Permanent variational wave functions for bosons

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We study the performance of permanent states (the bosonic counterpart of the Slater determinant state) as approximating functions for bosons, with the intention to develop variational methods based upon them. For a system of \( N \) identical bosons, a permanent state is constructed by taking a set of \( N \) arbitrary (not necessarily orthonormal) single-particle orbitals, forming their product and then symmetrizing it. It is found that for the one-dimensional Bose-Hubbard model with the periodic boundary condition and at unit filling, the exact ground state can be very well approximated by a permanent state, in that the permanent state has high overlap (at least 0.96 even for 12 particles and 12 sites) with the exact ground state and can reproduce both the ground state energy and the single-particle correlators to high precision. For a generic model, we have devised a greedy algorithm to find the optimal set of single-particle orbitals to minimize the variational energy or maximize the overlap with a target state. It turns out that quite often the ground state of a bosonic system can be well approximated by a permanent state by all the criterions of energy, overlap, and correlation functions. And even if the error is apparent, it can often be remedied by including more configurations, i.e., by allowing the variational wave function to be a combination of multiple permanent states. The algorithm is used to study the stability of a two-particle system, with great success. All these suggest that permanent states are very effective as variational wave functions for bosonic systems, and hence deserve further studies.

I. INTRODUCTION

The Hartree-Fock approximation for fermions is a paradigm in quantum mechanics [1, 2]. Conceptually, it is very simple. It is a variational method. For an \( N \)-fermion system, one just takes \( N \) orthonormal single-particle orbitals \( \{\phi_i| 1 \leq i \leq N\} \), constructs the product state \( \phi_1(x_1)\phi_2(x_2)\ldots\phi_N(x_N) \), and then anti-symmetrizes it to obtain the Slater determinant wave function

\[
\Xi(x_1, x_2, \ldots, x_N) = \frac{1}{\sqrt{N!}} \sum_{p \in S_N} (-1)^p \phi_{P_1}(x_1)\phi_{P_2}(x_2)\ldots\phi_{P_N}(x_N)
= \frac{1}{\sqrt{N!}} \det(\phi_i(x_j)).
\]

(1)

Here \( S_N \) denotes the symmetric group of degree \( N \). By construction, the determinant state satisfies the anti-symmetry condition and constitutes a legitimate wave function for a collection of identical fermions. With the variational wave function built in this way, the rest work is an optimization problem. One has to choose the \( N \) orthonormal orbitals optimally so as to minimize the energy expectation value of the \( N \)-body variational state.

It is a natural idea to generalize this approach to bosons. One can take \( N \) single-particle orbitals \( \{\phi_i| 1 \leq i \leq N\} \), form their product, but then symmetrize it to obtain the following state,

\[
\Phi(x_1, x_2, \ldots, x_N) = \frac{1}{\sqrt{N!}} \sum_{p \in S_N} \phi_{P_1}(x_1)\phi_{P_2}(x_2)\ldots\phi_{P_N}(x_N)
= \frac{1}{\sqrt{N!}} \det(\phi_i(x_j)),
\]

(2)

which we shall refer to as a permanent state. Unlike the fermionic case, here because of the symmetry instead of anti-symmetry condition, the single-particle orbitals are not necessarily orthogonal to each other [12], but could even be identical. In the extremal case in which all the orbitals are constrained to be the same, we have the Gross-Pitaevskii approximation [8, 9], which has been proven to be very successful for weakly interacting Bose gases [10, 11]. However, for more general systems, such as the Bose-Hubbard model which we shall study below, the Gross-Pitaevskii approximation is too restrictive and we had better allow more freedom for the \( N \) orbitals.

The idea seems very simple. However, probably because mathematically the permanent of a matrix lacks many of the nice properties of the determinant, such an approach has rarely been put into practice. As far as we know, the very limited literature starts with two papers of Romanovsky et al. in 2004 and 2006 [12, 13]. They employed the permanent state as variational wave functions for some few-boson systems in two-dimensional harmonic traps. They went beyond the Gross-Pitaevskii approximation by allowing each particle to occupy a different orbital, for which they coined the term unrestricted Bose-Hartree-Fock approximation. However, because of the
perceived high complexity of the self-consistency equations, they did not seek self-consistent orbitals, but prescribed them as displaced Gaussians. Subsequently, the self-consistency equations for the orbitals were derived by Heimsoth [14, 15]. Unfortunately, the formalism was still unnecessarily complicated, and he did not even implemented the Ryser algorithm for permanent computation. Consequently, he could handle at most six particles.

In this paper, we resume research in this vein but with different perspectives. For some reason, people almost always look at orthonormal single-particle orbitals, and henceforth construct orthonormal Fock states as basis functions for a multi-particle system. But working with orthonormal orbitals is artificial—The bosons themselves have no notion of orthogonality and are totally happy to reside in non-orthogonal orbitals. Therefore, for studying the structure of a bosonic wave function, one should give up the obsession of orthogonal orbitals. It is absolutely possible that a bosonic wave function looking complex with respect to a Fock basis is actually of a simple and compact structure, namely, is equal or close to a permanent state built of non-orthogonal orbitals. As we shall see below, this is indeed the case for many models, in particular, for the one-dimensional Bose-Hubbard model at unit filling. An immediate implication of such fortunate facts is that permanent states can be used as building blocks for constructing variational wave functions and it is worthwhile to develop variational methods based upon them. Of course, besides this pragmatic purpose, permanent states are of interest in their own right. They are the simplest bosonic wave functions, yet of rich structures, with information (energy, correlation functions, etc.) not easy to extract sometimes.

This paper is organized as follows. First in Sec. II, we review the connection between the first and second quantization formalisms, and we shall establish some analytic facts about the permanent state. Then in Sec. II we show that for the one-dimensional Bose-Hubbard model with periodic boundary condition and at unit filling, which is the standard setting for studying the superfluid-Mott insulator transition, the permanent state can be a very good approximation of the exact ground state of the system. This enables us to use the algorithm to study the stability of a two-boson system, which is analogous to the negative ion of hydrogen. Finally, we conclude in Sec. VI with some open problems. We would like to mention that the whole paper is actually a by-product of studying these open problems.

II. PERMANENT WAVE FUNCTIONS

For the sake of simplicity, let us assume a finite-dimensional single-particle Hilbert space

$$\mathcal{H} = \text{span}\{|x\rangle, 1 \leq x \leq L\},$$

where $|x\rangle$ are orthonormal basis vectors. The associated creation (annihilation) operators will be denoted as $a_x^\dagger$ ($a_x$). They satisfy the usual commutation relations. A generic (not necessarily normalized) single-particle state or a single-particle orbital in this space is $|\phi\rangle = \sum_{x=1}^L |x\rangle\langle x|\phi\rangle = \sum_{x=1}^L \phi(x)|x\rangle$. The associated creation operator is $a_x^\dagger = \sum_{x=1}^L \phi(x)a_x^\dagger$.

For an $N$-boson system, the many-body Hilbert space is spanned by the orthonormal Fock states

$$|n\rangle = \frac{(a_1^\dagger)^{n_1}(a_2^\dagger)^{n_2} \cdots (a_L^\dagger)^{n_L}}{\sqrt{n_1!n_2! \cdots n_L!}} |\text{vac}\rangle,$$

where $n \equiv (n_1, n_2, \ldots, n_L)$ is an $L$-tuple with $n_x \geq 0$ and $\sum_{x=1}^L n_x = N$. The number of such Fock states or the dimension of the many-body Hilbert space is

$$D = \binom{N + L - 1}{N} = \frac{(N + L - 1)!}{N!(L - 1)!}.$$  \hfill (5)

A generic $N$-boson state $|\Psi\rangle$ expands as $|\Psi\rangle = \sum_n C(n)|n\rangle$. In first quantization, the same state is expressed by the wave function $\Psi(x_1, x_2, \ldots, x_N)$, with $1 \leq x_i \leq L$. Under the action of particle permutations, the coordinate tuples $x \equiv (x_1, x_2, \ldots, x_N)$ break into different equivalent classes labelled by the occupation tuple $n$. The wave function should be constant on each class. We say a coordinate tuple $x$ belongs to $n$ and denote...
it as $\mathbf{x} \vdash \mathbf{n}$ if in $\mathbf{x}$, the value $x$ appears $n_x$ times. The cardinality of the class $\mathbf{n}$ is $N! / \prod_{x=1}^{L} n_x!$, and thus by considering the norm of $\Psi$ in both the first and second quantization form, we have \[ \Psi(\mathbf{x}) = \frac{n_1! n_2! \ldots n_L!}{N!} C(\mathbf{n}), \quad \mathbf{x} \in \mathbf{n}. \]

With this formula, one can convert a wave function in the first quantization form to the second quantization form, and vice versa.

Now suppose we have a set of $N$ arbitrary orbitals $\{\phi_i(x), 1 \leq i \leq N\}$. The simplest symmetric $N$-particle wave function one can construct out of them is the permanent state according to \[ \varepsilon = \hat{S}(\phi_1, \phi_2, \ldots, \phi_N) \]

This form should remind us of the standard Fock states. However, such a perfect cancellation cannot occur for the permanent state in \[ \text{(2)} \] or \[ \text{(8)} \], which in turn means that $\Phi$ must be non-vanishing.

The second one is about uniqueness. Given a set of orbitals $\{\phi_i, 1 \leq i \leq N\}$, one can construct a permanent state according to \[ \Phi \], or \[ \Phi \] in the case of bosons. Here of course, two sets of orbitals should be deemed equivalent if they differ just by a permutation or some linear scaling. The answer is no as we have \[ \Phi = \hat{S}(f_1, f_2, \ldots, f_N) = \hat{S}(g_1, g_2, \ldots, g_N), \]

then for some permutation $\sigma \in S_N$, $f_i \propto g_{\sigma(i)}$ for all $1 \leq i \leq N$.

At least five simple facts about a permanent state can be easily established. Although some of them are not much used in this paper, we collect them all here for completeness.

In the fermionic case, because of a well-known property of the determinant, the determinant of \[ \Phi \] vanishes identically if the $N$ orbitals are not linearly independent. However, such a perfect cancellation cannot occur for the permanent state in \[ \text{(2)} \] or \[ \text{(8)} \] in the case of bosons. Although this might not be as surprising as the terms in \[ \text{(2)} \] are all of the same sign, we have formulated it as a proposition, which will be referred to later.

**Proposition 1.** The $N$-particle permanent state $\Phi$ constructed with $N$ nonzero orbitals $\{\phi_i, 1 \leq i \leq N\}$ according to \[ \text{(2)} \] is necessarily non-vanishing.

**Proof.** Consider the inner product of $\Phi$ with the condensate-type symmetric state

\[ \Omega = v(x_1)v(x_2) \ldots v(x_N), \]

where $v \in \mathcal{H}$ is an arbitrary single-particle state. We have

\[ \frac{1}{\sqrt{N!}} \langle \Phi | \Omega \rangle = \prod_{i=1}^{N} \langle \phi_i | v \rangle. \]

The product on the right hand vanishes if and only if $v$ is orthogonal to some $\phi_i$, or $v \in \bigcup_{i=1}^{N} \ker(\phi_i)$. Here by an abuse of notation, we have also used $\phi_i$ to denote the linear functional $\langle \cdot | \phi_i \rangle$. The kernel of this functional $\ker(\phi_i)$ is simply the hyperplane orthogonal to $\phi_i$. By the well-known folklore in mathematics that a vector space over $\mathbb{C}$ cannot be the union of a finite number of proper subspaces \[ \bigcup_{i=1}^{N} \ker(\phi_i) \]

we know the union \[ \bigcup_{i=1}^{N} \ker(\phi_i) \]

cannot cover the whole space $\mathcal{H}$, and for some $v$ the product is non-vanishing, which in turn means that $\Phi$ must be non-vanishing.

At least five simple propositions about a permanent state can be easily established. Although some of them are not much used in this paper, we collect them all here for completeness.

In the fermionic case, because of a well-known property of the determinant, the determinant of $\Phi$ vanishes identically if the $N$ orbitals are not linearly independent. However, such a perfect cancellation cannot occur for the permanent state in \[ \text{(2)} \] or \[ \text{(8)} \] in the case of bosons. Although this might not be as surprising as the terms in \[ \text{(2)} \] are all of the same sign, we have formulated it as a proposition, which will be referred to later.
Again by the well-known folklore that a vector space cannot be the union of a finite number of proper subspaces \[19\], we know there must be some \(i\) such that \(\ker(f_1) = \ker(f_j) \cap \ker(g_1)\), or \(\ker(f_1) = \ker(g_1)\), which means \(f_1 \propto g_1\). By permutation or relabeling, we can assume \(i = 1\) and simply \(f_1 = g_1\).

We then wish to factor out \(f_1\) and \(g_1\) in \(12\) to get

\[
\prod_{i=2}^{N} (f_i|v) = \prod_{i=2}^{N} (g_i|v)
\]

for all \(v \in \mathcal{H}\). This already holds for \(v \notin \ker(f_1)\). To show that it actually holds also for \(v \in \ker(f_1)\), consider \(v \in \ker(f_1)\) and \(w \notin \ker(f_1)\). For any nonzero \(t \in \mathbb{C}\), \(v + tw \notin \ker(f_1)\), otherwise \(w = [(v + tw) - v]/t\) would be in \(\ker(f_1)\). By \(14\), we have then

\[
\prod_{i=2}^{N} (f_i|v + tw) = \prod_{i=2}^{N} (g_i|v + tw)
\]

for all \(t \neq 0\). That is, two polynomials of \(t\) evaluate to the same value for all \(t \neq 0\). This means that the two polynomials are actually the same. In particular, their constant terms are identical. We have thus proven that \(14\) holds for all \(v \in \ker(f_1)\) too. The proposition is then proven by induction.

