Symmetry in Auxiliary-Field Quantum Monte Carlo Calculations

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We show how symmetry properties can be used to greatly increase the accuracy and efficiency in auxiliary-field quantum Monte Carlo (AFQMC) calculations of electronic systems. With the Hubbard model as an example, we study symmetry preservation in two aspects of ground-state AFQMC calculations, the Hubbard-Stratonovich transformation and the form of the trial wave function. It is shown that significant improvement over state-of-the-art calculations can be achieved. In unconstrained calculations, the implementation of symmetry often leads to shorter convergence time and much smaller statistical errors, thereby a substantial reduction of the sign problem. Moreover, certain excited states become possible to calculate which are otherwise beyond reach. In calculations with constraints, the use of symmetry can reduce the systematic error from the constraint. It also allows release-constraint calculations, leading to essentially exact results in many cases. Detailed comparisons are made with exact diagonalization results. Accurate ground-state energies are then presented for larger system sizes in the two-dimensional repulsive Hubbard model.

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I. INTRODUCTION

The study of interacting quantum many-body systems remains an outstanding challenge, especially systems with strong particle interactions, where perturbative approaches are ineffective. Numerical simulations provide a promising approach for studying such systems. One of the most general numerical approaches is quantum Monte Carlo (QMC) methods based on auxiliary-fields, which are applied in condensed matter physics, nuclear physics, high-energy physics, and quantum chemistry. These methods allow essentially exact calculations of ground-state and finite-temperature equilibrium properties of interacting many fermion systems. As is well known, however, they suffer from the sign problem, which severely limits their applicability. A key issue in algorithm development in this context is to reduce the convergence time or variance reduction for fixed computing cost.

Considerable progress has been achieved in circumventing this problem by constraining the random walks in sampling the space of auxiliary-fields. These methods have come under the name of Constrained Path Monte Carlo (CPMC) for systems where there is a sign problem (for example, Hubbard-like models where the local interactions lead to auxiliary-fields that are real). For electronic systems where there is a phase problem (as the Coulomb interaction leads to complex fields), the methods have been referred to as phaseless or phase-free auxiliary-field QMC. In both cases, the idea is to constrain the sign or phase of the overlap of the sampled Slater determinants with a trial wave function. It eliminates the sign or phase instability and restores low-power (typically to the third power of system size) computational scaling. Applications to a variety of systems have shown that the methods are very accurate, even with simple trial wave function taken directly from mean-field calculations (see, e.g. Refs. and references therein). However, these methods are approximate. For example, open-shell situations often result in larger systematic errors. It is thus important to understand and develop ways to improve the quality of the constraint.

Symmetry properties and projection have a long history in quantum many body problems (see, e.g. Refs. ). In this paper, we discuss the imposition of symmetry properties and their effects in AFQMC calculations. It is shown that symmetry can often be rigorously preserved in the AFQMC framework despite the stochastic nature of the calculations, and taking advantage of this can lead to dramatic improvements. We address the issue from two aspects: symmetry preserved projection and the symmetry of the trial wave function.

In the symmetry preserved projection, we study the form of the Hubbard-Stratonovich (HS) transformation and its effect on the computational efficiency and accuracy. We focus on the comparison between charge-decompositions which preserve SU(2) symmetry and the standard form of Hirsch spin-decomposition. We discuss the optimal form of the charge decomposition which involves subtracting the proper “mean-field” background prior to making the HS transformation.

With a trial wave function which imposes exact symmetry properties, one is able to reduce the projection time as well as the statistical error bar. Symmetry alleviates the sign problem, since the convergence time to the ground state is reduced by ensuring the right symmetry in the trial wave function used to initiate the projection. Furthermore, the correct symmetry can help sustain projections of excited states which have different symmetry from that of the ground state. The most important feature of preserving symmetry in the trial wave function is in constrained AFQMC calculations, where we will show that the quality of the approximation improves significantly.

We consider the effect of symmetry in three different forms of ground-state AFQMC calculations, unconstrained Free Projection (FP), constrained path (CP) calculations, and released-constraint (RC) calculations. The
first, FP, corresponds to the standard algorithm which is formally exact but has the sign problem. The third, RC, can be thought of in the present context as a more sophisticated version of FP, in which we start from a much better initial state, namely a converged CP run, and release (i.e., remove) the constraint and continue the projection. In FP and RC, the focus is on the use of symmetry to reduce the statistical fluctuations and prolong the imaginary-time projections. We show that improvements are seen in both methods. Especially in RC the use of symmetry can lead to a dramatic effects and significantly ameliorate the sign problem.

We are interested in studying the effect of symmetry in CP calculations, in order to further improve the constraints. We find that the use of trial wave functions which preserve symmetry can greatly reduce the systematic bias from the constraint. In open-shell systems, the use of simple trial wave functions which restore symmetry properties eliminates most of the bias from the constraint. Another key finding is that one can switch between different forms of the HS transformation in CP and the following RC calculations. This is motivated by considering the different behaviors of statistical errors and sensitivity to the constraint.

We will use the two-dimensional repulsive Hubbard model as an example in our discussions. The model provides an ideal test ground for the present study. On the one-hand, it is computationally simple where exact diagonalization results are available in non-trivial system sizes. On the other hand, it is a fundamental model which captures many of the key aspects of simulating correlated materials and which, in fact, contains many unanswered questions. Through the approach presented in this paper, significantly more accurate calculations can be done on larger systems. For example, essentially exact results can now be obtained for $>100$ sites, as shown below.

The remainder of the paper is organized as follows. In section II we summarize the relevant features of the AFQMC method in three different forms: the unconstrained FP, CP, and RC, in order to facilitate the ensuing discussions. In section III we describe separately the two aspects of symmetry improved AFQMC, namely symmetry-preserving HS transformation and symmetry in the trial wave function. Results are first shown for demonstration and benchmark, then followed by new, near-exact ground-state energies for larger system sizes. Finally we conclude in Sec. IV.

II. AUXILIARY FIELD QUANTUM MONTE CARLO METHODS

We first summarize the key features of ground state AFQMC methods that are relevant to the symmetry study to follow. The three different forms of AFQMC are summarized in three subsections below. As mentioned, we will use the two-dimensional repulsive Hubbard model as a concrete example; however most of our discussions will apply to other Hamiltonians, including more realistic materials Hamiltonians.

