Coupling quantum Monte Carlo and independent-particle calculations: Self-consistent constraint for the sign problem based on the density or the density matrix

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Quantum Monte Carlo (QMC) methods are one of the most important tools for studying interacting quantum many-body systems. The vast majority of QMC calculations in interacting fermion systems require a constraint to control the sign problem. The constraint involves an input trial wave function which restricts the random walks. We introduce a systematically improvable constraint which relies on the fundamental role of the density or one-body density matrix. An independent-particle calculation is coupled to an auxiliary-field QMC calculation. The independent-particle solution is used as the constraint in QMC, which then produces the input density or density matrix for the next iteration. The constraint is optimized by the self-consistency between the many-body and independent-particle calculations. The approach is demonstrated in the two-dimensional Hubbard model by accurately determining the ground state when collective modes separated by tiny energy scales are present in the magnetic and charge correlations. Our approach also provides an ab initio way to predict effective interaction parameters for independent-particle calculations.

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I. INTRODUCTION

The study of interacting quantum many-body systems presents a major challenge in modern physics. Quantum Monte Carlo (QMC) methods\textsuperscript{1–4} are a key numerical approach for solving such systems. The dimension of the Hilbert space involved in a quantum many-body system grows exponentially with the system size. QMC methods can in principle provide stochastic evaluations of expectation values in such systems with computer times that scale polynomially with system size. However, with a few exceptions\textsuperscript{5,6}, direct QMC calculations in fermion systems suffer from the minus sign problem\textsuperscript{7,8}, which breaks this scaling. The most effective approach for dealing with the sign problem in general has been by a bias-variance trade-off. A constraint is applied in some space to restrict the Monte Carlo sampling, which introduces a systematic bias but in turn removes the exponential growth in variance and restores the algebraic complexity of the algorithm. The majority of QMC calculations have employed this approach, including many on spin and fermion models\textsuperscript{9,10}, and almost all on realistic systems in condensed matter physics\textsuperscript{11–13}, nuclear physics\textsuperscript{14}, and quantum chemistry\textsuperscript{15–17}.

A missing link in such an approach is that it has been difficult to make the constrained QMC calculations systematically improvable without drastically changing its computational scaling or complexity\textsuperscript{18}. Although the calculations are often among the most accurate possible for many-fermion systems\textsuperscript{10}, the accuracy cannot be assessed internally, and there has not been a conceptual framework which allows one to build on the outcome of the calculation in a practical way to further reduce the systematic error from the constraint. The constraint typically relies on a trial wave function which is provided by an external source (e.g., an independent-electron calculation or a variational Monte Carlo optimization\textsuperscript{19}), and a “one-shot” answer is obtained from the QMC.

In this paper we introduce a self-consistent constraint in QMC using the auxiliary-field QMC (AFQMC) framework. The approach couples the AFQMC calculation to an independent-electron calculation which provides the trial wave function for the constraint. The spin densities (or density matrix) obtained from the QMC are then fed back into the independent-electron calculation, whose effective interaction strength (or more generally, exchange-correlation functional) is tuned to best match the QMC densities. The output wave function is then used for a new AFQMC calculation, and the process is iterated to convergence. We show that this procedure allows the calculations to systematically improve. The QMC can recover from an initial constraint in a wrong state, i.e., one with an incorrect magnetic order, and provides the correct prediction at convergence even when a small residual constraint error is still present.

In an alternative, complementary view, the self-consistent approach is motivated by the fundamental role of the electron density or density matrix in many-fermion systems\textsuperscript{20}. The constraining wave function in AFQMC is usually taken as a solution from the Hartree-Fock (HF) or a density-functional theory (DFT) calculation based on the same many-body Hamiltonian. What is the optimal independent-electron wave function? Our approach defines a procedure for determining the answer. We will show that, by varying the strength of the Coulomb repulsion, the HF calculation can give better order parameters (spin densities here). The self-consistency procedure with QMC thus provides an optimal effective “U” parameter. This can potentially be used to derive effective Hamiltonians to be used by less computing-intensive methods in larger system sizes. Similarly, in the context of DFT, the procedure would define a way to find an optimal functional (in the spirit of hybrid functionals, for example).