The second proof is more direct. Let \(|f_i\rangle = \sum_{j=1}^{L} f_{ij} |j\rangle\) and similarly \(|g_i\rangle = \sum_{j=1}^{L} g_{ij} |j\rangle\). By \(8\) and \(11\),

\[
\Phi = \prod_{i=1}^{N} \left( \sum_{j=1}^{L} f_{ij} a_j^{\dagger} \right) |\text{vac}\rangle = \prod_{i=1}^{N} \left( \sum_{j=1}^{L} g_{ij} a_j^{\dagger} \right) |\text{vac}\rangle .
\]

The fact that the \(a_j^{\dagger}\) operators commute and the Fock basis states are linearly independent implies the polynomial equality

\[
\prod_{i=1}^{N} \left( \sum_{j=1}^{L} f_{ij} z_j \right) = \prod_{i=1}^{N} \left( \sum_{j=1}^{L} g_{ij} z_j \right),
\]

where \(z_j\) are indeterminates. The proposition is proven by the unique factorization theorem of multivariate polynomials over \(\mathbb{C}\) \(21\).

To appreciate Proposition \(8\) one should note that for fermions, a Slater determinant state constructed out of \(N\) orbitals is determined by the subspace (a point on the so-called Grassmannian manifold \(21\)) spanned by the orbitals, not by the orbitals themselves. The orbitals are just a basis of the subspace. Another basis would yield the same Slater determinant state up to a global constant.

By Proposition \(8\), the permanent state \(\Phi\) is invariant under the permutation \(f_i \rightarrow f_{\sigma i}\), with \(\sigma\) being an arbitrary permutation, and the scaling \(f_i \rightarrow \lambda_i f_i\), with the scaling factors satisfying the condition \(\prod_{i=1}^{N} \lambda_i = 1\). This allows us to count the degrees of freedom \(22\) of a permanent state as

\[
d = NL - (N - 1) = N(L - 1) + 1.
\]

Apparently, like not every \(N\)-fermion state is a Slater determinant state, not every \(N\)-boson state is a permanent state. However, in the special case of \(\dim \mathcal{H} = L = 2\), this is indeed the case. We have

**Proposition 3.** Suppose the single-particle Hilbert space is of dimension 2, i.e., \(\dim \mathcal{H} = L = 2\), then every \(N\)-boson state is a permanent state.

**Proof.** In this case, the \(N\)-boson Hilbert space is of dimension \(N + 1\) and a basis is the Fock states \(\{|a_1^\dagger\}^i |\text{vac}\rangle, 0 \leq i \leq N\}. But by \(18\), the number of degrees of freedom of a permanent state is also \(N + 1\), so the proposition is anticipated by dimension counting. To prove it rigorously, we expand an arbitrary state \(\Psi\) as

\[
\Psi = (a_2^\dagger)^m \sum_{k=0}^{N-m} c_k (a_1^\dagger)^{N-m-k} (a_2^\dagger)^k |\text{vac}\rangle.
\]

Here \(m\) is an integer between 0 and \(N\), and \(c_0 \neq 0\). The fact that \(a_1^\dagger\) and \(a_2^\dagger\) commute motivates us to consider the polynomial \(P(z) = \sum_{k=0}^{N-m} c_k z^{N-m-k}\). Let it factorize as

\[
P(z) = c_0 \prod_{j=1}^{N-m} (z - c_j).
\]

Then it is easy to see that \(\Psi\) factorize as

\[
\Psi = c_0 (a_2^\dagger)^m \prod_{j=1}^{N-m} (a_1^\dagger - c_j a_2^\dagger) |\text{vac}\rangle.
\]

By \(8\), we see it is a permanent state and can read off the single-particle orbitals. \(\square\)

Proposition \(8\) means that for a two-site Bose-Hubbard model \(23\), which is a canonical model for studying the Bose-Josephson effect, any state is a permanent state. The proposition also reminds us of a similar proposition for fermions \(24, 25\). Although an arbitrary fermionic wave function is not generally a Slater determinant state, for the case of \(N\) fermions in \(L = N + 1\) orbitals, the wave function is necessarily a Slater determinant.

Proposition \(8\) is about the case when the dimension of the single-particle Hilbert space is minimal (but still nontrivial). Similarly, when the particle number is minimal (but still nontrivial), i.e., when \(N = 2\), we have

**Proposition 4.** Suppose \(N = 2\) and \(\dim \mathcal{H} = L\), then every \(N\)-boson state can be written as the sum of at most \([ (L + 1)/2 \] permanent states. Here \(|x\rangle\) is the floor function denoting the greatest integer less than or equal to \(x\).
Proof. For \( N = 2 \), the bosonic wave function \( \Psi(x_1, x_2) \) with \( 1 \leq x_i \leq L \) can be considered as a complex symmetric matrix. By the Autonne-Takagi theorem \([26]\), it can be factorized as

\[
\Psi(x_1, x_2) = \sum_{j=1}^{L} \sqrt{D_j} f_j(x_1) f_j(x_2), \tag{22}
\]

where \( f_j \) are a set of orthonormal functions and \( D_j \) are a set of non-negative numbers in decreasing order. The functions \( f_j \) are actually the so-called natural orbitals, i.e., eigenvectors of the one-body reduced density matrix \( \rho = 2\Psi\Psi^\dagger \) associated with \( \Psi \), and \( 2D_j \) are the occupation numbers. In the following, we shall assume that \( \Psi \) is normalized, i.e., \( \langle \Psi | \Psi \rangle = 1 \), so that \( \sum_{j=1}^{L} D_j = 1 \).

In the form of \( (22) \), the wave function \( \Psi \) is already a sum of \( L \) permanent states. Now for any pair \( j \neq k \), we can combine \( \sqrt{D_j} f_j(x_1) f_j(x_2) \) and \( \sqrt{D_k} f_k(x_1) f_k(x_2) \) into a single permanent state, i.e.,

\[
\hat{S}(u, v) = \frac{1}{\sqrt{2}} [u(x_1)v(x_2) + v(x_1)u(x_2)]
\]

with (note that by Proposition \( 2 \) this is essentially the only solution)

\[
u \equiv \frac{1}{\sqrt{2}} (\sqrt{D_j} f_j + i \sqrt{D_k} f_k), \quad v \equiv \frac{1}{\sqrt{2}} (\sqrt{D_j} f_j - i \sqrt{D_k} f_k).
\]

The proposition is then proven by noting that in \( (22) \), when \( L \) is even, we have \( L/2 \) pairs and when \( L \) is odd, we have \( (L - 1)/2 \) pairs and an extra unpaired term, which is already a permanent state. \( \square \)

It is easy to see that the number \( [(L + 1)/2] \) in Proposition \( 3 \) is not only sufficient but also necessary for a generic 2-boson state. For a generic state, the matrix \( \Psi \) is nonsingular or full-ranked. On the other hand, a permanent state is at most 2-ranked. Hence, by the subadditivity property of the rank of a matrix \( \text{rank}(A + B) \leq \text{rank}(A) + \text{rank}(B) \), we need at least \( [(L + 1)/2] \) permanent states to fully recover the original state. However, in practice, one might just need a sufficiently good approximation of the original state. The question is then, by taking the sum of \( M < [(L + 1)/2] \) permanent states, i.e., by constructing a state in the form of

\[
\Omega = \sum_{\alpha=1}^{M} \hat{S}(u^{(\alpha)}, v^{(\alpha)}), \tag{23}
\]

where \( u^{(\alpha)} \) and \( v^{(\alpha)} \) are arbitrary orbitals, to what extent can we approximate a target function \( \Psi \)? Quantitatively, what is the maximal value of the overlap

\[
O = \frac{||\Omega|\Psi|^2}{\Omega(\Omega)^*|\Psi^\dagger |\Psi|^2} \tag{24}
\]

achievable with such an \( M \)-configuration state? For this problem, we have

**Proposition 5.** Suppose \( N = 2 \), \( \dim \mathcal{H} = L \), and \( M \leq [(L + 1)/2] \). For the target state \( \Psi \) in \( \Omega \), the largest possible value of the overlap of an \( M \)-configuration state in the form of \( \Omega \) with it is

\[
O_{\text{max}} = \max_{\Omega} \frac{||\Omega|\Psi|^2}{\Omega(\Omega)^*|\Psi^\dagger |\Psi|^2} = \sum_{j=1}^{2M} D_j. \tag{25}
\]

**Proof.** Like \( \Psi \) in \( \Omega \), the \( M \)-configuration state \( \Omega \) in \( \Omega \) is also symmetric and can also be factorized as

\[
\Omega = \sum_{j=1}^{2M} \sqrt{C_j} \phi_j(x_1) \phi_j(x_2), \tag{26}
\]

where \( \phi_{1 \leq j \leq 2M} \) are a set of orthonormal vectors which can be extended into a complete orthonormal basis \( \phi_{1 \leq j \leq L} \), and \( C_j \geq 0 \) are ordered in decreasing order. Without loss of generality, let us assume \( \Omega \) is normalized so that \( \sum_{j=1}^{2M} C_j = 1 \). Note that by construction \( \Omega \) is at most of rank \( 2M \), and thus here in \( \Omega \) we have at most \( 2M \) nonzero terms. For the overlap between \( \Omega \) and \( \Psi \), we have

\[
O = \sum_{j=1}^{2M} |\langle \phi_j | \Psi \rangle|^2 \leq \sum_{j=1}^{2M} \sum_{k=1}^{L} |\langle \phi_j | \phi_k \rangle|^2 = \text{Tr}(A A^\dagger),
\]

where \( A \equiv V^\dagger \Psi W^\dagger \), with \( V \) the \( L \times 2M \) matrix whose columns are \( \phi_{1 \leq j \leq 2M} \), and \( W \) the \( L \times L \) matrix whose columns are \( \phi_{1 \leq k \leq L} \). Note that \( W \) is unitary. We have

\[
\text{Tr}(A A^\dagger) = \text{Tr}(V^\dagger \Psi W^\dagger W^T \Psi^\dagger V) \leq \sum_{j=1}^{2M} D_j. \tag{27}
\]

Here we used the Ky-Fan inequality \( \dagger \) for the matrix \( \Psi^\dagger \), which states the for an \( L \times L \) hermitian matrix, the sum of its \( m \) diagonal elements is less than or equal to the sum of its \( m \) largest eigenvalues, for all \( 1 \leq m \leq L \). We have thus proven that the overlap is upper bounded by \( \sum_{j=1}^{2M} D_j \). That this upper bound can be achieved is obvious, as we can just take the first \( 2M \) terms in \( \Omega \), and by Proposition \( 3 \) it is an \( M \)-configuration state. \( \square \)

In general, the occupation numbers \( D_j \) decrease fast and a truncation of \( \Omega \) with a very limited number of terms can yield a good approximation of the original state.
B. Basic formulae

A generic many-body Hamiltonian is of the form

\[ H = H_1 + H_2 = \sum_{i=1}^{N} K(i) + \sum_{1 \leq i < j \leq N} U(i,j). \tag{28} \]

Here the first sum is over each particle, with \( K(i) \) denoting the sum of the kinetic energy and the external potential of the \( i \)th particle, while the second sum is over each pair, with \( U(i,j) \) denoting the interaction between the \( i \)th and \( j \)th particle. The quantity of primary interest is the expectation value of \( H \) with respect to a permanent state \( |\Phi^{(2)}\rangle \), i.e.,

\[ E_{\text{var}} = \frac{\langle \Phi | H | \Phi \rangle}{\langle \Phi | \Phi \rangle} = \frac{\langle \Phi | H_1 | \Phi \rangle + \langle \Phi | H_2 | \Phi \rangle}{\langle \Phi | \Phi \rangle}. \tag{29} \]

It is straightforward to calculate the denominator and the numerator here. But for the sake of generality and in view of the extension to the multiconfigurational case below, let us consider the off-diagonal matrix elements instead of the diagonal matrix elements. Suppose \( \Phi^{(1)} \) and \( \Phi^{(2)} \) are two permanent states constructed with two sets of orbitals \( \{ \phi^{(a)}_{\alpha} \}_{1 \leq \alpha \leq N}, \alpha = 1, 2 \), i.e.,

\[ \Phi^{(a)} = \hat{S}(\phi^{(a)}_1, \phi^{(a)}_2, \ldots, \phi^{(a)}_N), \quad a = 1, 2. \tag{30} \]

Let us first define the \( N \times N \) overlap matrix of the orbitals,

\[ A_{ij} = \langle \phi^{(1)}_i | \phi^{(2)}_j \rangle, \quad 1 \leq i, j \leq N. \tag{31} \]

We have then

\[ \langle \Phi^{(1)} | \Phi^{(2)} \rangle = \frac{1}{N!} \sum_{P,Q \in S_N} \langle \phi^{(1)}_{P_1} \phi^{(1)}_{Q_1} \phi^{(1)}_{P_2} \phi^{(2)}_{Q_2} \ldots \phi^{(1)}_{P_N} \phi^{(2)}_{Q_N} \rangle = \sum_{R \in S_N} \langle \phi^{(1)}_{R_1} \phi^{(2)}_{R_1} \phi^{(1)}_{R_2} \phi^{(2)}_{R_2} \ldots \phi^{(1)}_{R_N} \phi^{(2)}_{R_N} \rangle = \text{per}(A). \tag{32} \]

Here per(\( A \)) denotes the permanent of the matrix \( A \). As for the matrix elements of \( H_1 \),

\[ \langle \Phi^{(1)} | H_1 | \Phi^{(2)} \rangle = \frac{N}{N!} \sum_{P,Q \in S_N} \langle \phi^{(1)}_{P_1} | K | \phi^{(2)}_{Q_1} \rangle \prod_{i=2}^{N} \langle \phi^{(1)}_{P_i} | \phi^{(2)}_{Q_i} \rangle = \sum_{i_1=1}^{N} \sum_{j_1=1}^{N} \langle \phi^{(1)}_{i_1} | K | \phi^{(2)}_{j_1} \rangle \text{per}(A; i_1 | j_1), \tag{33} \]

where per(\( A; i_1 | j_1 \)) denotes the permanent of the \( (N-1) \times (N-1) \) minor obtained by deleting the \( i_1 \)th row and the \( j_1 \)th column from \( A \). Similarly, for the matrix elements of \( H_2 \),

\[ \langle \Phi^{(1)} | H_2 | \Phi^{(2)} \rangle = \frac{N(N-1)}{2(N!)^2} \sum_{P,Q \in S_N} \langle \phi^{(1)}_{P_1} \phi^{(1)}_{P_2} | U | \phi^{(2)}_{Q_1} \phi^{(2)}_{Q_2} \rangle \prod_{i=3}^{N} \langle \phi^{(1)}_{P_i} | \phi^{(2)}_{Q_i} \rangle = \frac{1}{2} \sum_{i_1 \neq i_2, j_1 \neq j_2} \sum_{i_1=1}^{N} \sum_{j_1=1}^{N} \langle \phi^{(1)}_{i_1} \phi^{(1)}_{i_2} | U | \phi^{(2)}_{j_1} \phi^{(2)}_{j_2} \rangle \times \text{per}(A; i_1, i_2 | j_1, j_2). \tag{34} \]

Here in the last line, the summation \( \sum_{i_1 \neq i_2, j_1 \neq j_2} \) means that \( i_1 \) and \( i_2 \) both run from 1 to \( N \), but they must take different values. The second summation is interpreted similarly. By per(\( A; i_1, i_2 | j_1, j_2 \)) we mean the permanent of the \( (N-2) \times (N-2) \) minor of \( A \) obtained by deleting row \( i_1, i_2 \) and column \( j_1, j_2 \).