The Hubbard model is written in second-quantized form as $^{19}$:

$$\hat{H} = \hat{K} + \hat{V} = -t \sum_{(i,j)\sigma} c_{i\sigma}^\dagger c_{j\sigma} + U \sum_{i} n_{i\uparrow} n_{i\downarrow}.$$  \hspace{1cm} (1)

Here $L$ is the number of lattice sites, $c_{i\sigma}^\dagger$ and $c_{i\sigma}$ are creation and annihilation operators of an electron of spin $\sigma$ on the $i$-th lattice site, $t$ is the nearest-neighbor hopping energy and $U$ is the interaction strength. Throughout this paper, we will use $t$ as units of energy and set $t = 1$. We assume that there are $N_{\uparrow}$ spin-up electrons and $N_{\downarrow}$ spin-down electrons on the lattice.

The Hamiltonian in Eq. (1), whose Hilbert space grows exponentially in size with $L$, presents an enormous challenge. Questions remain open about its properties. Ground state projection with Monte Carlo (MC) sampling is one of the candidates to solve the problem. The projection method is:

$$|\psi_0\rangle \propto \lim_{\beta \rightarrow \infty} e^{-\beta(\hat{H} - E_T)}|\psi_T\rangle,$$  \hspace{1cm} (2)

where $E_T$ and $|\psi_T\rangle$ are guesses of the ground state energy and wave function, and $\langle \psi_0 | \psi_T \rangle \neq 0$ in order for the projection to yield the ground state asymptotically. To target a lowest energy excited state of a different symmetry from $|\psi_0\rangle$ is similar to doing a ground-state calculation, except one would choose $|\psi_T\rangle$ which is not orthogonal to the targeted excited state but satisfies $\langle \psi_0 | \psi_T \rangle = 0$.

The propagator can be evaluated using a Trotter-Suzuki breakup $^{20,21}$:

$$(e^{-\Delta \tau (\hat{K} + \hat{V})})^n = (e^{-\frac{1}{2}\Delta \tau \hat{K}} e^{-\Delta \tau \hat{V}} e^{-\frac{1}{2}\Delta \tau \hat{K}})^n + O(\Delta \tau^2).$$  \hspace{1cm} (3)

Here we have $\Delta \tau n = \beta$, and a Trotter error arises from the omission of the higher order terms. We will not be concerned with the Trotter error here, other than to note that it can be controlled by extrapolating to $\Delta \tau \rightarrow 0$ with separate calculations using different $\Delta \tau$ values. In the results shown in this paper, we either perform such an extrapolation explicitly, or have checked via separate calculations that the Trotter error is within the statistical error. We also mention that, for the Hubbard interaction, decompositions without Trotter errors are possible $^{22}$.

The two-body propagator is then decoupled into one-body propagator by auxiliary fields, using the HS transformation $^{23,24}$. The general form is:

$$e^{-\Delta \tau \hat{V}} = \sum_x p(x)e^{\hat{O}(x)},$$  \hspace{1cm} (4)

where $\hat{O}(x)$ is a one-body operator that depends on the auxiliary field $x$, $p(x)$ is a probability density function with the normalization $\sum_x p(x) = 1$. In general the sum in Eq. (4) is an integral, and $x$ is a many-dimensional vector whose dimension is of the order of the size of the
one-particle basis. In the Hubbard model, \( x \) typically has \( L \) components, one for each lattice site. By setting:

\[
\hat{B}(x) = e^{-\frac{1}{2}\Delta tK}e^{i\phi(x)}e^{-\frac{1}{2}\Delta tK},
\]

we rewrite the projection as

\[
|\psi_0\rangle = \sum_{\vec{x}} P(\vec{x}) \prod_{i=1}^{n} \hat{B}(x_i)|\psi_T\rangle.
\]

The vector \( \vec{x} \) means \( (x_1, x_2, \cdots, x_n) \), and \( P(\vec{x}) = \prod_i p(x_i) \).

Then the ground state properties can be evaluated by:

\[
\langle \hat{A} \rangle_0 = \frac{\langle \psi_0|\hat{A}|\psi_0\rangle}{\langle \psi_0|\psi_0\rangle},
\]

which are many-dimensional integrals (e.g., \( 2nL \)-dimensions in the Hubbard model). MC methods are used to calculate the high dimension integrals, by sampling the probability density function using the Metropolis algorithm or a related method. The sign problem emerges because the integrand in the denominator, \( P(\vec{x})\langle \psi_T| \prod \hat{B}(x_i)|\psi_T\rangle \) is not always positive, which causes the MC signal to be eventually lost in the sampling noise.

### A. Unconstrained Free Projection (FP)

We carry out the FP calculation with an open-ended random walk similar to the CP approach. However, the discussion on symmetry in FP calculations in this paper will apply directly to standard ground-state AFQMC calculations using Metropolis sampling of paths. In our approach, a population of \( N_w \) random walkers is carried, which are typically initialized by the trial wave function. Each walker will have a weight \( w \) whose value is set as one at the beginning of the projection:

\[
|\psi^{(0)}\rangle = \sum_i w_i^{(0)}|\phi_i^{(0)}\rangle
\]

We apply the projection in Eq. 8 by random walks in Slater determinant space, instead of using the usual approach of Metropolis sampling of auxiliary-field paths. In each step, we sample the auxiliary field \( x \) according to \( p(x) \) by MC, and apply \( \hat{B}(x) \) to the Slater determinant wave function. Since the operators only contain one-body terms, they will generate another Slater determinant:

\[
|\psi^{(1)}\rangle = \sum_i w_i^{(1)}|\phi_i^{(1)}\rangle
\]

During the projection we multiply the constant (non-operator) values of the formula, e.g., the overall normalization \( e^{\beta H_T} \), to the weight of the walker. The weight of each walker will fluctuate in the random walk and after a few steps, some walkers can have large weights and some walkers will have small weights. We apply a population control procedure, splitting the walkers with large weights and eliminating walkers with small weights with the appropriate probability, such that the overall probability distribution is preserved but the weights are made more uniform. It will introduce a population control bias. In our calculations, the population control bias is much smaller than the statistical error. This is easy to accomplish in CP calculations. In FP and RC calculations, large populations are necessary as discussed below. Modified Gram-Schmidt orthogonalization is applied to each walker periodically as well.