For concreteness, we will use the Hubbard model to
describe the self-consistent AFQMC procedure:
\[
\hat{H} = - \sum_{i,j,\sigma} t_{ij} c_{i\sigma}^{\dagger} c_{j\sigma} + \sum_{i} U \hat{n}_{i\uparrow} \hat{n}_{i\downarrow} + \sum_{i} v_{i,\sigma} \hat{n}_{i\sigma},
\]
where \( c_{i\sigma}^{\dagger} (c_{i\sigma}) \) is the creation (annihilation) operator on lattice site \( i \), \( \sigma = \uparrow, \downarrow \) is the spin of the electron, \( \hat{n}_{i\sigma} = c_{i\sigma}^{\dagger} c_{i\sigma} \) is the number operator. The hopping matrix elements \( t_{ij} \), on-site interaction strength \( U \), and spin-dependent external potential \( v_{i,\sigma} \) are parameters. The overall electron density is given by the parameter \( n \equiv (N_{\uparrow} + N_{\downarrow})/N \), with \( N \) being the total number of lattice sites, and the hole density is then \( h = 1 - n \).

II. SELF-CONSISTENT METHOD COUPLING WITH INDEPENDENT- ELECTRON CALCULATIONS

The corresponding independent-particle (IP) calculation treats a Hamiltonian of the form:
\[
\hat{H}_{\text{IP}}^{\sigma} = - \sum_{i,j} t_{ij} c_{i\sigma}^{\dagger} c_{j\sigma} + \sum_{i} U_{\text{eff}}(\hat{n}_{i\uparrow}) \hat{n}_{i\downarrow} + \sum_{i} v_{i,\sigma} \hat{n}_{i\sigma},
\]
where \( \sigma \) denotes the opposite of \( \sigma \). In the standard unrestricted HF (UHF) calculation, \( U_{\text{eff}} \) takes the “bare” value of \( U \), and the input mean-field is a set of expectation values, \( \langle \hat{n}_{i\sigma} \rangle \), computed with respect to the solution from the previous IP step. In DFT with a local spin-density type of approach, \( U_{\text{eff}}(\hat{n}_{i\sigma}) \), in Eq. (2) is replaced by an exchange-correlation functional, \( V_{\text{xc}}(\langle \hat{n}_{i\sigma} \rangle) \).

The wave function from the IP solution is a single Slater determinant, \( \left| \psi \right\rangle = \left| \psi_{\uparrow} \right\rangle \otimes \left| \psi_{\downarrow} \right\rangle \), with \( \left| \psi_{\sigma} \right\rangle = \phi_{\uparrow}^{j_{\uparrow}} \phi_{\downarrow}^{j_{\downarrow}} \cdots \phi_{N_{\sigma}}^{j_{N_{\sigma}}} (0) \) where \( \phi_{\sigma}^{j_{\sigma}} = \sum_{\sigma} \phi_{\sigma}^{j_{\sigma}} c_{j_{\sigma}} \) creates a \( \sigma \)-spin electron in the single-particle orbital given by the vector \( \{ \phi_{\sigma}^{j_{\sigma}} \}_{j = 1, \ldots, N} \).