Above we see that to calculate the norm and physical expectation values of a permanent state, we have to calculate the permanent of the overlap matrix \( A \) and those of its minors. This is the price we have to pay for working with non-orthonormal orbitals.

As is generally believed, unlike the determinant of a matrix, the permanent of the matrix cannot be calculated in polynomial time. Currently, the best known general exact algorithm is the Ryser algorithm \( \text{[28]} \), which reduces the naive \( N \cdot N! \) evaluations to \( O(N^2 2^{N-1}) \). By using the Gray code, a further reduction by a factor of \( N \) can be achieved \( \text{[29]} \). This is the algorithm we use in this work \( \text{[30]} \). With this algorithm, it takes about 1 sec (0.01 sec) to calculate the permanent of a \( 22 \times 22 \) (15 \times 15, respectively) real-valued matrix on a commercial laptop computer and with MATLAB. We mention that in this paper, all calculation is done with MATLAB but without invoking the parallel computing toolbox.

III. BOSE-HUBBARD MODEL AT UNIT FILLING

To see whether a permanent wave function can be a good approximation of the ground state of a bosonic system, we take the one-dimensional Bose-Hubbard model with the periodic boundary condition and at unit filling as a case study. The Hamiltonian, as is often written in the second-quantization formalism, reads

\[ H = -\sum_{x=0}^{L-1} (a_x^d a_{x+1} + a_{x+1}^d a_x) + g \sum_{x=0}^{L-1} a_x^d a_x a_x a_{x+1}. \tag{35} \]

Here the single-particle Hilbert space \( \mathcal{H} \) is spanned by the orthonormal site (Wannier) states \( \{|x\} \), \( 0 \leq x \leq L - 1 \). The periodic boundary condition means \( |x\rangle = |x + L\rangle \). Note that we have taken the hopping strength as the unit of energy, and the Hamiltonian depends only on the parameter \( g \geq 0 \), which characterizes the on-site interaction strength.
For our purpose, it is often more convenient to work 
with the first-quantization formalism of \(28\). The corresponding \(K\) and \(U\) operators have matrix elements as

\[
K_{x,x'} = -(\delta_{x,x'+1} + \delta_{x,x'-1}),
\]

\[
U_{x_1x_2x'_1x'_2} = g\delta_{x_1x_2}\delta_{x_1x'_2}\delta_{x_1x'_1},
\]

In this section, we shall confine ourselves to the unit filling case, namely, the case when the particle number \(N = L\). The commensurate condition and the translation symmetry suggest putting the \(i\)th particle in an orbital centered at site \(i\), with all the orbitals of the same shape and related to each other by translations. Specifically,

\[
\phi_i(x) = \phi(x - i), \quad 1 \leq i \leq N,
\]

for some function \(\phi\). Here the periodic boundary condition requires \(\phi(x) = \phi(x + L)\). The total variational wave function (VWF) is then determined by the single-particle orbital \(\phi\). To construct an \(L\)-periodic function \(\phi\), we can choose an arbitrary primitive function \(\chi\) defined on the whole axis \(-\infty < x < +\infty\), and form the superposition

\[
\phi(x) = \sum_{j = -\infty}^{\infty} \chi(x - jL).
\]

We have tried two types of primitive functions, i.e., the Lorentz function and the exponential function,

\[
\chi^{(l)}(x; \lambda) = (1 + \lambda^2 x^2)^{-1},
\]

\[
\chi^{(e)}(x; \lambda) = e^{-\lambda |x|},
\]

where \(\lambda \geq 0\) is a parameter controlling the width of the functions. For these two simple types of primitive functions, the summation in (39) can be carried out analytically and yields

\[
\phi^{(l)}(x; \lambda) = \frac{1 - e^{-2\pi \lambda}}{1 - e^{-4\pi \lambda}},
\]

\[
\phi^{(e)}(x; \lambda) = \frac{\lambda e^{-\lambda x} + e^{-\lambda (L - x)}}{1 - e^{-\lambda L}}, \quad 0 \leq x \leq L.
\]

Note that in the limit of \(\lambda \to 0\), both \(\phi^{(l)}\) and \(\phi^{(e)}\) reduce to the zero-momentum Bloch state on the periodic lattice, while in the opposite limit of \(\lambda \to \infty\), both of them reduce to the Kronecker delta function \(\delta_{x,0}\).

The strategy is then simply to vary the parameter \(\lambda\) and calculate the variational energy \(E_{\text{var}}(\lambda; g)\) as a function of \(\lambda\) by using the formulae in Sec. II B. Here and henceforth, the dependence of \(E_{\text{var}}\) on the primitive orbitals should be understood tacitly. By the variational principle, \(E_{\text{var}}\) is always above the exact ground state energy \(E_{\text{gs}}\). The concern is whether the minimum of \(E_{\text{var}}\) can be sufficiently close to \(E_{\text{gs}}\). A case study with \(N = L = 10\) and \(g = 4\) is shown in Fig. 1. We see that for both types of primitive orbitals, at some value of \(\lambda\), \(E_{\text{var}}\) dips towards the horizontal line indicating \(E_{\text{gs}}\). At the minima, the relative error is 2% and 4%, respectively, for the Lorentz-type and exponential-type orbital. This is very encouraging and turns out to be typical.

By determining the minimum of the curve \(E_{\text{var}}(\lambda; g)\) for a fixed value of \(g\), we can get a variational estimate (denoted as \(E_{\text{var}}\)) of the ground state energy and an optimal permanent variational wave function \(\Phi_e\). In Fig. 2,
the variational energy $E_{\text{var}}$ is compared with the exact ground state energy $E_{\text{gs}}^\text{ext}$ obtained by exact diagonalization (ED) [32]. We see that over the full range of $0 \leq g < \infty$, the variational estimates agree with the exact values very well. The discrepancy is apparent only for the exponential-type variational wave function in the region around $g = 4$, with a relative error about 5.6%.

However, in this region, the Lorentz-type wave function is a much better approximation, reducing the relative error to about 2%. The general observation is that for $g \lesssim 8$, the Lorentz-type wave function yields a better upper bound for the ground state energy, while for $g \gtrsim 8$, the exponential-type wave function is better, although the latter fact is hardly visible in Fig. 2. Here in passing, we mention that if we take the Gross-Pitaevskii approximation, the orbital occupied by all the particles should be the zero-momentum Bloch state, as this choice minimizes the kinetic energy and the interaction energy simultaneously. The energy per particle would be $-2 + g(L - 1)/L$, i.e., linear in $g$, which is qualitatively wrong for large values of $g$.

Other than energy, a more stringent test for the accuracy of the variational wave function is its overlap with the exact ground state. We have thus Fig. 3(a), in which the overlap between the exact state $|\Psi_\text{gs}\rangle$ is obtained by ED, and the energy-minimizing (hence the subscript $e$) VWF $|\Phi_e\rangle$, is shown as a function of $g$. We see that in the full range, the overlap is at least 0.78 for the exponential-type VWF for a system as large as $N = L = 12$, and this number is even higher (0.95) for the Lorentz-type VWF. To appreciate these numbers, one should note that the dimension of the many-body Hilbert space is as large as $1.352 \times 10^8$. We also note that Fig. 3(a) and Fig. 2 are consistent with each other. In Fig. 3(a), all the curves show minima in the proximity of $g = 4$. This is exactly where the variational energies deviate most significantly from the exact one in Fig. 2. In this region, the Lorentz-type VWF has a much higher overlap than the exponential-type VWF with the exact ground state and accordingly, in this region the former has a lower energy as shown in Fig. 2. On the other hand, when $g \gtrsim 8$, the exponential-type VWF becomes better by the overlap criterion and accordingly, its energy is lower as shown in Fig. 2. The fact that in the small-$g$ region, the Lorentz-type VWF wins over the exponential-type VWF while in the large-$g$ region, the exponential-type VWF takes over might be understandable in view of the superfluid-Mott insulator transition. The large-$g$ region corresponds to the insulator phase, in which because of the strong particle-particle repulsion, each particle tends to be localized in its own site and the tunneling into neighboring sites should be exponentially small. The small-$g$ region corresponds to the superfluid phase, in which the particles are more mobile and a more extended orbital should be more appropriate.

So far, we have been taking the energy minimizing state $\Phi_e$ among either class of VWFs as an approximation of the exact ground state $|\Psi_\text{gs}\rangle$. Usually, this is the only thing one can do if the exact ground state is unavailable. However, if the exact ground state is available in a certain way, say, by exact diagonalization as we do here, a natural alternate approximation of it should be the variational state having maximal overlap with it. Let us denote it as $\Phi_o$, with the subscript meaning overlap. Hence, we have two related by different optimization problems. One is energy minimization and the other overlap maximization. There is no reason that the two solutions $\Phi_e$ and $\Phi_o$ should be the same, and by definition, we have $|\langle \Psi_\text{gs}|\Phi_e\rangle|^2 \leq |\langle \Psi_\text{gs}|\Phi_o\rangle|^2$ necessarily. In Fig. 3(b), we show $|\langle \Psi_\text{gs}|\Phi_o\rangle|^2$ as a function of $g$. In comparison with Fig. 3(a), we see that all the curves shift upwards as...
expected. For the exponential-type states, the increase of the overlap is quite apparent. For instance, while in Fig. 3(a), the minimum of \( |\langle GS|\Phi_o \rangle|^2 \) is about 0.78 for \( N = L = 12 \), in Fig. 3(b), the minimum of \( |\langle GS|\Phi_o \rangle|^2 \) is about 0.85. This strongly indicates that the two optimization problems are related but really different. For the Lorentz-type states, the increase of the overlap is less apparent but still visible. For \( N = L = 12 \), the minimum of the overlap increases from 0.95 to 0.96. From the curves in Fig. 3 by extrapolation one can infer that even for a system as large as \( N = L = 20 \), with a many-body Hilbert space of dimension about \( 6.9 \times 10^{10} \), the minimal values of the overlaps \( |\langle GS|\Phi_e \rangle|^2 \) and \( |\langle GS|\Phi_o \rangle|^2 \) would be about 0.9 if we take the Lorentz-type orbital. These numbers are very impressive.

We also note that in Fig. 3 all the curves, regardless of the primitive orbital type or the criterion, show minima in the vicinity of \( g = 4 \). This should be anticipated in view of the superfluid-Mott insulator transition. According to previous works \cite{33, 34}, the transition occurs at about \( g_c = 3.61 \). Close to the transition, the exact ground state should be most complex and it is hardest to approximate it with some simple functions.

In hindsight, the large overlap between the variational states and the exact ground state should be reasonable. There are at least three reasons that are in favor of such a welcome result. First, both VWFs can reproduce the exact ground state in either limit of \( g = 0 \) and \( g = \infty \). Second, it is well-known that for an arbitrary value of \( g \), by the Perron-Frobenius theorem \cite{26}, the ground state is strictly positive everywhere in the Fock-state basis. This property is shared by both VWFs by construction. Third, it is also known that the ground state belongs to the trivial representation of the symmetry group of the model (the dihedral group), or more specifically, it is invariant under all translations and reflections, a property again shared by both VWFs by construction.

Finally, as yet another check of the quality of the VWFs, we consider the single-particle correlator \( \langle a^\dagger_o a_x \rangle \). This expression is convenient for the exact ground state, which is obtained by ED in the Fock-state basis. For the VWFs, which are in the first-quantization formalism, we note that the corresponding single-particle operator \( \hat{C} \) has matrix elements \( \langle m|\hat{C}|n \rangle = \delta_{m,0}\delta_{n,x} \), and we can use (32) and (33) to calculate its expectation value

\[
\langle a^\dagger_o a_x \rangle = \sum_{i,j=1}^N \phi_i^*(0)\phi_j(x) \frac{\text{per}(A;ij)}{\text{per}(A)}. \tag{42}
\]
In Fig. 3, $\langle a_i^\dagger a_x \rangle$ is plotted against $x$ for the energy minimizing state $\Phi_e$ and the exact ground state $|GS\rangle$. We see a picture consistent with Fig. 2 and Fig. 3. In Fig. 3(a)-(c), when $g$ is small and the lorentz-type VWF is better, the correlator predicted by the lorentz-type VWF is very close to the exact one. In Fig. 3(c)-(d), when $g$ gets large, the exponential-type VWF is better, and accordingly the correlator predicted by the exponential-type VWF is close to the exact one. For any value of $g$, either the lorentz-type or the exponential-type VWF will be a good approximation by all the three criterions. Here it is also interesting to note that while the correlator predicted by the lorentz-type VWF is close to the exact one, for any value of $g$, the exponential-type VWF always underestimates the correlator, the lorentz-type VWF slightly underestimates it in the small-$g$ region, while overestimates it in the large-$g$ region. This might be related to the superfluid-Mott insulator transition.