As mentioned, the FP calculations are done in the same framework as the CP calculations. This is different from standard Metropolis sampling, which keeps an entire path of a fixed length \( n \) as the object to sample. This framework does not have any ergodicity issues, and it is straightforward to project to longer imaginary-time in order to approach the ground state. We typically turn off importance sampling in FP, sampling the fields according to \( P(\vec{x}) \) instead of using either the force bias or direct importance-sampling of discrete fields as is done in CP calculations. Empirically we find that this tends to give smaller statistical errors than invoking importance sampling and then lifting the constraint. The use of population control helps to reduce the noise but ultimately the shortcoming of this approach is that the lack of importance sampling will cause large noises as system size or \( n \) increases. Since in these situations the sign problem, when uncontrolled, tends to overwhelm the calculation anyway, the shortcoming is not of major practical relevance, and we find this mode of sampling to often be the more efficient in practice.

At the \( n^{th} \) step in the propagation, we measure the energy by:

\[
E = \frac{\sum_i w_i^{(n)} \langle \psi_T|H|\phi_i^{(n)}\rangle}{\sum_i w_i^{(n)} \langle \psi_T|\phi_i^{(n)}\rangle},
\]

If the projection has equilibrated, we can combine the populations at multiple \( n \) values in the estimator above to improve statistics on the ground-state energy. The energy measure in Eq. (10) is variational if we set \( |\psi^{(0)}\rangle = |\psi_T\rangle \), since

\[
E = \frac{\langle \psi_T|He^{-\beta H}|\psi_T\rangle}{\langle \psi_T|e^{-\beta H}|\psi_T\rangle} = \frac{\langle \psi_T|e^{-\beta H/2}He^{-\beta H/2}|\psi_T\rangle}{\langle \psi_T|e^{-\beta H/2}e^{-\beta H/2}|\psi_T\rangle}.
\]

To calculate the expectation value of an observable which does not commute with the Hamiltonian, we can use
back-propagation using part of the path and projecting the trial wave function $|\psi_T\rangle$. Because of the lack of importance sampling, back-propagation will tend to be very noisy in FP, and large population size will typically be needed.

The application of symmetry in either the trial wave function or the propagator $\hat{B}(x)$ is straightforward in FP. In Sec. \ref{sec:symmetry} we show how the use of symmetry can improve FP calculations and allow longer projection time with smaller statistical fluctuations. Furthermore, some excited states can be accessed.

\subsection{Constraint Path (CP)}

The constrained path (CP) approximation allows one to eliminate the sign problem present in FP. During the FP steps, the overlap between the ground state and the projected wave function, $\langle \psi_0 | \phi^{(l)} \rangle$, will in general approach zero, because intrinsically the projection is symmetric about $|\psi^{(l)}\rangle$ and $-|\psi^{(l)}\rangle$. In other words, at any given imaginary time $l$, the projection would proceed identically if each random walker $|\phi^{(l)}\rangle$ were switched to $-|\phi^{(l)}\rangle$, for example by a permutation of two of its orbitals. This means that, unless the random walks are somehow strictly confined to only one kind of “sign”, it will invariably become a random and equal mixture of both, given sufficiently large $l$. Thus measurements from the MC sampling will eventually have infinite variance. This is how the sign problem appears in an FP calculation.

The CP approach is based on the observation that $\Delta$, if any particular walker has the zero overlap with the ground state at imaginary time $\tau_l \equiv l\Delta\tau$ in the projection:

\begin{equation}
\langle \psi_0 | \phi^{(l)} \rangle = 0 \tag{12}
\end{equation}

this walker will contribute zero at any future time $\beta > \tau_l$, because:

\begin{equation}
\langle \psi_0 | e^{-(\beta-\tau_l)\hat{H}} | \phi^{(l)} \rangle = 0 . \tag{13}
\end{equation}

Then we are able to discard the walker once its path reaches a point where the overlap becomes zero. With this constraint the sign problem is eliminated, and the projection will still lead to the exact ground state. However, we obviously do not know the exact ground state wave function. In CP calculations, a trial wave function, $|\psi_T\rangle$, is chosen for determining the sign of the overlap. A walker which develops a zero overlap with $|\psi_T\rangle$ during the projection is discarded.

Importance sampling can be introduced in CP calculations both as a natural way to impose the constraint and for variance reduction. With importance sampling, the wave function during the projection can be written as:

\begin{equation}
|\psi^{(l)}\rangle = \sum_{i} w_i^{(l)} \frac{|\phi_i^{(l)}\rangle}{\langle \psi_T | \phi_i^{(l)} \rangle} . \tag{14}
\end{equation}

Instead of $p(x)$, one samples the auxiliary-fields from

\begin{equation}
\bar{p}(x) = p(x) \frac{\langle \psi_T | \hat{B}(x) | \phi_i^{(l)} \rangle}{\langle \psi_T | \phi_i^{(l)} \rangle} , \tag{15}
\end{equation}

which can be accomplished either directly for discrete fields using a heat bath-like approach or more generally via a force bias. This will automatically prevent the random walks from sampling any determinants with zero (or negative) overlap with the trial wave function. Those with larger overlap will be sampled more, although the weight from importance sampling will ensure that the exact distribution defined by Eq. (14) is sampled. The energy can be calculated by the mixed estimate similar to Eq. (10), although now with importance sampling it has the form:

\begin{equation}
E = \frac{\sum_{i} w_i^{(n)} \langle \psi_T | \hat{H} | \phi_i^{(n)} \rangle}{\sum_{i} w_i^{(n)} \langle \psi_T | \phi_i^{(n)} \rangle} . \tag{16}
\end{equation}

Following diffusion Monte Carlo (DMC), we refer to the quantity in the numerator, $E_L(\phi) \equiv \langle \psi_T | \hat{H} | \phi \rangle / \langle \psi_T | \phi \rangle$ as the local energy. An important characteristic of the constrained path approximation is that the mixed estimate in Eq. (10) is not variational.

The CP approximation has proved very accurate in the Hubbard model, especially for closed shell systems. For instance, the energy at $U = 4$ is typically within $< 0.5\%$ of the exact diagonalization result. It is, however, approximate. The systematic error in the energy tends to be larger in open-shell systems. Here we show that this error can be significantly reduced with trial wave functions which observe the correct symmetry.

