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Fermion Monte Carlo without fixed nodes: A game of life, death, and annihilation in Slater determinant space

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We have developed a new quantum Monte Carlo method for the simulation of correlated many-electron systems in full configuration-interaction (Slater determinant) spaces. The new method is a population dynamics of a set of walkers, and is designed to simulate the underlying imaginary-time Schrödinger equation of the interacting Hamiltonian. The walkers (which carry a positive or negative sign) inhabit Slater determinant space, and evolve according to a simple set of rules which include spawning, death and annihilation processes. We show that this method is capable of converging onto the full configuration-interaction (FCI) energy and wave function of the problem, without any a priori information regarding the nodal structure of the wave function being provided. Walker annihilation is shown to play a key role. The pattern of walker growth exhibits a characteristic plateau once a critical (system-dependent) number of walkers has been reached. At this point, the correlation energy can be measured using two independent methods—a projection formula and a energy shift; agreement between these provides a strong measure of confidence in the accuracy of the computed correlation energies. We have verified the method by performing calculations on systems for which FCI calculations already exist. In addition, we report on a number of new systems, including CO, O₂, CH₄, and NaH—with FCI spaces ranging from 10⁹ to 10¹⁴, whose FCI energies we compute using modest computational resources. © 2009 American Institute of Physics. [DOI: 10.1063/1.3193710]

I. INTRODUCTION

It has long been known that the only major obstacle preventing the exact numerical simulation of many-electron systems via stochastic methods such as diffusion quantum Monte Carlo (DMC) or the related Green's function Monte Carlo² (GFMC) is the Fermion sign problem.³ This problem stems from the antisymmetry property of many-electron wave functions to electron exchange, which leads to wave functions which have both positive and negative amplitudes. Since the Schrödinger equation can be viewed as a diffusion equation in imaginary time, its lowest energy solution is, in general, nodeless and symmetric, and therefore does not satisfy the required Fermion antisymmetry. Stochastic propagation leads exponentially quickly to this undesired solution. One way to prevent this "Boson catastrophe" is to constrain the propagation to disjoint areas of similar sign using the fixed-node approximation, 4-7 a procedure which would be exact if the applied nodal boundaries coincided with the exact nodal hypersurface of the ground-state electronic wave function. However, in practice, this is not the case, and it has proven extremely difficult to improve the fixed-node surface toward the exact one. The Fermion sign problem is thus recognized as one of the most important unsolved problems of computational theoretical physics and chemistry. A key question is whether the exact nodal hypersurface can emerge during the course of a simulation. Such a simulation would therefore not require any a priori information regarding the nodes of the exact wave function. In this paper we describe a new quantum Monte Carlo (QMC) method in which this highly desirable property is shown to arise, for systems described with basis sets commonly used in quantum chemistry. Our method is shown to converge on to the full configuration-interaction (FCI) solution, i.e., the exact wave function and energy for the basis set under consideration.^{8–11} In addition to reproducing existing FCI calculations, we have used this method to predict the FCI energies of several molecules which have not been reported to date. We also report a benchmark study of the Ne atom (correlating all electrons) in several basis sets up to cc-pCVQZ, which has a FCI basis exceeding 10¹⁴ determinants. That is over 10 000 times larger than the largest FCI calculation reported to date. ¹² Our method unifies QMC and FCI in a way which profoundly extends the scope of both techniques.

There are three ingredients to the new method, which take elements from DMC and FCI: (i) In common with DMC, we perform a long-time integration of the imaginary-time Schrödinger equation; however, in contrast to DMC, this is achieved in a space of Slater determinants. In addition, the propagation step in our algorithm differs from DMC in that it consists purely of population dynamics (i.e., walker birth and death processes). There are no diffusive moves as such. (ii) In common with DMC, the instantaneous wave function is represented using "walkers," rather than ampli-

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tude coefficients, the latter being the case in FCI. The representation using walkers enables us to describe stochastically the FCI wave function without storing all amplitudes simultaneously. (iii) Each walker carries a positive or negative sign. The key ingredient in our algorithm is walker annihilation, in which pairs of walkers of opposite sign which coincide on the same determinant are removed from the simulation. We show that an algorithm based on these three ideas converges on the exact solution to the Hamiltonian expressed in the basis, which owing to the Fermion antisymmetry of the Slater determinant basis, will be the lowest Fermionic solution available in this basis. It should be noted that working in Slater determinant space does not circumvent the sign problem, even though it prevents convergence to a Bosonic solution. This is because the FCI wave function does not (in general) have strictly non-negative amplitudes in this space. We note that walker annihilation has been previously suggested in the context of nodal release GFMC and DMC. 13-16 However, as noted in these studies, only low-dimensional phase spaces could be treated owing to the formidable difficulties in achieving cancellation in continuum spaces. Instead, by working in discrete Slater determinant spaces, we find that walker annihilation proves an essential and effective component of a Fermion Monte Carlo method.