In Fig. 4, $\langle a_i^\dagger a_x \rangle$ is plotted against $x$ for the overlap-maximizing states $\Phi_o$. It is normal to expect that $\Phi_o$ reproduces the correlation function better than $\Phi_e$. This is indeed the case. We see that for $g \leq 6$, when $|\langle GS|\Phi_o\rangle|^2$ is significantly higher than $|\langle GS|\Phi_e\rangle|^2$, the curves of $\langle a_i^\dagger a_x \rangle$ shift closer to the exact curve for both types of orbitals. In particular, the curves with the Lorentz-type orbitals almost coincide with the exact curves. For $g \geq 8$, $|\langle GS|\Phi_o\rangle|^2 \simeq |\langle GS|\Phi_e\rangle|^2$, indicating that $\Phi_e \simeq \Phi_o$, we do not see any significant change of the curves.

**IV. OPTIMIZATION ALGORITHM**

In the proceeding section, we have seen that for the one-dimensional Bose-Hubbard model with the periodic boundary condition and at unit filling, it is possible to construct some permanent state out of some simple orbitals to approximate its exact ground state very well. This is checked by examining the variational energy, the overlap with the exact ground state, and the single-particle correlation function.

The close approximation is achieved with some preasigned orbitals depending on a single parameter $\lambda$. A natural question is whether the numbers can be further improved by allowing more freedom of the orbitals. This leads to two optimization problems. First, for a given bosonic system with Hamiltonian $H$, how can we find a set of orbitals $\{\phi_i, 1 \leq i \leq N\}$, such that the energy expectation value of the permanent state $\Phi$ constructed in (2)

$$E = \frac{\langle \Phi | H | \Phi \rangle}{\langle \Phi | \Phi \rangle},$$

is minimized? Second, for a given normalized bosonic wave function $|\Psi\rangle$, how can we find the permanent state which is the optimal approximation of it? That is, how can we find the permanent state as in (2) such that the overlap

$$O = \frac{\langle \Phi | \Psi \rangle}{\langle \Phi | \Phi \rangle}$$

is maximized?

There exists a common simple strategy for both problems [24, 25]. For clarity, let us focus on the first problem for the present. Let us fix $N - 1$ orbitals, say, the orbitals $\phi_{2 \leq i \leq N}$, and try to find an optimal $\phi_1$. To this end, we note that with the orbitals $\phi_{2 \leq i \leq N}$ fixed, the numerator and denominator in (43) are both hermitian forms of $\phi_1$. That is, one can find operators $F$ and $G$ such that

$$\langle \phi_1 | F | \phi_1 \rangle = \langle \Phi | H | \Phi \rangle$$

and

$$\langle \phi_1 | G | \phi_1 \rangle = \langle \Phi | \Phi \rangle.$$  

We then can rewrite the ratio as

$$E(\phi_1) = \frac{\langle \phi_1 | F | \phi_1 \rangle}{\langle \phi_1 | G | \phi_1 \rangle},$$

(45)

By definition, $F$ and $G$ are hermitian single-particle operators depending on the orbitals $\phi_{2 \leq i \leq N}$. Importantly, $G$ is even positive definite as long as $\phi_{2 \leq i \leq N}$ are all nonzero, as by definition $\langle \phi_1 | G | \phi_1 \rangle = \langle \Phi | \Phi \rangle \geq 0$, with the equality achieved only if $\phi_1 = 0$. Here we recall Proposition 1 which asserts that $\Phi$ is necessarily nonzero if $\phi_{2 \leq i \leq N}$ are nonzero. It is a straightforward but lengthy calculation to derive the explicit expressions of $F$ and $G$, so we defer it to the Appendix. Suppose we have prepared the operators $F$ and $G$ (this is the most time-consuming part of the iteration). The optimal $\phi_1$ that will minimize the ratio in (45) is just the solution of the following generalized eigenvalue problem

$$\tilde{F}\phi = \epsilon \tilde{G}\phi$$

(46)

corresponding to the smallest eigenvalue $\epsilon_{\min}$, and the minimum of the ratio is just $\epsilon_{\min}$. Note that we just need the smallest eigenvalue. Therefore, we can resort to the Lanczos algorithm.

Once we have updated $\phi_1$, we can turn to $\phi_2$, and then to $\phi_3$, and so on. However, for convenience of programming, we can just make a circular shift of the orbitals $\phi_i \rightarrow \phi_{i-1}$, and continue to update $\phi_1$. In this process, the variational energy decreases monotonically, and as it is lower bounded by the exact ground state energy, it will definitely converge.

We have to mention that (46) is essentially the self-consistency equation derived by Heimsloth before by the method of performing variational differentiation of abstract Hilbert space vectors [14, 15]. However, hopefully here our different point of view has led to a more compact and transparent formalism. As we shall see in the subsection below, this formalism allows easy extension to the multiconfiguration case, which was not considered previously.

Now it should be clear that the second problem can be treated similarly. The ratio (44) can be written as

$$O = \frac{\langle \phi_1 | \gamma | \gamma | \phi_1 \rangle}{\langle \phi_1 | G | \phi_1 \rangle},$$

(47)

where the single-particle orbital $\gamma$ is defined by the summation or partial contraction

$$\gamma(x) = \sqrt{\frac{N}{L}} \sum_{x_2, \ldots, x_N=1}^{L} \Psi(x, x_2, \ldots, x_N) \prod_{i=2}^{N} \phi_i^* (x_i).$$

(48)
Once \( \gamma \) is calculated (again, this is the most time-consuming part), the optimal \( \phi_1 \) can be obtained by solving a generalized eigenvalue equation similar to (16), with \( \{ \gamma \} \) replacing \( \tilde{F} \).

Naively, the summation in (18) has the complexity of \( L^{N-1} N \), as \( x_{2 \leq i \leq N} \) run independently from 1 to \( L \). However, one should note that \( \Psi \) is invariant under permutations of \( x_{2 \leq i \leq N} \). Making use of this fact and making the dummy variables from \( x_i \) to \( y_{i-1} \), (18) can be written as

\[
\gamma(x) = \sqrt{N!} \sum_{y} \Psi(x, y) \frac{\per(\phi^*(y))}{n(y)!},
\]

where the summation is over the ordered \((N - 1)\)-tuple \( y \equiv (y_1, y_2, \ldots, y_{N-1}) \) with \( 1 \leq y_1 \leq y_2 \leq \cdots \leq y_{N-1} \leq L \), and \( \phi^*(y) \) denotes the \((N - 1) \times (N - 1) \) matrix with its \( i \)th row being the \( y_i \)th row of the \( L \times (N - 1) \) matrix \((\phi_2^*, \phi_3^*, \ldots, \phi_N^*)\). In the denominator, \( n(y)! \equiv \prod_{i=1}^{N-1} n_i! \), with \( n_i \) denoting the times \( i \) appears in \( y \). The computational complexity is now on the order of \((L^{N-2}N)^2 \). \( N \)

A. The multiconfiguration case

So far, we have assumed a single configuration. This is yet the only case people have considered [12-15]. For better approximation, one can try \( M > 1 \) sets of orbitals \( \{ \phi_1^{(\alpha)} \}, 1 \leq \alpha \leq M, 1 \leq i \leq N \), and let the variational wave function be the sum of the permanent wave functions constructed by each set of orbitals. Specifically,

\[
\Phi = \sum_{\alpha=1}^{M} \Phi_{\{ \alpha \}} = \sum_{\alpha=1}^{M} S(\phi_1^{(\alpha)}, \ldots, \phi_N^{(\alpha)}).
\]

Fixing the orbitals \( \phi_{2 \leq i \leq N} \) in each set, the variational energy [19] can be written as

\[
E = \frac{\sum_{\alpha, \beta=1}^{M} \langle \phi_{\{ \alpha \}} | H | \phi_{\{ \beta \}} \rangle}{\sum_{\alpha, \beta=1}^{M} \langle \phi_{\{ \alpha \}} | \phi_{\{ \beta \}} \rangle} = \frac{\sum_{\alpha, \beta=1}^{M} \langle \phi_{\{ \alpha \}} | \tilde{F}(\alpha) | \phi_{\{ \beta \}} \rangle}{\sum_{\alpha, \beta=1}^{M} \langle \phi_{\{ \alpha \}} | \tilde{G}(\alpha \beta) | \phi_{\{ \beta \}} \rangle},
\]

where \( \tilde{F}(\alpha \beta) \) and \( \tilde{G}(\alpha \beta) \) depend on the fixed orbitals \( \phi_{2 \leq i \leq N}^{(\alpha)} \) and \( \phi_{2 \leq i \leq N}^{(\beta)} \), and can be calculated with essentially the same formulae as in the Appendix—Just add the superscript \( \alpha \) to the orbitals in the bras and \( \beta \) to the orbitals in the kets. It is easily seen that

\[
(\tilde{F}(\alpha \beta))^\dagger = \tilde{F}(\beta \alpha), \quad (\tilde{G}(\alpha \beta))^\dagger = \tilde{G}(\beta \alpha).
\]

Apparently, (51) can be cast in the same form as (15), if we identify \( \phi_1 \) as the concatenated vector

\[
\phi_1 \equiv (\phi_1^{(1)}; \phi_1^{(2)}; \ldots; \phi_1^{(M)}),
\]

which is of length \( LM \), and define the block operators \( \tilde{F} \equiv (\tilde{F}(\alpha \beta)) \) and \( \tilde{G} \equiv (\tilde{G}(\alpha \beta)) \), which are of size \( LM \times LM \). The same update and iteration procedures can then be carried out.

Similarly, in the multiconfiguration case, the overlap (11) can be written as

\[
O = \frac{\sum_{\alpha, \beta=1}^{M} \langle \phi_{\{ \alpha \}} | \gamma(\alpha) \rangle \langle \gamma(\beta) | \phi_{\{ \beta \}} \rangle}{\sum_{\alpha, \beta=1}^{M} \langle \phi_{\{ \alpha \}} | \tilde{G}(\alpha \beta) | \phi_{\{ \beta \}} \rangle},
\]

where the single-particle orbital \( \gamma(\alpha) \), like \( \gamma \) in (18), is defined by the same summation with \( \phi_{\{ \alpha \}} \) replacing \( \phi \). Again, the same strategy as above can be applied.

B. A pitfall with \( N = 2 \)

A pitfall is to be avoided in implementing the multiconfiguration scheme. In the single-configuration case, the hermitian operator \( \hat{G} \) is strictly positive definite as long as the orbitals \( \phi_{2 \leq i \leq N} \) are nonzero. This is because by definition \( \langle \phi_1 | \hat{G} | \phi_1 \rangle = \langle \Phi | \Phi \rangle \), and by Proposition 1 \( \Phi \) is nonzero if \( \phi_{1 \leq i \leq N} \) are nonzero. In contrast, in the multi-configuration case (50), we do not necessarily have \( \Phi \neq 0 \) even if all the orbitals \( \{ \phi_1^{(\alpha)} \}, 1 \leq \alpha \leq M, 1 \leq i \leq N \) are nonzero—the configurations could cancel each other out. This happens particularly in the two-particle case of \( N = 2 \). For illustration, let us consider the two-particle, two-configuration case. For any value of \( \{ \phi_2^{(1)}, \phi_2^{(2)} \} \), if we choose \( \{ \phi_1^{(1)}, \phi_1^{(2)} \} \) as \( \{ \phi_2^{(2)}, -\phi_2^{(1)} \} \),

\[
\Phi = \hat{S}(\phi_1^{(1)}, \phi_1^{(2)}) + \hat{S}(\phi_2^{(1)}, \phi_2^{(2)}) = \hat{S}(\phi_2^{(2)}, -\phi_2^{(1)}) - \hat{S}(\phi_2^{(1)}, \phi_2^{(2)}) = 0.
\]

That the total wave function \( \Phi \) could vanish means that \( \hat{G} \) in the multi-configuration case is just positive semi-definite—it could have zero eigenvalues. Theoretically, this does not cause any problem because when the denominator of (51) vanishes, its numerator vanishes too. In other words, the eigenvectors of \( \hat{G} \) with the zero eigenvalue are also eigenvectors of \( \hat{F} \) with the zero eigenvalue. In numerics, one has to restrict \( \hat{F} \) and \( \hat{G} \) to the subspace spanned by the eigenvectors of \( \hat{G} \) with nonzero (hence positive) eigenvalues. This can be easily implemented once \( \hat{G} \) is diagonalized (but a better approach is described below).

In practice, this cautious extra effort is necessary only for \( N = 2 \). In our extensive numerical simulations, we have never encountered a case of \( \hat{G} \) becoming singular for \( N \geq 3 \). The reason is yet to be understood but it is not surprising as when \( N \geq 3 \), for fixed \( \phi_2^{(\alpha)} \), unlike (55), it is hard to find \( \phi_1 \equiv (\phi_1^{(1)}; \phi_1^{(2)}; \ldots; \phi_1^{(M)}) \) to make \( \Phi \) vanish; likely there is no solution. But for \( N = 2 \), it occurs necessarily. With \( M \) configurations, generally \( \hat{G} \) has \( \left( \frac{M}{2} \right) \) eigenvectors with the eigenvalue zero. This number can be understood in view of (55). For fixed orbitals \( \phi_2^{(\alpha)} \), one can find \( \left( \frac{M}{2} \right) \) linearly independent vectors \( \phi_1 \) such that \( \Phi \) vanishes.
The setting is a one-dimensional Bose-Hubbard model in a harmonic trap. The Hamiltonian is

\[ H_{\text{trap}} = -\sum_{x=-L_0}^{L_0} (\hat{a}_x^\dagger \hat{a}_x + \text{h.c.}) + \frac{g}{2} \sum_{x=-L_0}^{L_0} \hat{a}_x^\dagger \hat{a}_x^\dagger \hat{a}_x \hat{a}_x + \kappa \sum_{x=-L_0}^{L_0} x^2 \hat{a}_x^\dagger \hat{a}_x. \]  

(58)

Here \( \kappa > 0 \) is the stiffness of the harmonic potential. For symmetry, we have assume a lattice of size \( L = 2L_0 + 1 \). We take the open boundary condition.