\subsection{Release Constraint (RC)}

From a converged CP calculation, one can release the constraint and continue with the projection. Calculations of similar character have been done in the framework of DMC, under the name of released node. Since the CP result is already very close to the ground state and FP in AFQMC tends to have a reduced sign problem in general, one can expect that releasing the constraint in AFQMC will be effective and will allow the removal or reduction of systematic bias in more systems.

In principle the idea of releasing the constraint is straightforward. As mentioned, the RC calculation can theoretically be viewed as an FP calculation, with a much better starting point. Technically, however, the implementation of RC can be challenging. The initial population, namely that from CP, is obtained with importance sampling, which automatically imposes the constraint. On the one hand, the importance function must be modified in RC to allow the random walks to have a significant chance to sample the region with $\langle \psi_T | \phi^{(l)} \rangle$ being
negative (or develop different phases in the more general case). Indeed, the more efficient approach in FP in AFQMC seems to be with no importance function, as discussed earlier. On the other hand, since the initial population from CP has the form of Eq. (13). “undoing” the importance sampling will cause large statistical fluctuations and accelerate the onset of the sign problem. Thus we need to find the best way to interface the RC part with CP so as to minimize the growth of statistical noise.

RC calculations within AFQMC have been published by Sorella recently. Our approach is somewhat different and we will publish the method and further results elsewhere. The algorithmic details do not affect the issues that are the focus of the present paper, namely the use of symmetry properties to improve the calculations. A key aspect is our use of different forms of the HS transformation in the CP and the RC portions of the calculations. That is, we switch to a different HS decomposition in the RC in order to impose exact symmetry properties which drastically change the behavior of the RC calculations, as described below.

In RC calculations, we will use a mixed estimator similar to Eq. (10) to measure the energy:

$$E_{\text{RC}}(\beta) = \frac{\sum_{i} w_{i}^{\text{RC}} \langle \psi_{T} | H e^{-\beta \hat{H}} | \phi_{i}^{\text{CP}} \rangle}{\sum_{i} w_{i}^{\text{CP}} \langle \psi_{T} | e^{-\beta \hat{H}} | \phi_{i}^{\text{CP}} \rangle}.$$  

Thus $E_{\text{RC}}(\beta = 0) = E_{\text{CP}}$. As mentioned, the mixed estimate in CP is not variational. The RC energy will asymptotically converge to the exact ground-state energy. However, it can converge from below or above. Indeed, as we further discuss below, the convergence can be non-monotonic for poorer trial wave functions.

III. SYMMETRY IMPROVED AFQMC: RESULTS AND DISCUSSION

In this section we describe approaches to impose symmetry in AFQMC calculations, and study their effects on computational efficiency and, more importantly, on the sign problem. We divide the discussion into two parts: the choice of the Hubbard-Stratonovich transformation and its effect on symmetry, and symmetry in the trial wave function. While we will consider all three flavors of AFQMC introduced in Sec. II, our focus will be on CP and RC, since these are the most general methods which will allow calculations to scale to large system sizes. It is shown that the use of symmetry can lead to large reductions in statistical and systematic errors, and alleviate the sign problem.

A. Hubbard-Stratonovich Transformation

For each form of the two-body interaction, there are different ways to decompose the propagator, leading to different forms of Eq. (1). Decompositions based on Hartree, Fock, and pairing mean-fields are all possible. Even within each mean-field framework, the details can affect the final form of the one-body propagator and the computational efficiency. When a constraint is imposed, the form of the HS transformation chosen can affect the systematic accuracy given a form of the trial wave function. For the Hubbard interaction, for example, the most commonly used HS transformation involves discrete auxiliary-fields, due to Hirsch. The spin form of this decomposition is:

$$e^{-\Delta \tau U_{\text{it}} n_{i}^{\uparrow}} = e^{-\Delta \tau U(n_{i}^{\uparrow} + n_{i}^{\downarrow})/2} \sum_{x_{i}=\pm 1} \frac{1}{2} e^{\gamma x_{i}(n_{i}^{\uparrow} - n_{i}^{\downarrow})},$$

which results in an Ising-like auxiliary-field for each lattice site. The constant $\gamma$ is determined by

$$\cosh(\gamma) = e^{\Delta \tau U/2}. \quad (19)$$

It can also be mapped to a charge density form:

$$e^{-\Delta \tau U_{\text{it}} n_{i}^{\uparrow}} = e^{-\Delta \tau U(n_{i}^{\uparrow} + n_{i}^{\downarrow} - 1)/2} \sum_{x_{i}=\pm 1} \frac{1}{2} e^{\gamma x_{i}(n_{i}^{\uparrow} + n_{i}^{\downarrow} - 1)},$$

with

$$\cosh(\gamma) = e^{-\Delta \tau U/2}. \quad (20)$$
A more general HS transformation

\[ e^{\tilde{A}^2} = \frac{1}{\sqrt{2\pi}} \int_{-\infty}^{\infty} e^{-x^2/2 + \sqrt{\pi}\tilde{A}} dx, \]  

(22)

applies to any two-body operators written in the form of a square. To apply this to the Hubbard interaction, we write:

\[ n_{i\uparrow} n_{i\downarrow} = \frac{1}{2} [(n_{i\uparrow} + n_{i\downarrow} - \bar{n})^2 - \bar{n}^2 - (1 - 2\bar{n})(n_{i\uparrow} + n_{i\downarrow})] , \]  

(23)

where \( \bar{n} \) can take any value (including acquiring a dependence on \( \beta \)). We then let \( \tilde{A} = \sqrt{-\Delta \tau U/2} (n_{i\uparrow} + n_{i\downarrow} - \bar{n}) \) and use Eq. (22) to obtain an HS transformation with continuous fields. The constant \( \bar{n} \) can be thought of, physically, as a background term that one subtracts from the one-body operator prior to applying the HS transformation. This has been pointed out before for Hubbard interactions and for Coulomb interactions. As discussed below, the optimal choice for \( \bar{n} \) is to remove all background interactions from the mean-field, by minimizing the quadratic first term on the right-hand side in Eq. (23). Over typically densities, this choice leads to significant improvement over other choices including the standard Hirsch discrete decomposition.

