2, \ldots, y_m$ describe the quantum fluctuations.

[15] This is the case for all $V(x)$ that are polynomial in $x$.