We start from \( MN \) random orbitals [see Eq. (50)], where \( M \) is the configuration number and \( N \) is the particle number. Essentially, we just generate an \( ML \times N \) matrix with each element chosen randomly from the interval \([0,1]\) according to the uniform distribution. The orbitals (the columns) are then updated in a circular way. The variational state after \( n \) updates (steps) will be denoted as \( \Phi_n \), and its energy will be denoted as \( E_n \). Note that \( E_n \) is obtained simultaneously in solving the generalized eigenvalue problem (46).

By construction, \( E_n \) decreases monotonically and will definitely converge. The concern is in which way and with what rate it converges to its limit \( E_{\infty} \). This is studied in Fig. 6, where for a set of values of the parameters \( (N, L, \kappa, g) \) and with \( M = 1 \) or 2, some typical trajectories of \( E_n - E_{\infty} \) (here we just approximate \( E_{\infty} \) by some \( E_n \) with a large enough \( n \)) are displayed. For each value of \( M \), we have three trajectories corresponding to three different sets of initial orbitals.

We see that often the trajectory is not very regular—It is neither pure exponential nor pure power law, but clearly divides into different parts. In many cases, after some relaxation or transition stage, which can last for a long time, the energy eventually enters an exponentially decreasing mode. In our extensive numerical simulations, the observation is that the trajectory of the energy depends not only on the model and the model parameters, but also on the initial conditions and can differ significantly from run to run. We cannot draw any definite rule for the convergence rate of the energy, but the feeling is that the convergence tends to be slower with more configurations and weaker interactions.

1. The non-interacting case

Possibly the most embarrassing and surprising thing is that the convergence is slowest in the non-interacting limit. In this trivial case, the exact ground state \( |\text{GS}\rangle \) is simply a condensate-type (and hence a permanent) state with all particles occupying the single-particle ground state. We do not need to invoke the algorithm for its calculation, however, if we do, the convergence is as slow as a power law. In Figs. 7(a) and 7(b), with \( g = 0 \) but the other parameters the same as in Fig. 6, trajectories of the energy error \( E_n - E_{\infty} \) and the infidelity \( 1 - |\langle \text{GS} | \Phi_n \rangle|^2 \) are shown respectively for a particular run. Here we take

C. Convergence of the algorithm

We take a concrete model to illustrate the convergence behavior of the algorithm. Consider a one-dimensional Bose-Hubbard model in a harmonic trap. The Hamiltonian is

\[ \hat{H}_{\text{trap}} = -\sum_{x} (\hat{a}_x^\dagger \hat{a}_x + \text{h.c.}) + \frac{g}{2} \sum_{x} \hat{a}_x^\dagger \hat{a}_x^\dagger \hat{a}_x \hat{a}_x + \kappa \sum_{x} x^2 \hat{a}_x^\dagger \hat{a}_x. \]

The variational state after \( n \) updates (steps) will be denoted as \( \Phi_n \), and its energy will be denoted as \( E_n \). Note that \( E_n \) is obtained simultaneously in solving the generalized eigenvalue problem (46).

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$E_n$ to be the exact ground state energy. In either figure, the curve drops down steeply at about $n \sim N$, and afterwards it follows a straight line in the log-log plot. Basically, the picture is that after the first round of update, i.e., when each orbital has been updated once, the variational wave function is already very close to the exact state (the overlap is over 0.99 in the particular case). Afterwards, it improves slowly by a power law.

The exact reason behind the slow convergence is yet to be understood. Here we just emphasize that the convergence is slow only in the asymptotic sense. In the initial phase, the algorithm can already deliver the state $\Phi_n$ close enough to its limit $\Phi_\infty = | GS \rangle$.

In Figs. 7(c) and 7(d), we show two snapshots of the constituent orbitals, the particle density distribution $\rho(x, x)$, and the correlation function $\rho(0, x)$, which are respectively the diagonal and off-diagonal parts of the single-particle reduced density matrix $\rho$ defined as

$$\rho(x_1, x_2) = \langle a_{x_2}^\dagger a_{x_1} \rangle. \quad (59)$$

We see that even for an $n$ as small as $n = N = 3$, the permanent variational state $\Phi_n$ can already reproduce the exact values of $\rho(x, x)$ and $\rho(0, x)$ to high precision. The intriguing thing is that while the total wave function is already very close to the exact many-body ground state in terms of energy, overlap, and some most relevant correlation functions, the constituent orbitals are still far away or at least visibly different from the exact single-particle orbital. In view of Proposition 2 which states that different sets of orbitals necessarily result in different many-body wave functions, the current observation implies that the latter is not necessarily very sensitive to perturbations of the former in some cases. This in turn implies that it might not be a good idea to use convergence of the orbitals as a criterion in determining the termination of the algorithm [15].

2. Local minima

It should be no wonder that the algorithm can get stuck in a local minimum like many other greedy algorithms. This is illustrated in Fig. 8(a) with a model of site Bose-Hubbard and some specific value of $g$. We see that many trajectories of $E_n$ settle down on a secondary minimum. Note that for clarity, we have shown only the first 100 steps, but actually the horizontal lines extend all the way up to $n = 1500$.

In Fig. 8(b), we plot the possible eventual values of the variational energy $E_n$ and the overlap $|\langle GS | \Phi_n \rangle|^2$ as functions of $g$. For each value of $g$, like in Fig. 8(a), we have run the algorithm 20 times, each time up to $n = 1500$. We see that for $g$ smaller than some critical value $g_c \approx 8$, we get only a single value for either of $E_\infty$ and $|\langle GS | \Phi_\infty \rangle|^2$, which indicates that there is only a global minimum, however for $g$ larger than $g_c$, we get two different values for either of $E_\infty$ and $|\langle GS | \Phi_\infty \rangle|^2$, which indicates the presence of a second, local minimum.

Our experience is that local minima are ubiquitous. Generally, their number increases with the number $M$ of configurations. To avoid them and enhance the probability of hitting the global minimum, we simply run the algorithm multiple times, say 20 times for $M = 4$ configurations.

3. Real versus complex

So far, we have assumed the optimal orbitals to be real. For many systems with the time reversal symmetry, the many-body ground state is real and the assumption that the optimal orbitals should also be real seems very reasonable. However, this is not the case.

We take a minimal model to illustrate the possibility that the optimal orbitals could be complex although the total wave function is real. Consider a two-particle, two-site Bose-Hubbard model, i.e., a model with $(N, L) =$
FIG. 8. (Color online) (a) Twenty trajectories of the variational energy $E_n$. Each trajectory starts with a different set of randomly generated initial orbitals. (b) Possible limiting values of the variational energy $E_\infty$ and the overlap $\langle GS|\Phi_\infty \rangle^2$. Note that for each value of $g$, we have 20 runs as in (a) and in each run the number of iteration steps is 1500. The setting is a one-dimensional Bose-Hubbard model in a harmonic trap with the Hamiltonian (58). The fixed parameters are $(N, L, \kappa, M) = (3, 13, 0.2, 1)$.

FIG. 9. (Color online) Trajectories of the energy of the permanent variational state with either (a) real or (b) complex orbitals. In each case, we have five runs starting with five different sets of random orbitals. The number of configurations is $M = 1$. The horizontal dotted lines indicate the exact ground state energy. The setting is a two-particle, two-site Bose-Hubbard model with the Hamiltonian (60). The parameter $g = -1$.

(2, 2). The Hamiltonian is

$$H_{ds} = -(a_1^\dagger a_2 + a_2^\dagger a_1) + \frac{g}{2}(a_1^\dagger a_1 a_1^\dagger a_1 + a_2^\dagger a_2 a_2^\dagger a_2). (60)$$

Consider a real wave function $\Psi(x_1, x_2)$, with $x_{1,2} = 1, 2$, of this model. By Proposition 9 or Proposition 10, $\Psi$ can be written as a permanent state

$$\Psi = \hat{S}(u, v)$$

(61)

with two orbitals $u$ and $v$. The question is whether $u, v$ can both be real. Componentwise, (61) means

$$\Psi(1, 1) = \sqrt{2}u_1v_1, \quad \Psi(2, 2) = \sqrt{2}u_2v_2,$$

$$\Psi(1, 2) = \frac{1}{\sqrt{2}}(u_1v_2 + v_1u_2).$$

We have then

$$\Delta \equiv \Psi(1, 2)^2 - \Psi(1, 1)\Psi(2, 2) = \frac{1}{2}(u_1v_2 - v_1u_2)^2.$$

We thus see that for (61) to have real solutions, a necessary condition is that the quantity $\Delta$ be non-negative. It is easy to verify that this is also sufficient. Hence, when $\Delta < 0$, in the single configuration approximation, the wave function $\Psi$ can be exactly recovered with complex orbitals but not with real orbitals. Note that such a condition is satisfied by a cat-type state, which can be realized as the ground state of the model (60) with an attractive on-site interaction $g < 0$. In Fig. 9 we show the variational energies of the ground state of such a model, calculated with either real or complex orbitals. We see that while complex orbitals can deliver the exact value, real orbitals miss it with some overestimation. Therefore, we see that at least theoretically, complex orbitals are superior to real orbitals for energy minimization. However, the observation is that for all the models we consider in this paper, if the interaction is repulsive ($g \geq 0$) and if we take the single configuration approximation ($M = 1$), the complex approach leads to identical results with the real approach. More specifically, in these circumstances, even if we start with complex orbitals, the iteration will result in real orbitals. The reason is yet to be understood. For multiple configurations ($M > 1$), with complex orbitals, often we do get lower energies, however, the improvement is not very significant.
thus often confine ourselves to real orbitals in the following. Anyway, real arithmetics are four times fast than complex arithmetics, and the disadvantage in accuracy can be compensated by including more configurations.

V. APPLICATION OF THE ALGORITHM

A. The Bose-Hubbard model revisited

With the numerical optimization algorithm above, we can handle more general models. But let us start from the Bose-Hubbard model with the periodic boundary condition and at unit filling, and see how much it can improve over the results in Sec. [II]

For this particular model, we take a single configuration (\(M = 1\)). We never encounter any local minimum, and the symmetry of the model is perfectly preserved by the optimal orbitals. That is, although we always start from random orbitals, the orbitals we eventually get are always of the same shape and differ from each other just by translations, as described by [35].

In Fig. [11] we show the estimated ground state energy \(E_{gs}\) and the overlap \(|\langle GS|\Phi_e\rangle|^2\) obtained with unrestricted orbitals. For comparison, also shown are the results with Lorentz or exponential orbitals. We see that in the region \(g \leq 6\), the VWF with unrestricted orbitals does not improve much over the VWF with Lorentz orbitals neither by energy nor by overlap. Accordingly, the predicted correlation function \(\langle a_0^\dagger a_x \rangle\) is close to that predicted by the Lorentz VWF, as can be seen by comparing Figs. [11](a)-(c) with Figs. [4](a)-(c). However, in the region \(g \geq 6\), unrestricted orbitals do lead to improvement over both the Lorentz and the exponential orbitals, both by the criteria of energy and overlap. For instance, at \(g = 8\) and with \(N = L = 12\), the overestimate in energy (difference between the variational and exact ground state energy) reduces from 0.0127 (Lorentz) and 0.0106 (exponential) to 0.0039 (unrestricted), and simultaneously the deficiency in overlap \((1 - |\langle GS|\Phi_e\rangle|^2)\) reduces from 0.2023 (Lorentz) and 0.0203 (exponential) to 0.0026 (unrestricted). These numbers indicate that the algorithm produces really good approximation of the exact ground state. Indeed, as Figs. [11](d)-(f) show, now the VWF-predicted correlation function \(\langle a_0^\dagger a_x \rangle\) almost coincides with the exact values for \(g \geq 8\).

Overall, in Figs. [10] and [11] we see that for a system as large as \(N = L = 12\) and in the whole range of \(g\), the ground state can be very well approximated by a permanent state. This is very impressive in view of the dimension of the many-body Hilbert space, which is as large as \(D = 1352078\). The exact ground state is obtained by exact diagonalization and is a vector of this size with the Fock states as a basis. In contrast, the permanent variational state is constructed with \(N = 12\) orbitals, each of which is a vector of size \(L = 12\). That the orbitals are not even independent but related to each other by translations means that the permanent state is encoded with a \(12 \times 12\) circulant matrix with only 12 independent variables. From the data compression point of view, with the permanent state as an approximation of the exact ground state, the compression ratio is very high while the fidelity is still very good.

B. More general models

We now turn to more general models. First of all, let us break the periodic boundary condition of the Bose-Hubbard model above and replace it with the open boundary condition. The Hamiltonian is

\[
H_{obc} = - \sum_{x=-L}^{L-1} (a_x^\dagger a_{x+1} + \text{h.c.}) + \frac{g}{2} \sum_{x=-L}^L a_x^\dagger a_{x+1} a_x a_{x+1} \tag{62}
\]

We still assume unit filling, so the particle number \(N = L = 2L_0 + 1\). By the mere change of the boundary condition, the translation symmetry is lost and now the orbitals should differ in shape. It is then unclear what orbitals to choose to construct the permanent state—We have to resort to the numerical algorithm.

In Fig. [12](a), with \(N = L = 11\), the permanent estimated ground state energy is compared with the exact diagonalization result, and in Fig. [12](b), the overlap between the permanent variational state \(\Phi_e\) and the exact ground state \(|\Phi_e\rangle\) is shown. We see that like the periodic boundary condition case, across the full range of the on-site interaction \(g\), the permanent state is a very good approximation of the exact ground state. The overlap is as large as 0.964 even in the worst case, and the relative error in energy is at most 2%.

In Fig. [13] for three different values of \(g\), we show the constituent orbitals, the density distribution function \(\rho(x, x)\) and the correlation function \(\rho(0, x)\). For each value of \(g\), we start from \(N = L = 12\) random orbitals, and then update each orbital 100 times. In the top panels, we see that besides the bulk orbitals which are similar to each other in shape, there are two edge orbitals, which are maximal on the edges and decay into the bulk. As the permanent state is very close to the exact ground state by overlap, these orbitals provide a very good picture of the exact ground state. We also see that the permanent state predicted values of the density distribution and the correlation function agree with the exact results very well.