The AFQMC method projects the many-body ground state of \( \hat{H} \) in Eq. (1) by an iterative process:
\[
\lim_{m \to \infty} \left( e^{-\tau \hat{H}} \right)^{m} \left| \psi_{T} \right\rangle \propto \left| \Psi_{0} \right\rangle,
\]
where \( \tau > 0 \) is a small parameter. For convenience, we take the initial state, which must be non-orthogonal to \( \left| \Psi_{0} \right\rangle \), to be a single Slater determinant trial wave function, \( \left| \psi_{T} \right\rangle \). The many-body propagator is written as \( e^{-\tau \hat{H}} = \int p(x) e^{\hat{h}(x)} dx \) where \( \hat{h}(x) \) is a general IP “Hamiltonian” dependent on the multi-dimensional vector \( x \), and \( p(x) \) is a probability density function. The interacting many-body system is thus mapped into a linear combination of many IP systems in fluctuating auxiliary fields, \( x \). The AFQMC method represents the many-body wave function as an ensemble of Slater determinants, i.e., \( \left| \Psi_{0} \right\rangle = \sum_{k} \omega_{k} \left| \psi_{k} \right\rangle \).

The iterative projection is realized by a random walk in Slater determinant space, in which for each walker \( \left| \psi_{k} \right\rangle \), an auxiliary field \( x \) is sampled from \( p(x) \), and the walker is propagated: \( e^{\hat{h}(x)} \left| \psi_{k} \right\rangle \to \left| \psi'_{k} \right\rangle \). Computationally this is similar to a step in the IP calculation.

Because the propagator \( e^{\hat{h}(x)} \) contains stochastically fluctuating fields, the random walks will, except for special cases protected by symmetry, reach Slater determinants with arbitrary sign or phase. In representing the ground state, only one from each pair of Slater determinants \( \{ \pm \left| \psi \right\rangle \} \) (or from the set \( \{ e^{i\alpha} \left| \psi \right\rangle \} \)) is needed. When both (all) are present in the samples, the wave function signal is lost in noise, because the Monte Carlo weights, \( w_{k} \), are always positive. This is the sign (phase) problem.

For the Hubbard Hamiltonian, \( \hat{h}(x) \) is real, so “only” a sign problem appears. To control this problem, we use the trial wave-function \( \left| \psi_{T} \right\rangle \) for importance sampling, which guides the random walk and constrains it to only half of the Slater determinant space:
\[
\left\langle \psi_{T} \left| \psi_{k} \right\rangle > 0 \right\rangle^{22}.
\]
This approach has been referred to as the constrained-path Monte Carlo (CPMC) method. For a general Hamiltonian with two-body interactions, a generalized gauge condition allows a similar framework for the phase problem.

This framework eliminates the sign or phase problem, at the cost of introducing a systematic bias. Previous studies in a variety of systems have shown that the bias tends to be small, in both models and realistic materials, making this one of the most accurate many-body approaches for general interacting fermion systems. In this work, we introduce a self-consistent method to further reduce the bias introduced by the constraint from the trial wave function.

To start the self-consistent procedure, we first carry out a CPMC calculation for the many-body Hamiltonian, Eq. (1), using any typical choice of \( \left| \psi_{T} \right\rangle \), for example a non-interacting wave function or the UHF solution. We use back-propagation to compute the expectation values of the quantities that do not commute with \( \hat{H} \). After the CPMC calculation, we solve the IP Hamiltonian in Eq. (2), using the densities obtained from the preceding QMC calculation as the input mean field, i.e., \( \langle \hat{n}_{i\sigma} \rangle_{\text{QMC}} \to \langle \hat{n}_{i\sigma} \rangle \). An effective interaction, \( U_{\text{eff}} \), is applied instead of the “bare” \( U \) value. We vary \( U_{\text{eff}} \) to find an optimal value whose solution gives densities closest to the input from QMC, i.e., the \( U_{\text{eff}} \) which minimizes:
\[
\delta = \frac{1}{N} \sum_{i\sigma} \left( \langle \hat{n}_{i\sigma} \rangle_{\text{IP}} - \langle \hat{n}_{i\sigma} \rangle_{\text{QMC}} \right)^{2}^{1/2}.\]

The IP solution with the optimal \( U_{\text{eff}} \) determined from Eq. (3) is then used as the \( \left| \psi_{T} \right\rangle \) in a new CPMC calculation. This procedure is iterated until convergence.