In Fig. 1, we illustrate the effect of the background \( \bar{n} \) in the continuous charge decomposition of Eqs. (23) and (24). The logarithm of the statistical error bar is plotted versus projection time for different values of the background \( \bar{n} \). The calculations are all FP so the sign problem is present, as indicated by the growing error bars, which are essentially linear in the log-plot with projection time. It is seen that the minimum statistical error is achieved when \( \bar{n} = (n_{i\uparrow} + n_{i\downarrow})_{\text{MF}} = (N_{i\uparrow} + N_{i\downarrow})/L \). The efficiency of the decomposition decreases as \( \bar{n} \) deviates from the optimal value. It is a symmetric function of the deviation: a background value which is larger or smaller than the optimal value by the same amount gives comparable results. We point out that, although we have illustrated this with the repulsive model, the same applies to the attractive case. For example, in dilute Fermi gas simulations, Eqs. (23) with a small \( \bar{n} \) will be much more efficient than Eqs. (24) which corresponds to \( \bar{n} = 1 \).

It is often thought that the use of a discrete HS field, compared to continuous fields, leads to significant performance advantages. This is not the case, as shown in Fig. 1. The discrete charge decomposition of Eq. (23) is shown in the figure. We see that it is almost the same as the continuous decomposition with \( \bar{n} = 1 \). This is because the interaction term in the discrete charge decomposition is mapped to \( (n_{i\uparrow} + n_{i\downarrow} - 1) \), identical to the continuous transformation when \( \bar{n} \) is set to 1. The discrete decomposition is ideal near half-filling, but will be inefficient in dilute systems, for example, in Fermi gas simulations.

Because the decomposition in Eq. (22) preserves SU(2) symmetry, it can be more efficient than the discrete spin decomposition of Eq. (18), which is the most commonly used form in simulations of systems with repulsive interactions. This point is more subtle, however, as it is intertwined with the sign/phase problem. In Fig. 2 the
two decompositions are compared in a situation free of the sign problem, namely the half-filled repulsive Hubbard model. It can be seen that the Gaussian charge decomposition leads to much smaller statistical fluctuations. This has also been pointed out by Meng et. al, using the discrete charge decomposition of Eq. (20) at half filling.

In Fig. 2 we study the case when a phase problem is present ($U>0$, so $\tilde{A}$ is imaginary): $4 \times 4$, with $N_\uparrow = N_\downarrow = 7$ and $U = 8$. This system has a severe sign problem for the discrete spin decomposition. The continuous Gaussian decomposition leads to a phase problem. The latter decomposition initially has smaller error bars, benefiting from the preservation of symmetry, but after some time, its error bars exceed that of the spin decomposition. So in systems with a sign/phase problem, the efficiency is a balance of two competing aspects. On the one hand, the charge decomposition has an advantage for preserving symmetry. On the other hand, the phase problem tends to result in fast deterioration of the statistical signal and is a disadvantage. Below we discuss how to exploit these characteristics in different calculations.

We can use the advantage of the spin decomposition in longer time projections, as shown in Fig. 4 in a FP calculation of the periodic $8 \times 8$ supercell with $N_\uparrow = N_\downarrow = 13$ and $U = 4$. A FE trial wave function is used. The number of walkers is $5 \times 10^5$, with total 60 runs to obtain the average and estimate the statistical error bars. The red horizontal line gives the final CPMC result. The inset shows a magnified view of the last part of the projection. The spin decomposition calculation provides a reliable upper bound to the ground-state energy.

![FIG. 4: (Color online) Total energy versus projection time in FP calculations using discrete spin and continuous charge decompositions. The system is a $8 \times 8$ lattice under periodic boundary conditions, with $N_\uparrow = N_\downarrow = 13$ and $U = 4$. A FE trial wave function is used. The number of walkers is $5 \times 10^5$, with total 60 runs to obtain the average and estimate the statistical error bars. The red horizontal line gives the final CPMC result. The inset shows a magnified view of the last part of the projection. The spin decomposition calculation provides a reliable upper bound to the ground-state energy.](image)

Comparison, the CP calculation using a free-electron trial wave function gives $-66.857(2)$, as indicated by the line in Fig. 4 running for minutes on a single core. )

The charge decomposition can offer a significant advantage in short projection time, however. A main application of this is in RC calculations. Since we start from a population of a converged CP run, the initial state is close to the true ground state. One can expect a short projection in the RC calculation to recover a significant fraction of the correction to CP. In Fig. 5 we show an example RC calculation, for the same system as in Fig. 4. In this calculation, the CPMC portion always uses the standard spin decomposition, which has a severe sign problem. As can be seen from the inset, the CP energy obtained from the mixed-estimate is not variational $\frac{1}{2}$. (Typical CP calculations will run to much larger $\beta$ than shown in the main figure, in order to collect statistics.) In the RC portion, two different calculations are shown, one continuing to use the discrete spin decomposition while the other switching to the symmetry charge-decomposition. It is seen that the latter leads to much smaller statistical fluctuations and allows the RC calculation to reach convergence. The RC with discrete spin decomposition has much larger errors, and also displays a population control bias $\frac{5}{5}$. (The statistical error and the bias could, of course, be reduced by increasing the population size further.)

![FIG. 5: (Color online) Effect of symmetry decomposition in release constraint calculations. The main figure shows the convergence of CPMC energy with projection time, compared with the exact ground-state energy. The CPMC calculation uses the discrete spin decomposition. The inset shows RC calculations, starting from a converged CP state, using two different forms of the HS decomposition. The system is the same as in Fig. 4, $4 \times 4$, $N_\uparrow = N_\downarrow = 7$ and $U = 8$, using a symmetry multi-determinant trial wave function. The number of walkers is $1 \times 10^5$, with total 100 runs in the RC portion to collect statistics.](image)
B. Symmetry of the Trial Wave Function

In this section we discuss the other aspect of symmetry in AFQMC calculations, the use of trial wave functions which preserve symmetry. We generate the trial wave function with particular symmetries: total spin $S^z$, total momentum $\vec{K}$, rotational symmetry in momentum space $R$, mirror reflection $\sigma$ along the line $K_y = K_x$ in momentum space. When the total momentum $\vec{K} = 0$, we use the $C_{4v}$ point group irreducible representation to label different symmetry state. In the present paper, these properties are imposed in the trial wave function by a brute-force approach, making a linear combination of Slater determinants, as discussed in further details below.

1. Trial Wave Function in FP Calculations

Imposing the proper symmetry in the trial wave function can accelerate convergence and reduce the equilibration time in the FP calculations. As mentioned in Sec. II A the trial wave function is often also used to generate the initial population in FP. In all the FP calculations in this section, we use the HS transformation given in Eqs. (22) and (23), using optimal background values from simple mean-field calculations.