As a second example, let us consider the one-dimensional Bose-Hubbard model in a harmonic trap, with the Hamiltonian of [63]. In Fig. [14] we show the variational ground state energy \(E_{gs}\) and the overlap \(|\langle GS|\Phi_e\rangle|^2\), calculated with various numbers of configurations and with either real or complex orbitals, as functions of \(g\). We see that in the single configuration case \((N = M = 1)\), the real and complex approaches agree with each other exactly. The discrepancy between the variational energy and the exact value is apparent, and the overlap drops to 0.76 at \(g = 15\). However, the situation improves dramatically if we take \(M = 3\) configurations.
FIG. 10. (Color online) (a) Energy per particle of the energy-minimizing permanent states $\Phi_e$ constructed with either the Lorentz-type (40a) or the exponential-type (40b) primitive orbitals, or unrestricted orbitals. The solid line indicates the exact ground state energy calculated by exact diagonalization (ED). (b) Overlap between the energy-minimizing permanent states $|\Phi_e\rangle$ and the exact ground state $|GS\rangle$. In this figure, the parameters are $N = L = 12$.

With three configurations, both the real and the complex approaches get the ground state energy so accurate that the difference with the exact value is hardly visible in the figures. Accordingly, the overlap $|(GS|\Phi_e)^2$ is very close to unity throughout the range of $g$. Actually, the minimal value of the overlap is as large as 0.995 in the two figures. This means that with three configurations, be the orbitals real or complex, we can recover the exact ground state to very high precision. In the intermediate case of $M = 2$ configurations, the energy and the overlap are in-between. A peculiarity is that with real orbitals, the curve of the overlap is discontinuous at some point. This is due to the existence of local minima. At the critical point, two local minima change order in energy, or more precisely, the originally global minimum is surpassed by another minimum which was originally just a local one.

In Figs. 16 and 17, which correspond to $M = 1$, $M = 2$, and $M = 3$, respectively, we show the (real) orbitals $\phi_i^{(m)}$, the density distribution $\rho(x,x)$, and the correlation function $\rho(0,x)$ for three different values of $g$. In Fig. 15, we see that the single configuration approximation, in accord with Fig. 14, is quantitatively not very accurate for large values of $g$. However, the orbitals are consistent with the fermionization picture in the large-$g$ limit [35–37]. In Fig. 16, with two configurations, the similarity between the variational results and exact results improves, but the difference is still apparent for $g = 10$ and 15. A notable feature of the two-configuration approximation is that it breaks the parity symmetry of the model—the density distribution $\rho(x,x)$ and the correlation function $\rho(0,x)$ are asymmetric. This kind of phenomena is quite common in the conventional Hartree-Fock approximation, and is equally so with us. Below we shall see more examples. In Fig. 17, we have $M = 3$ configurations, and now the variational results almost coincide with the exact results, as is anticipated by the energy and overlap information in Fig. 14.

In the three models above, we see that the permanent approach with a very limited number of configurations can yield very accurate results. The observation is that...
regardless of the strength of the interaction, this is often the case if the system is not very dilute, or more precisely, if the ratio $N/L_{\text{eff}}$ is not too small, where by $L_{\text{eff}}$ we mean the effective volume of the system, i.e., the volume accessible to the particles. In the Bose-Hubbard model at unit filling, regardless of the boundary condition, the ratio is 1; in the Bose-Hubbard model in a harmonic trap above, although the lattice is of size $L = 13$, by the shapes of the orbitals or the density distribution, we infer that the effective lattice size $L_{\text{eff}} \approx 7$, so the ratio is about $3/7$.

Our experience is that the most challenging situation for the permanent variational approach is a dilute gas in the Tonks-Girardeau limit, in which the system is dilute (i.e., $N \ll L_{\text{eff}}$) and the on-site interaction $g$ is strong. We consider such a Bose-Hubbard model in Fig. 18, where we have $N = 4$ particles on a flat, open lattice of size $L = 13$. When $g$ is as small as 1, the single-configuration approximation can get the density distribution $\rho(x, x)$ and the correlator $\rho(0, x)$ accurately. As $g$ increases to 2, the error becomes visible but is still small. However, if $g$ further increases to 4, the discrepancy becomes very apparent.

The natural remedy is to take multiple configurations. In Fig. 19, we show how the variational energy and the overlap improve as the number of configurations increases. The improvement is steady but slow in comparison with Fig. 14. At $g = 4$, which is small by the scale of Fig. 19, even with $M = 5$ configurations, the deviation of the variational energy from the exact value and the deviation of the overlap from unity are still visible. In Fig. 20, we show how the situation in Fig. 18(c) improves by taking more and more configurations. We see that unlikethe situation in Fig. 17 with $M = 3$ configurations, the variational curves are still manifestly different from the exact ones. Only with $M = 5$, do the two almost coincide.

The trend of the curves in Fig. 19 suggests that for even larger values of $g$, we would need even more configurations to get a good approximation of the exact ground state. In Fig. 21 we examine the case of $g = 15$. For this value of $g$, the exact ground state is already very close to its fermionization limit at $g = +\infty$. We see that the single configuration approximation fails blatantly and it even breaks the parity symmetry of the model. With $M = 5$ configurations, the situation improves but only with $M = 10$ configurations do the variational predicted density distribution and correlation function agree with the exact values very well.

We thus see that generally taking multiple configurations can effectively reduce the error. The concern is how much price we have to pay. From Sec. IV A we see that with $M$ configurations, the number of blocks that we have to calculate for preparation of $\hat{F}$ and $\hat{G}$ increases by a factor of $\frac{1}{2}M(M + 1)$. This polynomial growth is mild.

C. Symmetry breaking and restoration

It is a common observation in the practice of Hartree-Fock approximation that the solution often spontaneously breaks the symmetries of the Hamiltonian $\hat{H}$. This could also happen with us, as we have seen in Fig. 19 and Fig. 21 above, where the permanent variational states do not respect the $\mathbb{Z}_2$ symmetry of the models.

We have two options to restore symmetry, i.e., to construct a state sharing the same symmetry with the exact ground state. The first approach is by brute force, we can simply take more configurations. Hopefully, the accuracy of the approximation will improve and the symmetry is restored alongside. This happens in Fig. 17 and Fig. 21(c). The second approach is based on the group representation theory. We can construct a projection operator corresponding to the irreducible representation of the exact ground state, and let it act on the variational state. The resultant state is also a permanent variational state, but generally with more configurations.

Below we take two concrete models to illustrate and compare the two approaches. In the first model, we have $N = 3$ bosons in a double-well potential. The Hamiltonian is

$$H_{\text{dw}} = H_{\text{obc}} + \sum_{x=-L_0}^{L_0} V(x)a_x^\dagger a_x,$$

where the double-well potential $V(x)$ is the superposition of a harmonic trap and a Gaussian bump, i.e.,

$$V(x) = kx^2 + \hbar \exp(-x^2/\sigma^2),$$
FIG. 13. (Color online) The orbitals $\phi_i(x)$, the particle density distribution $\rho(x, x)$, and the one-particle correlator $\rho(0, x)$ for three different values of $g$. The circles are for the permanent state, while the * markers are for the exact diagonalization results. The setting is a one-dimensional Bose-Hubbard model with the open boundary condition as defined in (62). The fixed parameters are $(N, L) = (11, 11)$.

where $\kappa$, $h$, and $\sigma$ are parameters. The system has a $Z_2$ symmetry, and it can be easily proven that the parity of the ground state is even and consequently the particle density distribution and the correlator are both even functions of $x$. However, in Fig. 22(a1), in the single configuration approximation, we see that the orbitals are apparently asymmetric, with one orbital located in the left well and the rest two in the right well. Consequently, the particle density and the correlator are both asymmetric, and the deviation from the exact values is significant.

To reinstall symmetry by the first approach, we can simply take one more configuration as in Fig. 22(c), where we get symmetric orbitals, and the variational results agree with the exact ones perfectly. We can also try the second approach. Let the permanent state in Fig. 22(a) be $\Phi_e = \hat{S} (\phi_1, \phi_2, \phi_3)$. An even-parity state can then be easily constructed as

$$\Phi_e = \hat{S} (\phi_1, \phi_2, \phi_3) + \hat{S} (\hat{P} \phi_1, \hat{P} \phi_2, \hat{P} \phi_3),$$  

where the inversion operator $\hat{P}$ is defined as $(\hat{P} \phi)(x) = \phi(-x)$. The newly constructed variational state $\Phi_e$ consists of two configurations, with the new configuration transformed from the old one by inversion. The physical quantities calculated with the projected state $\bar{\Phi}_e$ are shown in Fig. 22(b). We see significant improvement over the pro-projection state $\Phi_e$ in Fig. 22(a). Qualitatively, the symmetry is restored; quantitatively, the overlap with the exact ground state has increased from 0.91786 to 0.99255, and the predicted ground state energy per particle has decreased from 0.98983 to 0.97049 (the exact value is 0.95831). Of course, by construction, we do not expect $\bar{\Phi}_e$ to be optimal in energy among all the two-configuration variational states. Indeed, its energy is slightly higher than that of the optimal state in Fig. 22(c). However, its advantage is that it is obtained for free and is still a fairly good approximation.

The second model is simply the Bose-Hubbard model...
FIG. 15. (Color online) The orbitals $\phi_i(x)$, the particle density distribution $\rho(x, x)$, and the one-particle correlator $\rho(0, x)$ for three different values of $g$. The circles are for the permanent state, while the * markers are for the exact diagonalization results. The setting is a one-dimensional Bose-Hubbard model in a harmonic trap as defined in (58). The fixed parameters are $(N, L, \kappa) = (5, 13, 0.25)$ as in Fig. 6. In the top panels, the dotted line sketches the harmonic potential.

FIG. 16. (Color online) Same as Fig. 15, but with $M = 2$ configurations. Note that the variationally calculated density distribution $\rho(x, x)$ and the correlation function $\rho(0, x)$ are asymmetric.

with the periodic boundary condition. Because of the boundary condition, the lattice can be visualized as a closed lattice ring, and the symmetry group of the model is recognized as that of a regular polygon, i.e., the dihedral group $D_L$. The group consists of $L$ translations (or rotations) and $L$ reflections

$$D_L = \{\hat{S}^m \hat{T}^n, 0 \leq m \leq 1, 0 \leq n \leq L - 1\}. \quad (66)$$

Here the generating operators $\hat{S}$ and $\hat{T}$ are defined as $(\hat{S}\phi)(x) = \phi(-x)$ and $(\hat{T}\phi)(x) = \phi(x-1)$. By the Perron-Frobenius theorem, the exact ground state $|GS\rangle$ is non-degenerate and positive everywhere in the Fock basis. It then follows easily that $|GS\rangle$ belongs to the trivial representation of the dihedral group. The projection operator for this irreducible representation is simply

$$\hat{P} = \frac{1}{2L} \sum_{m=0}^{L-1} \sum_{n=0}^{L-1} \hat{S}^m \hat{T}^n. \quad (67)$$

Therefore, if we obtain an $M$-configuration variational state in the form of (60) by the algorithm, by projection we obtain immediately the following state invariant under all the symmetry transforms of the model,

$$\Phi_e = \sum_{m=0}^{1} \sum_{n=0}^{L-1} M_{\alpha} \hat{S}^m \hat{T}^n \phi_1^{(\alpha)}, \ldots, \hat{S}^m \hat{T}^n \phi_N^{(\alpha)}). \quad (68)$$

which is a $2LM$-configuration state.

In Fig. 23, we consider such a model with $(N, L, g) = (3, 11, 5)$. In the top row, we see that with $M = 1$ to $M = 3$, the variational state always breaks the symmetry of the model completely, i.e., all the translation and reflection symmetries are lost. This is most easily seen from the density distribution. In the bottom row, with the projected states, the symmetry of the exact ground state is recovered. We see that generally, the projection process not only restores the expected symmetry, but also lowers the energy and increases the overlap. Remarkably, it is most effective in the single configuration case.

From Fig. 22 and Fig. 23, we see that the symmetry restoration procedure is a worthwhile post-processing to the time-consuming iteration procedure. It is fairly cheap and reasonably effective.
D. When multiple configurations are perfect

In the proceeding sections, we have seen that taking multiple configurations can get us very close to the target state. Here, we discuss the scenario that a generic target state can be exactly recovered by multiple configurations.

By Proposition 4, if the single-particle Hilbert space is of dimension $L = 2$, then for arbitrary $N$, an $N$-boson state is a permanent state, i.e., it consists of a single configuration. For higher values of $L$, a generic state is not a permanent state and a natural question is, at least how many configurations we need to recover it exactly. A quick lower bound is obtained by mere dimension counting. The dimension $\mathcal{D}$ of the many-body Hilbert space is given by $\lfloor x \rfloor$. By Proposition 4, a permanent state has $d = N(L-1) + 1$ degrees of freedom. Hence, we need at least

$$M_0 = \lceil \mathcal{D}/d \rceil$$

configurations to recover a generic state, where $\lceil x \rceil$ denotes the least integer no less than $x$. There is no reason that this lower bound can be achieved. Indeed, it is an underestimate in the special case of $N = 2$. By Proposition 4 we need $\lceil (L+1)/2 \rceil$ configurations to recover a generic 2-boson state. Here the number scales as $L/2$ for large $L$. However, the estimate of (69) is $M_0 \simeq L/4$ for large $L$.

Although (69) fails for $N = 2$, there are evidences that for many pairs of $(L, N)$, it does give the right answer. Specifically, extensive numerical experiments indicate that if the pair $(L, N)$ take values among the set of $\{(3, 2), (3, 3), (3, 4), (3, 5), (4, 3)\}$, the $N$-boson wave function can always be written as the summation of $M_0 = 2$ permanent states. Similarly, if $(L, N)$ take values among the set of $\{(4, 4), (5, 3)\}$, the $N$-boson wave function can always be written as the sum of $M_0 = 3$ permanent states.

The numerical experiment is done in the following way. We first generate a random $N$-boson state, then generate a set of $M_0$ random single-particle orbitals, with $M_0$ given by (69), and then use the overlap maximization algorithm to update the orbitals. The concern is whether the overlap will surpass the threshold $1 - 10^{-5}$ after 300 rounds of update. If not, a new set of random orbitals are generated and the optimization process is restarted. This process is repeated until in some run the threshold is surpassed. If so, we turn to a new random $N$-boson state and repeat the check.