III. RESULTS

We use the two-dimensional Hubbard model at density \( n = 0.875 \) as a test case. The hopping matrix element \( t_{ij} = t \) for nearest neighbors and 0 otherwise, and we set the interaction \( U = 8t \). This parameter regime, mimicking the situation in doped cuprates, is notoriously challenging, and its ground state in the thermodynamic limit remains unknown. We focus on the nature of the magnetic and charge correlations, which is crucial for un-
FIG. 1: Systematic improvement of the CPMC accuracy from the self-consistent procedure. The top panel plots staggered spin density along the $y$-direction vs. site label $i_y$ ($i_x = 1$). The bottom panel plots the corresponding hole density. The left and right columns show two self-consistent procedures starting from two different initial $|\Psi_T\rangle$’s, the free-electron wave function and the UHF solution at $U = 8t$, respectively. In the legend, the $U_{\text{eff}}$ value of the IP calculation is listed for each iteration step. In (a) and (b), the differences at different stages of the iteration, with respect to the reference DMRG results, are shown in the insets. The system is $4 \times 16$, with $U = 8t$, $h = 1/8$ doping, with pinning field applied to both edges along $L_y$.

Understanding the properties of lightly doped antiferromagnets.

In our test calculations, we will consider a cylindrical geometry, i.e., cells with periodic boundary conditions in the $x$-direction and open boundary conditions in the $y$-direction, for which density matrix renormalization group (DMRG) calculations can give very accurate benchmark results for systems of significant sizes. We denote the integer coordinates of the lattice site $i$ by $(i_x, i_y)$. Local antiferromagnetic (AFM) correlations are expected, and our calculations aim to probe what happens to the antiferromagnetic (AFM) correlations are expected, and our calculations aim to probe what happens to the AFM correlation in this regime of doping ($h = 1/8$) and strong interaction ($U = 8t$). To break degeneracy from translational symmetry, pinning fields$^{31-33}$ are applied at the edges: $v_y = -v_L = (-1)^{i_y}v_0$ for $i_y = 1$ and $i_y = L_y$. We set the pinning field strength $v_0 = t/4$ in all calculations. With pinning fields, two-body spin and charge correlation functions in periodic systems are turned into one-body spin and charge order parameters, which are simpler to measure in the calculation. Thus, in addition to the total energy, we will focus on the local spin and charge densities

$$h(i) = 1 - \langle \hat{n}_{i \uparrow} + \hat{n}_{i \downarrow}\rangle; \quad S_z(i) = \langle \hat{n}_{i \uparrow} - \hat{n}_{i \downarrow}\rangle/2.$$

We first illustrate the method in a $4 \times 16$ system. Two different choices of the trial wave function $|\psi_T\rangle$ are used for the initial CPMC calculation. The first is the ground state of the corresponding non-interacting Hamiltonian (referred to as free-electron hereafter). The other is the UHF solution$^{34}$ obtained with the “bare” $U$ value, i.e., $U = 8t$. In Fig. 1, we show the staggered spin densities, $(-1)^{i_y}S_x(i)$, and the hole densities $h(i)$ computed by the self-consistent QMC procedure, and compare them with DMRG results, which are essentially exact for this system. The results are, as expected, statistically invariant with respect to $i_x$, and are only shown for $i_x = 1^{35}$. The staggered spin densities, shown in the upper panel, depict a modulation of the AFM order. (The magnetic moments are the strongest at the edges because of the pinning fields.) At a node when the curve crosses zero, a $\pi$ phase shift is created in the AFM pattern. The holes tend to concentrate at the nodes, creating the peaks seen in the bottom panel.