![Figure 6](image)

**FIG. 6:** (Color online) The effect of symmetry trial wave functions in FP calculations, compared to exact results in $4 \times 4$ lattices with $U = 4$: (a) $N_x = N_y = 8$, no sign problem; (b) $N_x = N_y = 7$, severe phase problem. The UHF trial wave function is generated by a UHF calculation with $U = 0.5$, $0s0kx0ky1D$ is a single-determinant non-interacting trial wave function with $S^z = 0$ and $k_x = k_y = 0$, and $0s0kx0kyMD$ is a multi-determinant trial wave function which has rotational symmetry in momentum space in (a) and $B1$ symmetry in (b), in addition to $S^z = 0$, $k_x = 0$, $k_y = 0$.

We illustrate the effect of the symmetry in the trial wave function in Fig. 6. In the top panel, we consider a half-filled system which is thus sign-problem-free. The effects of three trial wave functions are compared: the unrestricted Hartree-Fock (UHF) wave function, a single-determinant trial wave function formed by occupying $k$-states, and a multi-determinant trial wave function which preserves additional symmetry. The UHF trial wave function builds in correlation effect via its static antiferromagnetic order, and is an excellent starting point, as can be seen from the variational energy values at the beginning of the projection. However, it is contaminated by higher spin eigenstates, and the FP with UHF exhibits a long convergence time, as seen in the inset. (The effect of spin-contamination in AFQMC calculations has been discussed in continuum systems by Purwanto et. al.) The single-determinant FE trial wave function has a very high variational energy. Its statistical error bars are larger since the $|\psi_T\rangle$ in Eq. (10) to evaluate the energy is much poorer, and preserves fewer symmetry properties. However, it eventually leads to a faster convergence than the UHF trial wave function because symmetry has properly removed certain excitations. The multi-determinant trial wave function with symmetry, which we obtain by diagonalizing in the subspace of the open-shell [in the spirit of a small complete active-space self-consistent field (CASSCF) calculation], leads to rapid convergence and small statistical errors. In the bottom panel, Fig. 6b, we show an example when there is a severe sign-phase problem. The same trends are seen, with the full symmetry trial wave function leading to rapid convergence of the projection.

With the mixed estimate, the symmetry trial wave function on the left will have zero overlap with any wave function component in a different symmetry space. This allows one to project out explicitly lower energy states of different symmetry, and thus an opportunity to study excited states. This has been used in QMC calculations before. In the AFQMC formalism, the walkers are full Slater determinants, so that the symmetry projection can be done rigorously and explicitly for each walker. We illustrate excited state calculations in Fig. 7 where the converged FP results show excellent agreement with results from exact diagonalization.

2. Trial Wave Function in CP Calculations

In CPMC, the sign or phase problem is controlled by the sign or gauge condition of the overlap with the trial wave function. The condition is approximate, and the resulting systematic error depends on the trial wave function. Thus the trial wave function has an especially important role in CP calculations. In this section, we study how trial wave functions which preserve symmetries impact the accuracy in CP calculations. As shown in prior studies, the systematic error from the constraint is in general small for Hubbard-like systems, even when an FE or UHF trial wave function is used. In closed-shell systems in particular, the error is often negligible, as seen in the example of $8 \times 8$ system with 26 electrons.
The energy is plotted versus projection time for a $4 \times 4$ lattice with $N_{\uparrow} = 3$ and $U = 4$. The exact results for the ground state and the three excited state energies are shown for comparison. The symmetry of each energy level is labeled. The trial wave functions are chosen with the correct symmetry using multi-determinant. The error bars are shown but are smaller than symbol size at most points.

in Sec. III A The systematic errors tend to be larger for open-shell systems. As we show below, the leading reason for the problem in open-shell systems seems to be symmetry in the trial wave function. The use of trial wave functions with proper symmetry often leads to a dramatic reduction in the CP error.

We use the discrete spin decomposition in Eq. (18) in the CP calculations, which causes “only” a sign problem, even when a twist angle is applied in the boundary condition of the supercell $\mathcal{S}$. We first focus on small system sizes where exact results are available to make detailed and systematic comparison. Larger systems are treated later, and compared with our best results from RC calculations. In Fig. 3 we study the systematic error in the case of $4 \times 4$ with $N_{\uparrow} = N_{\downarrow} = 7$, which has the most severe sign problem in systems that can be diagonalized presently. We study the systematic error as $U$ goes from 0 to 12, spanning weak to moderate to strong interactions. The CP results with an FE trial wave function is shown. (We use a small twist of opposite sign for $\uparrow$ and $\downarrow$ spins to generate the FE trial wave function, which breaks the SU(2) symmetry, but has translational symmetry and $\overline{R} = 0$. ) The CP systematic error tends to grow with $U$, reaching about 2% of the total energy, or about 1% of the correlation energy at $U = 12$. As we see, the use of symmetry wave functions (obtained by diagonalizing the open-shell, leading to a total of 10 determinants) makes the CP systematic error very small across the range of $U$. Figure 4 contains a zoomed-in view, at $U = 8$, of the CP/SYM run and the subsequent RC which leads to an exact result. In Table I we compile the results from a variety of systems where exact diagonalization can be done to provide a quantitative measure. The CP results are compared for FE (or UHF solution obtained from a weak $U$) trial wave functions and symmetry trial wave functions. Significant improvement is seen in open-shell systems, and accurate results are obtained from CP calculations.

In Table I we have included a set of results for $4 \times 4$ and $N_{\uparrow} = N_{\downarrow} = 8$. There is no sign problem at half-filling. As has been discussed before, the CP calculations can be easily made exact at half-filling (or for negative $U$) by re-defining the importance sampling to have a non-zero minimum. However, if this were ignored and the CP algorithm applied to half-filling literally, an artificial constraint would result because the random walk cannot tunnel from one side of $\langle \Psi_T|\phi \rangle = 0$ to the other, even though both sides are positive. The calculated energy would then show a bias, which is visible in the results shown in the table. With symmetry trial wave functions, this bias is removed even when running CP unmodified, and the CP results at half-filling are accurate.