For the set of values of $(L, N)$ mentioned above, we have checked over $10^5$ random $N$-boson states, and they all passed the check. This is strong evidence that for such $(L, N)$, the naive lower bound of (69) is achieved.

So far, we have failed to find a rigorous proof of the findings above. Here we just reformulate the problem in pure mathematics so that it might be more convenient for
that it can be written as the sum of \( M \) means

\[
\prod_{j=1}^{L} (a_j^\dagger)^{n_j} |\text{vac}\rangle.
\]

Here the summation is over all occupation tuple \( n = (n_1, n_2, \ldots, n_L) \) with \( n_j \geq 0 \) and \( \sum_{j=1}^{L} n_j = N \). By (3), that it can be written as the sum of \( M \) permanent states means

\[
\Psi = \sum_{n} C_n \prod_{j=1}^{L} (a_j^\dagger)^{n_j} |\text{vac}\rangle,
\]

where \( C_{ij}^{(\alpha)} \) are constants. Because the \( a_j^\dagger \) operators commute, and the (unnormalized) Fock states \( \prod_{j=1}^{L} (a_j^\dagger)^{n_j} |\text{vac}\rangle \) are linearly independent, this is equivalent to saying that the degree-\( N \) homogeneous polynomial in \( L \) variables

\[
P(z_1, z_2, \ldots, z_L) = \sum_{n} C_n \prod_{j=1}^{L} z_j^{n_j}
\]

is expressible as

\[
P(z_1, z_2, \ldots, z_L) = \sum_{\alpha=1}^{M} \prod_{j=1}^{N} (\sum_{i=1}^{L} C_{ij}^{(\alpha)} z_j).
\]

Formulated in this way, we see the problem is very similar to the polynomial Waring problem [40]. The difference is just that while in the Waring problem, one seeks to decompose a general degree-\( N \) homogeneous polynomial in \( N \)-th powers of linear forms (linear polynomials), here we seek a decomposition in terms of \( N \)-th products of linear forms.

E. A stability problem

Having checked the accuracy, reliability, and flexibility of the algorithm, we now apply it to a realistic problem.

In atomic physics, a famous problem is the stability of the negative ion of hydrogen [41]. The question is, can we add an extra electron to a hydrogen atom, or can a proton bind two electrons simultaneously? The delicacy is that the repulsion between the two electrons is as strong as the attraction of the proton to either of them. Historically, it took people much imagination, insights, and endeavors to devise appropriate variational wave functions to establish the stability of the system [42–44]. Among all the trial wave functions people came up with, the one by Chandrasekhar is the simplest—It is a permanent state! Explicitly, it is of the form [44]

\[
f(r_1, r_2) = e^{-\alpha r_1 - \beta r_2} + e^{-\beta r_1 - \alpha r_2},
\]

where \( r_{1,2} \) are the distances of the two electrons to the proton and \( \alpha, \beta \) are two variational parameters. The energy minimum is achieved at \((\alpha, \beta) = (1.03925, 0.28309)\). Note that \( \alpha \gg \beta \), which means that one orbital is “in”, while the other is far “out”. This configuration is of course reasonable. The outer electron feels an almost completely shielded potential, and therefore can only be loosely bound.

Here we consider a lattice variant of the problem [45]. Suppose we have two on-site interacting bosons on a cubic lattice with a single defect site. The Hamiltonian is

\[
H = -\sum_{\langle i,j \rangle} (a_i^\dagger a_j + \text{h.c.}) + g \sum_i a_i^\dagger a_i^\dagger a_i a_i - U a_i^\dagger a_i^\dagger a_i^\dagger a_i.
\]

The defect site is taken as the origin of the lattice, which theoretically should be infinite but in our numerical simulation will be assumed to be a \( L_1 \times L_1 \times L_1 \) one and will be put as symmetric as possible with respect to the defect site. Apparently, the defect site is introduced to mimic the proton attraction and the on-site interaction to mimic the electron-electron repulsion.

It is well known that if the defect potential \( U \) is strong enough, i.e., if \( U \) is larger than some critical value \( U_c \) which is about 3.957, the defect can induce a single-particle defect model localized around it [46]. The two bosons can then be trapped in this defect mode simultaneously if they are not interacting. The problem is whether they can still be bound by the defect potential if the repulsion \( g \) between them is turned on, and tuned to a certain strength.

Denote the energy of the defect mode as \( E_{df} \). At \( U = U_c \), \( E_{df} = -6 \). As \( U \) increases, \( E_{df} \) decreases mono
FIG. 20. (Color online) The orbitals $\phi_i^{(a)}(x)$, the particle density distribution $\rho(x, x)$, and the one-particle correlator $\rho(0, x)$ for three increasing values of $M$. The setting and parameters are the same as in Fig. 18(c).

![Diagram showing orbitals, density distribution, and one-particle correlator for different values of M.]

FIG. 21. (Color online) Same as Fig. 20, but the value of $g$ is 15 instead of 4.

![Diagram showing orbitals, density distribution, and one-particle correlator with g=15.]

...tonically. Its exact value can be easily calculated by exact diagonalization with a sufficiently large lattice, say, $L_1 = 101$. The ionization threshold of the double-bound state is apparently

$$E_{th} = -6 + E_{df}, \quad (76)$$

which corresponds to the state in which one boson is trapped in the defect mode and the other boson free and at the bottom of the energy band.

The ground state of the two-boson system can then be shown to be a double-bound state if we can show that the ground state energy is below the threshold energy. In Fig. 23 we study a concrete case with $(g, U) = (1.5, 4.5)$. Note that for this value of $U$, the defect mode is very shallow, with the energy $E_{df} = -6.136$, so the fate of the double-bound state is uncertain without calculation. In Fig. 23(a), we see that the variational calculation with a permanent state with a single configuration yields very accurate results—the difference with those obtained by exact diagonalization is hardly visible even by the minute scale of the plot. Limited by the memory capacity of our laptop, we can do exact diagonalization only up to $L_1 = 16$, which as we see is insufficient for proving binding. However, with the variational method, we can work up to $L_1 = 21$ and succeed in pushing the ground state energy $E_{gs}$ below the threshold, thus proving binding. For comparison, we have also checked the Gross-Pitaevskii approximation, which as a special case of the permanent approximation, forces the two orbitals to be the same. We see that the error is much larger—too large to prove binding actually. In Fig. 23(b), we examine the quality of the permanent and the Gross-Pitaevskii variational states by considering their overlap with the exact ground state $|GS\rangle$ obtained by exact diagonalization. Again, we see that the permanent state is an exceedingly good variational state—its overlap with the exact state is as high as 0.9994 even for $L_1 = 16$. This vividly demonstrates the relevance of the permanent state in bosonic systems. The Gross-Pitaevskii state is also good, but not as good as the permanent state. Its errors in energy and overlap are 15-50 times larger. We thus see how crucial it is to allow the orbitals to vary independently.

As a second case, we consider $(g, U) = (-2.5, 3.5)$ in Fig. 24. This value of $U$ is inadequate for the formation of a defect mode. However, as the interaction between the two bosons is now attractive, a double-bound state is still possible as the particle-particle attraction can reinforce the defect potential. This is indeed the case. As we see in Fig. 24(a), both exact diagonalization and the permanent variational state with $M = 2$ configurations can push the ground state energy below the threshold, which is now simply $E_{th} = -12$. Unlike in Fig. 23, here the permanent approach with a single configuration fails to prove binding. It yields identical results with the Gross-Pitaevskii approach. This indicates that the permanent state actually degenerates into the Gross-Pitaevskii state,
FIG. 22. (Color online) The orbitals \(\phi_i(x)\), the particle density distribution \(\rho(x,x)\), and the one-particle correlator \(\rho(0,x)\) for \(N = 3\) bosons in a double-well potential [see Eqs. (63) and (64)]. The parameters are \(L = 13\), \(g = 2\), \(\kappa = 0.5\), \(h = 4.5\), \(\sigma = 1\). In (a) and (c), the variational state consists of \(M = 1\) or \(M = 2\) configuration(s), respectively. In (b), the state is built out of the state in (a) by the projection process in (65). In each column, the energy \(E_{gs}\) of the variational state and its overlap \(O\) with the exact ground state are shown. The exact value of the ground state energy per particle is 0.9583.

FIG. 23. (Color online) The orbitals \(\phi_i(x)\), the particle density distribution \(\rho(x,x)\), and the one-particle correlator \(\rho(0,x)\) for \(N = 3\) bosons on a closed chain with \(L = 11\) sites. The on-site interaction strength \(g = 5\). The top row corresponds to the pro-projection variational states with \(M = 1\) to \(M = 3\) configurations. The bottom row corresponds to the projected variational states. As in Fig. 22, for each state, its energy and its overlap with the exact ground state are shown. The exact value of the ground state energy per particle is \(-1.8526\).

A fact further confirmed by studying the overlap of these states with the exact state in Fig. 25(b). In both panels of Fig. 25, we see that in this particular problem and with this particular set of parameters, by including one more configuration, the error in energy and overlap can be reduced by one order of magnitude. This exemplifies the necessity and effectiveness of taking multiple configurations.

Finally, we mention that for a two-body (or even a three-body) system like the present one, the permanent calculation is never an issue. The preparation of the operators \(F\) and \(\hat{G}\) can be done swiftly, and the most time-consuming part would be solving the generalized eigenvalue problem (46) or (57) if the lattice size is in the order of thousands as here. But here are some tricks. First, as we need just the smallest generalized eigenvalue, we can invoke a Lanczos-type algorithm (in Matlab, just call \texttt{eigs}); second, actually we do not need to construct \(\hat{F}\) as a matrix explicitly. In a Lanczos-type algorithm, we just need to know its action on a vector, which is very simple.
FIG. 24. (Color online) (a) Ground state energy (per particle) of the two-boson model \( \text{(75)} \) as calculated by exact diagonalization (ED), or the permanent variational state with a single configuration, or under the Gross-Pitaevskii approximation (GPA). The dashed line indicates the ionization threshold \( \text{(76)} \) of the double-bound state. The parameters are \((g, U) = (1.5, 4.5)\). (b) Overlap of the permanent state and the Gross-Pitaevskii state with the exact ground state \( |\text{GS}\rangle \). Note that the very limited memory of our laptop (less than 8 GB) allows us to do ED on a cubic lattice only up to the size \( L_1 = 16 \).

FIG. 25. (Color online) (a) Ground state energy (per particle) of the two-boson model \( \text{(75)} \) as calculated by exact diagonalization (ED), or the permanent approach with \( M = 1 \) or \( M = 2 \) configuration(s), or under the Gross-Pitaevskii approximation (GPA). The parameters are \((g, U) = (-2.5, 3.5)\). (b) Overlap between the exact ground state \( |\text{GS}\rangle \) and the various variational states.

VI. CONCLUSIONS AND OPEN PROBLEMS

We have explored the potential of the permanent state as variational wave functions for bosons. The result is very encouraging. First, we found that for the one-dimensional Bose-Hubbard model with periodic boundary condition and at unit filling, the exact ground state can be well approximated by a permanent state with translation-related orbitals. The permanent state overlaps well with the exact ground state, yields energy close to the exact value, and produces correlation functions close to the exact ones. Then with an iteration algorithm, we examined more general models. It is quite often that a single permanent state approximates the exact ground state very well, by all the criterions of energy, overlap, and correlation functions. In case the discrepancy is apparent, it can be remedied by including more configurations.

The algorithm has its advantages and disadvantages. Let us first address its disadvantages. The primary drawback of a permanent-based approach is of course the permanent computation, which scales unfavorably with the particle number \( N \). However, with current computational facilities, it is not prohibitively expensive. On our
laptop, in the single-configuration case, it takes about 5.1 seconds to update one orbital for $N = L = 12$, and the time reduces to 1.0 second if $N = 10$. Hence, it is totally feasible to study a large enough few-boson system with the algorithm. Note that there is still room for acceleration, as computing the permanents of the minors of the overlap matrix can be easily parallelized. Note also that in practice many models of interest have only $N = 2$ or $N = 3$ particles. For such small values of $N$, the permanent computation is of course not an issue at all. While the scaling of the complexity of the algorithm with respect to the particle number $N$ is not that favorable, the scaling with respect to the system volume $L$ is quite favorable. The observation is that for $L \lesssim N$, most time is spent on the permanent calculation and the $L$-dependence is negligible. Only for $L \gg N$, the time needed to update one orbital increases apparently with $L$, but still it increases at most in a polynomial way. For example, for $N = 12$, the time is 5.5, 7.6, and 21.4 seconds for $L = 25, 50, 100$, respectively. Roughly speaking, while we are indeed confined to a limited number of particles, we have essentially no limitation on the system volume, nor the dimensionality of the system. The memory needed by the algorithm is also minimal. While in this paper we have only considered the $(N, L)$ pairs for which the dimension of the many-body Hilbert space is at most on the order of one million, so that exact diagonalization is possible and we have exact results for comparison, the algorithm can handle other values of $(N, L)$ easily.

It might be a good idea to combine the current algorithm with the Lanczos algorithm. That is, one can think of post-processing the single- or multi-configurational variational state $\Phi$ obtained by the current algorithm by a Lanczos-type process. Starting from $\Phi$, we apply the Hamiltonian $H$ repeatedly to it, and at each stage, we try to approximate the resultant wave function with a combination of permanent states. In this way, we construct a Krylov-type subspace spanned by a set of multi-configurational states. A variational wave function better than $\Phi$ can then be sought in this subspace by solving a generalized eigenvalue problem. The procedures can be carried out most conveniently for the two-particle ($N = 2$) case, thanks to the Autonne-Takagi factorization (22). Some preliminary test yields encouraging results.

In this tentative work, we have focused on some one-dimensional lattice models in the Bose-Hubbard category. Generalization to continuum models (17, 18), higher dimensions, and multi-component systems should be straightforward. Moreover, the simple strategy of the algorithm apparently is also applicable to fermions—Actually, originally it was used for fermions (24, 25), although for some different problem. It should be worthwhile to check how it works for fermions.