We see from Fig. 1 that, independent of which $|\psi_T\rangle$ is used in the initial CPMC calculations, the self-consistent procedure leads to a systematic improvement of the spin densities which approach the DMRG results. Convergence from the free-electron $|\psi_T\rangle$ requires only about 3 iterations between the QMC and IP calculations. The UHF initial $|\psi_T\rangle$, which predicts a wrong phase (see below), gives results in the first-iteration CPMC with large errors. The self-consistency quickly recovers and converges in about 6 iterations. The difference between converged CPMC spin densities and those from DMRG, as seen in the insets, is very small. The hole density, which shows a slightly larger residual error, clearly gives the correct charge pattern. The self-consistent CPMC method thus accurately determines the ground state and its magnetic order in this system.

The convergence process of the self-consistent procedure is further illustrated in Fig. 2. The left panel shows the relative error in the CPMC ground-state energy per site, from the reference DMRG value of $-0.77127(2)$. The energy is seen to converge, independent of the initial $|\psi_T\rangle$, to a value with a residual relative error about $-0.4\%$. We note that the mixed estimate, which is used in CPMC to computed the energy, is not variational$^{25,36}$. The self-consistency actually leads in a slightly worse
The converged CPMC results after self-consistent procedure for a large system, of $16 \times 32$. In the upper panel, the staggered spin and hole densities are plotted. The red and blue horizontal lines represent zero spin-density and the average hole density, respectively. In the lower pane, the spin density for the cell is shown with a colormap. As in the earlier systems, $U = 8t$, $h = 1/8$, and pinning field is applied to both edges along $L_y$.

with the four nodal lines of modulation clearly visible (where the holes are more concentrated). This is consistent with a wavelength of $1/h$ seen at lower interaction strengths.

IV. DISCUSSION

We have also tested a different but related approach for constructing the trial wave function self-consistently from QMC. To encode the information on the one-body density matrix from CPMC, $\rho^{\text{CPMC}}$, in the next stage $|\Psi_T\rangle$, we seek a Slater determinant which gives a one-body density matrix closest to $\rho^{\text{CPMC}}$. This is done using natural orbitals, i.e., by diagonalizing:

$$\rho^{\text{CPMC}} = V \Lambda V^\dagger.$$  \hspace{1cm} (5)

The eigenvectors (natural orbitals) in $V$ corresponding to the $N_x$ largest eigenvalues in $\Lambda$ are chosen to construct a Slater determinant. This procedure is implemented for each spin species $\sigma$ separately, leading to a UHF-like $|\Psi_T\rangle$. We found that this procedure gave results similar to the self-consistent approach in the systems tested above.

Although we have used the Hubbard model as an illustration, the self-consistent procedure we have proposed can be generalized to AFQMC calculations in real materials, and opens new directions to further improve the predictive power of calculations in correlated electron systems. In real materials, the self-consistent iteration with IP calculations can be used to improve an exchange-correlation functional, for example to tune the optimal mixing parameter in a hybrid functional. The procedure could also be used to find a correct “$U$” param-
eter in the context of LDA+$U^{41}$. The idea of introducing an effective “$U$” can also be connected to embedding theories$^{42,43}$, although here the feedback of $U_{\text{eff}}$ to the real system (cluster) is less direct.

V. SUMMARY

In summary, we have developed a self-consistent constrained-path AFQMC method which allows the QMC calculation to systematically improve its accuracy, while fully controlling the fermion sign or phase problem. The paradigm coupling QMC with an IP calculation allows a feedback from the former into the latter. This provides not only a way to improve the constraining trial wave functions for the (next iteration) QMC, but also provides not only a way to improve the constraining trial wave functions for the (next iteration) QMC, but also an independent-particle framework which in itself gives a drastically improved description of the physical system. The approach can be applied to strongly correlated models in condensed matter, ultra-cold atoms and optical lattices, nuclear shell models, as well as \textit{ab initio} calculations in molecules and solids.

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References

For hole density, the results are the same for each row, while for spin density, there is a $\pi$ phase between even and odd rows.