The improvement of CP calculations with the symmetry trial wave function is not just to the ground-state energy. The CP bias in the observable is also significantly reduced. An example is shown in Fig. 5 in which we calculate the structure factor of the spin-spin correlation function in the ground state:

$$S(K) = \frac{1}{N} \sum_{ij} S_i^z S_j^z \exp[iK(R_i - R_j)].$$

As mentioned earlier, we use the back-propagation technique to calculate correlation functions. The result is plotted for the same $4 \times 4$ systems for three different values of $U$. The peak at $(\pi, \pi)$ indicates strong
anti-ferromagnetic correlations. We see that the CP result using UHF trial wave function shows a larger anti-ferromagnetic order, because the UHF state itself overestimates the order. The symmetry trial wave function removes the bias and leads to results in agreement with exact diagonalization.

![Diagram](image_url)

FIG. 9: (Color online) The structure factor $S(k)$ of the spin-spin correlation function for three interaction strengths. The system is $4 \times 4$ with $N_x = N_y = 7$, and the horizontal axis labels of $K$ are in units of $\pi/2$. The symmetry trial wave function has $S^2 = 0$ and $K_x = K_y = 0$ and B1 symmetry. CPMC has 10,000 walkers, with back-propagation $\beta = 1$.

3. Trial Wave Function in RC Calculations

Formally the role of symmetry in the trial wave function in RC calculations is similar to that in FP. However it is intimately connected to the discussion in the previous section on CP, since the initial state in RC is the converged solution from CP. The symmetry trial wave functions improves the CP approximation and the quality of the wave function sampled from CP, as indicated by the improvement in the energy and in the calculated observables. This means symmetry trial wave functions also allow better RC calculations, by providing a better initial state and by giving a better trial wave function in the mixed estimate in Eq. (17). As discussed in Sec. IIA we also impose symmetry with the HS transformation in RC, by switching from the Ising spin form in the CP calculation to the Gaussian form in the RC part. We find this combination to improve the quality of the RC calculations greatly in most cases. An example is shown in Fig. 5 Results from RC/SYM calculations are also shown in Table II for systematic comparisons with CP and with exact diagonalization results.

Figure 10 illustrates the behavior of RC calculations using two different trial wave functions, the UHF versus a symmetry trial wave function. A small system size of $3 \times 3$ with $2 \uparrow$ and $2 \downarrow$ electrons is chosen such that the RC calculation can also be carried out explicitly to allow direct comparison. (In the explicit calculation, we propagate the CP population of $|\phi_{CP}\rangle$ directly by applying $e^{-\Delta \tau \hat{H}}$. The propagation is carried out by expanding each walker in terms of exact eigenstates of $\hat{H}$.) We see that CPMC/UHF gives an energy closer to the exact value ($\sim 0.1\%$ error) compared to CPMC/SYM ($\sim -0.3\%$ error). The corresponding RC/UHF moves further away from the exact answer and shows no indication of convergence in the imaginary-time span in which RC/SYM is well-converged. The explicit RC calculation, as shown in the inset, reveals a highly non-monotonic behavior. The projection does converge to the correct ground-state energy, but requiring an imaginary time of $> 100$. This would be impossible to reach in a QMC RC calculation because of the sign problem. Thus non-monotonic behaviors could be difficult to detect and would yield misleading results. The improvement with the symmetry trial wave function, which leads to rapid and monotonic convergence, is then especially valuable.

![Diagram](image_url)

FIG. 10: (Color online) RC calculations with symmetry trial wave functions and without. The system is $3 \times 3$ with $N_x = N_y = 2$ and $U = 4$. The symmetry trial wave function has $S^2 = 2, K_x = 0, K_y = 0$ while the UHF wave function breaks these symmetries. CP/UHF is very accurate, but RC/UHF has non-monotonic behavior and slow convergence, as shown by the explicit propagation. RC/SYM converges rapidly and monotonically. The explicit propagation (EP) result of RC/UHF is shown to large projection time in the inset.

The use of proper symmetry can allow RC calculations of excited states, similar to the discussion in Fig. 7. Since CP allows one to start from an initial population much closer to the exact exact state, RC can be more accurate. An example is shown in Fig. 11 in which the many-body ground state and first excited state energies are calculated as a function of crystal momentum. Both CP and RC are done with the same trial wave function, in which
TABLE I: Computed ground-state energy per site ($E/L$) from CP, with FE or UHF trial wave function (CP/FE) and with multi-determinant symmetry trial wave function (CP/SYM) respectively, compared with release-constraint (RC/SYM) and exact diagonalization results. RC/SYM uses the same symmetry trial wave function as in CP/SYM. The symmetry of the ground state is given in the last column. The statistical error bars in the QMC results are on the last digit and are shown in parentheses.