Below are some open problems. The fact that for the one-dimensional Bose-Hubbard model with periodic boundary condition and at unit filling, the exact ground state can be well approximated by a permanent state with translation-related orbitals is very impressive. This should arouse one’s interest in the permanent state in its own right. In the field of cold atoms, the folklore is that Bose-Einstein condensation occurs as the temperature lowers, de Broglie wave lengths of the particles increase, and the wave-packets overlap. Now, a permanent state with translation-related orbitals is in accord with this picture. Given a primitive wave packet $\phi(\vec{x})$, we can imagine a many-body permanent state constructed with the wave packets $\phi(\vec{x} - \vec{R}_m)$, where $\vec{R}_m$ runs through an $n$-dimensional lattice. The overlap between adjacent orbitals can be well adjusted by changing the length scale of the primitive orbital $\phi$. The concern is, is it possible to realize a transition by tuning its length scale? How does the correlation function depend on the primitive orbital? What is the effect of the dimensionality of the lattice? In short, can we use a permanent state with regularly distributed identical wave-packets as a prototypical wave function to model the condensation transition? Note that the problem does not refer to a Hamiltonian.

That a single permanent state is often a very good approximation of the exact ground state motivates two problems. First, is it possible to construct an interacting Hamiltonian (49) whose ground state is exactly a permanent state? Second, how dense are the permanent states in the many-boson Hilbert space? Quantitatively, is there a number $\delta > 0$, such that for any many-boson state there exists a permanent state whose overlap with it is at least $\delta$? Preliminary study suggests that $\delta \geq 0.15$ for $(N, L) = (3, 13)$. This is not a small number in view of the fact that the dimension of the few-body Hilbert space is 455. The ground state of a realistic model is non-generic, so the largest possible value of the overlap of a permanent state with it should be much higher.

In this paper, we have focused on the energy-minimization problem. The other problem of overlap-maximization, i.e., the problem of finding the optimal single- or multi-configurational permanent approximation of a given bosonic wave function should also be a worthy one. It is about the structure of a bosonic wave function. At least for fermions, it is now well-known that the anti-symmetry condition entails deep structures of the fermionic wave function, with consequences far beyond the commonplace of the Pauli exclusion principle (50, 54), and the notion of optimal Slater approximation has proven to be useful in this study (21, 25, 52). It is fair to expect that for bosons, the symmetry condition also has far-reaching consequences and hopefully, the notion of optimal permanent approximation is a useful one too.

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Appendix: Expressions of \( \hat{F} \) and \( \hat{G} \) in Sec. [IV]

Let us start with \( \hat{G} \) which is simpler than \( \hat{F} \). By an analogy of the Laplace expansion for the determinant, we have

\[
\langle \Phi | \Phi \rangle = \text{per}(A) = \langle \phi_1 | \phi_1 \rangle \text{per}(A; 1|1) + \sum_{j_1=2}^{N} \langle \phi_1 | \phi_{j_1} \rangle \text{per}(A; 1|j_1)
\]

\[
= \langle \phi_1 | \phi_1 \rangle \text{per}(A; 1|1) + \sum_{j_1=2}^{N} \sum_{i_1=2}^{N} \langle \phi_1 | \phi_{j_1} \rangle \langle \phi_{i_1} | \phi_1 \rangle \text{per}(A; 1, i_1|1, j_1).
\]

(A.1)

From this expression, we can read off the operator \( \hat{G} \) defined by \( \langle \phi_1 | \hat{G} | \phi_1 \rangle = \langle \Phi | \Phi \rangle \). It is

\[
\hat{G} = \text{per}(A; 1|1) \hat{I} + \sum_{j_1=2}^{N} \sum_{i_1=2}^{N} \text{per}(A; 1, i_1|1, j_1) \langle \phi_{i_1} | \phi_1 \rangle,
\]

(A.2)

where \( \hat{I} \) is the identity operator. Note that \( \hat{G} \) depends on the orbitals \( \phi_{2 \leq j \leq N} \) but not on \( \phi_1 \).

We then turn to \( \hat{F} \). For the single-particle part, we have

\[
\langle \Phi | H_1 | \Phi \rangle = \sum_{i_1=1}^{N} \sum_{j_1=1}^{N} \langle \phi_{i_1} | K | \phi_{j_1} \rangle \text{per}(A; i_1|j_1)
\]

\[
= \langle \phi_1 | K | \phi_1 \rangle \text{per}(A; 1|1) + \sum_{j_1=2}^{N} \langle \phi_1 | K | \phi_{j_1} \rangle \text{per}(A; 1|j_1) + \sum_{i_1=2}^{N} \langle \phi_1 | K | \phi_1 \rangle \text{per}(A; i_1|1)
\]

\[
+ \sum_{i_1=2}^{N} \sum_{j_1=2}^{N} \langle \phi_1 | K | \phi_{j_1} \rangle \text{per}(A; i_1|j_1)
\]

\[
= \langle \phi_1 | K | \phi_1 \rangle \text{per}(A; 1|1) + \sum_{j_1=2}^{N} \sum_{i_1=2}^{N} \langle \phi_1 | K | \phi_{j_1} \rangle \langle \phi_{i_1} | \phi_1 \rangle \text{per}(A; 1, i_1|1, j_1)
\]

\[
+ \sum_{j_1=2}^{N} \sum_{i_1=2}^{N} \langle \phi_1 | K | \phi_{j_1} \rangle \langle \phi_{i_1} | \phi_{j_1} \rangle \langle \phi_{i_2} | \phi_1 \rangle \text{per}(A; 1, i_1, i_2|1, j_1, j_2).
\]

(A.3)

Here in the last line, the summation \( \sum_{i_1 \neq i_2 \neq j_1 \neq j_2}^{N} \) means that \( i_1 \) and \( i_2 \) both run from 2 to \( N \), but they must take different values. Similar summation expressions below should be interpreted similarly. We see that the contribution of \( H_1 \) to \( \hat{F} \) is

\[
\text{per}(A; 1|1)K + \sum_{j_1=2}^{N} \sum_{i_1=2}^{N} K | \phi_{j_1} \rangle \langle \phi_{i_1} | \ast \text{per}(A; 1, i_1|1, j_1) + \sum_{j_1=2}^{N} \sum_{i_1=2}^{N} | \phi_{j_1} \rangle \langle \phi_1 | K \ast \text{per}(A; 1, i_1|1, j_1)
\]

\[
+ \sum_{i_1=2}^{N} \sum_{j_1=2}^{N} \langle \phi_1 | K | \phi_{j_1} \rangle \hat{I} \ast \text{per}(A; 1, i_1|1, j_1) + \sum_{i_1 \neq i_2 \neq j_1 \neq j_2}^{N} \sum_{j_1=2}^{N} \sum_{i_1=2}^{N} \langle \phi_{i_1} | K | \phi_{j_1} \rangle \langle \phi_{i_2} | \phi_{j_2} \rangle \ast \text{per}(A; 1, i_1, i_2|1, j_1, j_2).
\]

(A.4)

Next we turn to the two-particle or the interaction term \( H_2 \). We have by \( (34) \)

\[
\langle \Phi | H_2 | \Phi \rangle = \frac{1}{2} \sum_{i_1 \neq i_2, j_1 \neq j_2}^{N} \sum_{j_1=2}^{N} \langle \phi_1, \phi_{i_2} | U | \phi_{j_1}, \phi_{j_2} \rangle \text{per}(A; i_1, i_2|j_1, j_2).
\]

(A.5)
For clarity, three cases will be considered separately. In the first case, two $\phi_1$’s are associated with $U$. We have

$$\frac{1}{2} \sum_{i_1=2}^{N} \sum_{j_1=2}^{N} \langle \phi_1 \phi_1 | U | \phi_1 \phi_2 \rangle \text{ per}(A; 1, i_1, j_1, j_2) + \frac{1}{2} \sum_{i_1=2}^{N} \sum_{j_1=2}^{N} \langle \phi_1 \phi_1 | U | \phi_1 \phi_1 \rangle \text{ per}(A; 1, i_1, j_1, j_1)$$

$$+ \frac{1}{2} \sum_{i_1=2}^{N} \sum_{j_1=2}^{N} \langle \phi_1 \phi_1 | U | \phi_1 \phi_1 \rangle \text{ per}(A; 1, i_1, j_1, j_1) + \frac{1}{2} \sum_{i_1=2}^{N} \sum_{j_1=2}^{N} \langle \phi_1 \phi_1 | U | \phi_1 \phi_1 \rangle \text{ per}(A; 1, i_1, j_1, j_2)$$

$$= \sum_{i_2=2}^{N} \sum_{j_2=2}^{N} \langle \phi_1 \phi_1 | U | \phi_1 \phi_2 \rangle \text{ per}(A; 1, i_2, j_2, j_2) + \sum_{i_1=2}^{N} \sum_{j_1=2}^{N} \langle \phi_1 \phi_1 | U | \phi_1 \phi_2 \rangle \text{ per}(A; 1, i_1, j_1, j_2)$$

$$= \sum_{i_2=2}^{N} \sum_{j_2=2}^{N} \langle \phi_1 \phi_1 | U | \phi_1 \phi_2 \rangle \text{ per}(A; 1, i_2, j_2, j_2) + \sum_{i_1=2}^{N} \sum_{j_1=2}^{N} \langle \phi_1 \phi_1 | U | \phi_1 \phi_2 \rangle \text{ per}(A; 1, i_1, j_1, j_2). \quad (A.6)$$

Contributions of these expressions to the operator $\hat{F}$ can be easily read off. For example, the operator $O$ corresponding to the matrix element $\langle \phi_1 \phi_1 | U | \phi_1 \phi_2 \rangle$ is defined by $\langle \phi_1 | O | \phi_1 \rangle = \langle \phi_1 \phi_1 | U | \phi_1 \phi_2 \rangle$. For the Bose-Hubbard model in which $U$ is an on-site interaction [57], the operator $O$ is actually a general complex potential with the explicit expression $O(x) = g \psi_1(x) \psi_1(x)$.

In the second case, one $\phi_1$ is associated with $U$. We have

$$\frac{1}{2} \sum_{i_1=2}^{N} \sum_{j_1=2}^{N} \langle \phi_1 \phi_1 | U | \phi_1 \phi_2 \rangle \text{ per}(A; 1, i_1, j_1, j_2) + \frac{1}{2} \sum_{i_1=2}^{N} \sum_{j_1=2}^{N} \langle \phi_1 \phi_1 | U | \phi_1 \phi_2 \rangle \text{ per}(A; 1, i_1, j_1, j_1)$$

$$+ \frac{1}{2} \sum_{i_1=2}^{N} \sum_{j_1=2}^{N} \langle \phi_1 \phi_1 | U | \phi_1 \phi_2 \rangle \text{ per}(A; 1, i_1, j_1, j_1) + \frac{1}{2} \sum_{i_1=2}^{N} \sum_{j_1=2}^{N} \langle \phi_1 \phi_1 | U | \phi_1 \phi_2 \rangle \text{ per}(A; 1, i_1, j_1, j_1)$$

$$= \sum_{i_2=2}^{N} \sum_{j_2=2}^{N} \langle \phi_1 \phi_1 | U | \phi_1 \phi_2 \rangle \text{ per}(A; 1, i_2, j_2, j_2) + \sum_{i_1=2}^{N} \sum_{j_1=2}^{N} \langle \phi_1 \phi_1 | U | \phi_1 \phi_2 \rangle \text{ per}(A; 1, i_1, j_1, j_1)$$

$$= \sum_{i_2=2}^{N} \sum_{j_2=2}^{N} \langle \phi_1 \phi_1 | U | \phi_1 \phi_2 \rangle \text{ per}(A; 1, i_2, j_2, j_2) + \sum_{i_1=2}^{N} \sum_{j_1=2}^{N} \langle \phi_1 \phi_1 | U | \phi_1 \phi_2 \rangle \text{ per}(A; 1, i_1, j_1, j_1). \quad (A.7)$$

In the third case, none $\phi_1$ is associated with $U$. We have

$$\frac{1}{2} \sum_{i_1=2}^{N} \sum_{j_1=2}^{N} \langle \phi_1 \phi_1 | U | \phi_1 \phi_2 \rangle \text{ per}(A; 1, i_2, j_2, j_2)$$

$$= \frac{1}{2} \sum_{i_1=2}^{N} \sum_{j_1=2}^{N} \langle \phi_1 \phi_1 | U | \phi_1 \phi_2 \rangle \text{ per}(A; 1, i_1, j_1, j_2)$$

$$+ \frac{1}{2} \sum_{i_1=2}^{N} \sum_{j_1=2}^{N} \langle \phi_1 \phi_1 | U | \phi_1 \phi_2 \rangle \text{ per}(A; 1, i_1, j_1, j_2). \quad (A.8)$$

In the last line, we see that we have to calculate the permanents of a series of $(N-4) \times (N-4)$ matrices $(N-3)^2$ times. By the improved Ryscer algorithm, the total evaluation is on the order of $N^7 2^{(N-5)}$. This is the most time-consuming part in the preparation of $\hat{F}$ and $\hat{G}$, if $N \geq 5$.

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[1] D. R. Hartree, Math. Proc. Camb. Philos. Soc. 24, 89 (1928).

[2] J. C. Slater, Phys. Rev. 32, 339 (1928).
While it is known that the Slater determinant states for $m$

is conjectured that the set of permanent states is not a

Therefore not interested in such trivial cases.

Note that by Proposition 3, any state of a two-site Bose-

some MATLAB subroutines for permanent computation

they lead to much poorer approximations, so we do not

we do not bother to mention them.

they are not interested in such trivial cases.

which is conjectured that the set of permanent states is not a

they are orthonormalized at the beginning.

results in the fermionic case, the orbitals $\phi_{1 \leq i \leq N}$ do not need to be orthonormal neither. However, the result-

In this case, the resulting Slater state would be essentially the same if the orbitals are orthonormalized at the beginning.

Some MATLAB subroutines for permanent computation using the Ryser algorithm and taking advantage of the Gray code are in the Supplementary Material.

We have actually also tried Gaussian functions. However, they lead to much poorer approximations, so we do not bother to mention them.

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