There have been several proposals to simulate Fermionic systems in determinantal spaces using auxiliary-field Monte Carlo (AFMC). 17-22 However, in their exact formulation, auxiliary-field methods suffer from exponentially large statistical noise in the limit of large imaginary-time propagation, which is necessary to project out the ground state. In order to stabilize this problem, a phaseless-approximation AFMC method has been proposed, 21 with promising energies (i.e., to within a few millihartrees of known FCI energies) when applied to molecular systems. 22 However, like the fixed-node approximation, the phaseless approximation is an uncontrolled approximation which may be difficult to improve upon.

Below we give motivation and elaborate on our algorithm, before showing applications of the method to several real physical systems.

II. MOTIVATION AND DERIVATION OF THE ALGORITHM

In the FCI method, we seek a wave function Ψ_0 which satisfies the time-independent Schrödinger equation, $H\Psi_0$ = $E_0\Psi_0$, as a variationally optimized linear combination of Slater determinants $\{|D_{\bf i}\rangle\}$. These are antisymmetric functions in which N orbitals are chosen out of 2M spin orbitals $\{\phi_1,\phi_2,\ldots,\phi_{2M}\}$ and constructed as follows:

$$|D_{\mathbf{i}}\rangle \equiv |D_{n_1,n_2,\dots,n_N}\rangle = a_{n_1}^{\dagger} a_{n_2}^{\dagger},\dots,a_{n_N}^{\dagger}|\rangle \tag{1}$$

$$= \frac{1}{\sqrt{N!}} |\phi_{n_1} \phi_{n_2}, \dots, \phi_{n_N}|, \quad n_1 < n_2 < \dots < n_N. \quad (2)$$

 a_i^{\dagger} is a Fermion creation operator for spin-orbital ϕ_i . In this work, the orbitals used are real, canonical Hartree–Fock orbitals. The size of the Slater determinant space $(N_{\rm FCI})$ is on the order of $\binom{M}{N/2}^2$ for a spin-unpolarized system, a number

that grows factorially with M and N (although symmetry restrictions for some systems can reduce this by up to an order of magnitude). The FCI wave function is expressed as

$$\Psi_0^{\text{FCI}} = \sum_{\mathbf{i}} C_{\mathbf{i}} | D_{\mathbf{i}} \rangle, \tag{3}$$

where the CI coefficients $\{C_i\}$ satisfy an eigenvector problem:

$$\sum_{\mathbf{i}} \langle D_{\mathbf{i}} | H | D_{\mathbf{j}} \rangle C_{\mathbf{j}} = E_0^{\text{FCI}} C_{\mathbf{i}}.$$
 (4)

 $E_0^{\rm FCI}$ is the lowest energy solution available in this basis, an upper bound to the exact energy. Owing to the fact that the off-diagonal Hamiltonian matrix elements are not all of the same sign, the CI coefficients can be positive or negative. The "sign" structure of the FCI wave function is given by a vector whose components are $sign(C_i)$ or 0 if $C_i=0$. A trial wave function has the correct sign structure only if the sign of every component matches those of this vector (up to an overall sign, since $-\Psi_0^{FCI}$ is an equally valid solution with the same energy). In practice, it turns out to be impossible to predict $sign(C_i)$ without a knowledge of C_i , and this, in essence, is the manifestation of the Fermion sign problem in the discrete Slater determinant space. In FCI, the CI coefficients are obtained via a (nonstochastic) iterative diagonalization method. While it is recognized that the FCI method is the most robust method to treat electron correlation, its scope is greatly limited by the prohibitive computational requirements (especially storage) of such iterative diagonalization in the full space of Slater determinants. For example the Davidson²³ method or the preconditioned conjugate-gradient method²⁴ requires at least two vectors of length equal to the FCI space to be stored, and often many more. The largest molecular FCI calculation to date is the N₂ molecule, ¹² and has $\approx 10^{10}$ determinants in D_{2h} symmetry. As Nicholas Handy has observed, 25 "unless something unexpected happens, it is unlikely that size will be much exceeded." Inspired by the DMC method, rather than attempt a direct diagonalization of the FCI Hamiltonian, we propose to simulate the imaginary-time Schrödinger equation by performing a stochastic population dynamics on an evolving set of walkers which live and propagate in Slater determinant space. However, we do not impose any prior knowledge on the signs of the Slater determinants.