| $L$ ($N^{\uparrow}, N^{\downarrow}, U$) | CP/FE  | CP/SYM | RC/SYM | ED  | $(S^2, K_x, K_y)$ |
|-------------------------------|--------|--------|--------|-----|-----------------|
| $2 \times 2$ (2,1,4) | -1.60564(5) | -1.60615(3) | -1.60465(5) | -1.60463 | (0.75,0,1) |
| $2 \times 3$ (2,2,4) | -1.38328(6) | -1.40129(4) | -1.40085(4) | -1.40087 | (2,0,0) |
| $2 \times 3$ (2,2,8) | -1.2239(2) | -1.2463(1) | -1.2443(3) | -1.2442 | (0,0,0) |
| $2 \times 4$ (2,2,4) | -1.36839(2) | -1.37387(2) | -1.37379(3) | -1.37383 | (0,0,0) |
| $2 \times 4$ (3,3,4) | -1.56939(4) | -1.56942(4) | -1.56944(5) | -1.56941 | (0,0,0) |
| $3 \times 3$ (4,4,4) | -0.7783(2) | -0.8127(1) | -0.8091(1) | -0.8094 | (0,0,0) |
| $4 \times 4$ (2,2,4) | -1.36839(2) | -1.37387(2) | -1.37379(3) | -1.37383 | (0,0,0) |
| $4 \times 4$ (2,2,8) | -1.2239(2) | -1.2463(1) | -1.2443(3) | -1.2442 | (0,0,0) |
| $4 \times 4$ (2,2,12) | -1.0948(1) | -1.1048(1) | -1.1042(2) | -1.1044 | (0,0,0) |
| $4 \times 4$ (3,3,4) | -0.93394(2) | -0.94622(1) | -0.94598(1) | -0.94600 | (6,0,0) |
| $4 \times 4$ (3,3,8) | -0.9034(1) | -0.9208(1) | -0.9203(1) | -0.9202 | (6,0,0) |
| $4 \times 4$ (3,3,12) | -0.8334(1) | -0.8428(1) | -0.8386(3) | -0.8388 | (0,0,0) |
| $4 \times 4$ (4,4,4) | -1.0948(1) | -1.1048(1) | -1.1042(2) | -1.1044 | (0,0,0) |
| $4 \times 4$ (4,4,8) | -0.9293(1) | -0.9376(1) | -0.9329(5) | -0.9328 | (0,0,0) |
| $4 \times 4$ (4,4,12) | -0.8439(1) | -0.8557(1) | -0.8507(6) | -0.8512 | (0,0,0) |
| $4 \times 4$ (5,5,4) | -1.1012(1) | -1.1104(1) | -1.1084(2) | -1.1080 | (0,0,0) |
| $4 \times 4$ (5,5,8) | -0.9793(1) | -0.9863(1) | -0.9840(1) | -0.9840 | (0,0,0) |
| $4 \times 4$ (5,5,12) | -0.8334(1) | -0.8428(1) | -0.8386(3) | -0.8388 | (0,0,0) |
| $4 \times 4$ (6,6,4) | -1.1012(1) | -1.1104(1) | -1.1084(2) | -1.1080 | (0,0,0) |
| $4 \times 4$ (6,6,8) | -0.9293(1) | -0.9376(1) | -0.9329(5) | -0.9328 | (0,0,0) |
| $4 \times 4$ (6,6,12) | -0.8439(1) | -0.8557(1) | -0.8507(6) | -0.8512 | (0,0,0) |
| $4 \times 4$ (7,7,2) | -1.19584(2) | -1.19992(1) | -1.19822(2) | -1.19821 | (0,2,2) |
| $4 \times 4$ (7,7,4) | -0.9793(1) | -0.9863(1) | -0.9840(1) | -0.9840 | (0,0,0) |
| $4 \times 4$ (7,7,6) | -0.8334(1) | -0.8428(1) | -0.8386(3) | -0.8388 | (0,0,0) |
| $4 \times 4$ (7,7,8) | -0.7361(1) | -0.7461(1) | -0.7417(8) | -0.7418 | (0,0,0) |
| $4 \times 4$ (7,7,10) | -0.6687(2) | -0.6782(1) | -0.673(2) | -0.6754 | (0,0,0) |
| $4 \times 4$ (7,7,12) | -0.6202(2) | -0.6296(2) | -0.627(4) | -0.6282 | (0,0,0) |
| $4 \times 4$ (8,8,4) | -0.84225(6) | -0.85140(6) | -0.85133(6) | -0.85137 | (0,0,0) |
| $4 \times 4$ (8,8,8) | -0.51625(4) | -0.5293(2) | -0.5291(2) | -0.5293 | (0,0,0) |
| $4 \times 4$ (8,8,12) | -0.364(3) | -0.374(2) | -0.373(4) | -0.3745 | (0,0,0) |

\(^a\) $B_1$ symmetry is also used in $|\psi_T\rangle$.
\(^b\) $A_1$ symmetry is also used in $|\psi_T\rangle$.
\(^c\) $R_{x/2}|\psi_T\rangle = \exp(i3\pi/2)|\psi_T\rangle$ symmetry is also used.

The correct symmetry is imposed. Consistent with prior experience, CP is very accurate for the ground state, although systematic error is visible at larger twist angles. The CP result is less accurate for the excited state. With RC, the CP error is removed and the results are seen to be essentially exact.

**IV. DISCUSSION AND CONCLUSION**

In this paper, we have studied the role of symmetry in AFQMC calculations, and discussed the imposition of symmetry from two key aspects of an AFQMC calculation, namely the HS transformation and the trial wave function. It is shown that major improvements in efficiency and accuracy can be achieved. To allow detailed and systematic benchmark and analysis, we have used smaller lattice sizes extensively, where exact results are available. The method applies straightforwardly to larger systems. CP calculations will scale as a low power with system size; FP and RC will of course have a sign/phase problem, albeit at a much reduced level with the proposed symmetry improvements. In Table II, we present CPMC energies in several open-shell systems.
horizontal axis gives the first excited state energies versus crystal momentum. RC greatly improves the calculation of excited states and band structures. The system is $4 \times 4$ with $N_{\uparrow} = N_{\downarrow} = 5$ and $U = 4$. QMC statistical error bars are smaller than symbol size. The horizontal axis gives $|k|$ along a line cut $k_y = 2k_x$. Exact diagonalization (ED) results are shown for comparison. The line is to aid the eye.

**TABLE II: Energy per site in some open-shell Hubbard square lattices at $U = 4$.** The symmetry of the ground state, $S^z = 0$, $K_x = K_y = 0$, is preserved in the trial wave function used in CP/SYM and RC/SYM. The UHF trial wave function is generated with $U = 0.5$ as has been done before.

| L    | $(N_\uparrow,N_\downarrow)$ | CP/UHF       | CP/SYM       | RC/SYM       |
|------|----------------------------|--------------|--------------|--------------|
| 6x6  | (12,12)                    | -1.18444(3)  | -1.18625(3)  | -1.18525(4)  |
| 6x6  | (24,24)                    | 0.14899(2)   | 0.14709(3)   | 0.14899(4)   |
| 8x8  | (14,14)                    | -1.07173(1)  | -1.07239(1)  | -1.07180(2)  |
| 8x8  | (22,22)                    | -1.18580(2)  | -1.18673(2)  | -1.1858(2)   |
| 10x10| (40,40)                    | -1.11378(3)  | -1.11468(2)  | -1.1135(2)   |
| 12x12| (58,58)                    | -1.10912(3)  | -1.11015(3)  | -1.1089(2)   |

$^a$ $B_2$ symmetry is also used in $|\psi_T\rangle$.

The development presented in this paper will allow many applications even in its current form. Although we have focused on zero-temperature methods, many of the ideas will also apply to finite-temperature calculations. The formula can be directly mapped from the U>0 case we discussed to U<0 with a particle-hole transformation. The CP/SYM calculations and the RC/SYM from it represent a major step forward, as we can now have internal checks and a systematically improvable computational method capable of reaching two- and three-dimensions and large system sizes.

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