The imaginary-time Schrödinger equation provides the fundamental starting point of all "projector" techniques. It states that the imaginary-time derivative of any wave function Ψ is given simply by the Hamiltonian acting on $-\Psi$, namely,

$$\frac{\partial \Psi}{\partial \tau} = -H\Psi. \tag{5}$$

Given a starting wave function $\Psi(\tau=0)$, the wave function $\Psi(\tau)$ for arbitrary τ is proportional to

$$\Psi(\tau) \propto e^{-\tau H} \Psi(\tau = 0). \tag{6}$$

In order to project onto the Fermionic ground state, $\Psi(\tau=0)$ must chosen to be a fully antisymmetric function with an overlap with the ground state. Usually a single de-

terminant function with the correct symmetry, such as the Hartree–Fock determinant D_0 , suffices. The long-time limit of Eq. (6) then projects out the components of D_0 on excited states, leaving the Fermionic ground state, Ψ_0 :

$$\Psi_0 = \lim_{\tau \to \infty} e^{-\tau (H - E_0)} D_0, \tag{7}$$

where the constant of proportionality $e^{+\tau E_0}$ has been introduced to keep the ground-state contribution from decaying to zero. The aspiration of all Monte Carlo projector methods is to realize this long-time limit by performing a stochastic integration of Eq. (5). In normal DMC, however, the integration is not performed on an antisymmetrized space, and therefore admits solutions with Bosonic character, which rapidly grow to dominate unless constrained in some manner, e.g., by the fixed-node approximation. An integration in a pure antisymmetric subspace would not suffer from this effect, and this is may be a help in reducing the sign problem for Fermion systems. Our first aim, therefore, is to develop an analog to Eq. (5) in a Slater determinant basis, in a form which can be integrated stochastically.

To this end, let us define a matrix K, whose elements are the matrix elements between Slater determinants of the Hamiltonian, with the Hartree–Fock energy $E_{\rm HF}$ subtracted from the diagonal elements:

$$K_{ii} \equiv \langle D_i | K | D_i \rangle = \langle D_i | H | D_i \rangle - E_{HF} \delta_{ii}. \tag{8}$$

The diagonal matrix elements of K are therefore all positive (or zero), while the off-diagonal elements are simply the off-diagonal matrix elements of the Hamiltonian. With this definition, the lowest energy eigenvalue of K is $E_0^{\rm FCI}-E_{\rm HF}$, which is the correlation energy, $E_{\rm corr}$ for the problem. The matrix elements of H can be computed in terms of the one-electron and two-electron integrals using standard methods, which are reviewed in Appendix A.

Writing

$$\Psi(\tau) = \sum_{i} C_{i}(\tau) |D_{i}\rangle \tag{9}$$

and substituting into Eq. (5), we obtain a set of coupled linear first-order differential equations for the CI coefficients in terms of the K matrix:

$$-\frac{dC_{\mathbf{i}}}{d\tau} = \sum_{\mathbf{j}} (K_{\mathbf{i}\mathbf{j}} - S\delta_{\mathbf{i}\mathbf{j}})C_{\mathbf{j}}$$
 (10)

$$=(K_{ii}-S)C_{i}+\sum_{\mathbf{j}\neq i}K_{i\mathbf{j}}C_{\mathbf{j}},$$
(11)

where an arbitrary "energy shift," S, has been introduced into the diagonal terms, whose role will be population control, to be discussed later. It is evident that if we instantaneously have a vector \mathbf{C} of amplitudes whose components C_i satisfy

$$\sum_{\mathbf{j}} K_{\mathbf{i}\mathbf{j}} C_{\mathbf{j}} = SC_{\mathbf{i}},\tag{12}$$

then $d\mathbf{C}/d\tau=0$, making this vector stationary. In addition, by virtue of Eq. (12), this vector is an eigenstate of the K matrix (and hence of the H), with eigenvalue S. It follows that if S equals the correlation energy $E_{\rm corr}$, then the stationary state is

the ground state of the H matrix. Furthermore, if we start from an arbitrary set of C_i amplitudes [which does not satisfy Eq. (12)] and integrate the above set of coupled differential equations, the long-time solution leads to the ground-state eigenvector. This can be proven by decomposing the starting vector into the eigenstates of H, and noting that the components on the excited states would decay exponentially with time, leaving the ground-state eigenvector.

A direct numerical integration of Eq. (11) requires the full set of C_i coefficients to be available at each time step, which is prohibitive. Instead, in the spirit of DMC, let us consider a population of N_w walkers; each walker α is located on a determinant \mathbf{i}_{α} , and has a sign $s_{\alpha} = \pm 1$. We now define the C_i amplitude on determinant $|D_i\rangle$ to be proportional to the *signed* sum of walkers (N_i) ,

$$C_{\mathbf{i}} \propto N_{\mathbf{i}} = \sum_{\alpha} s_{\alpha} \delta_{\mathbf{i}, \mathbf{i}_{\alpha}} \tag{13}$$

 $(\delta_{\mathbf{i},\mathbf{i}_{\alpha}})$ is the discrete Kronecker delta, and equals one if $\mathbf{i}_{\alpha} = \mathbf{i}$, and is otherwise zero). According to Eq. (13), $N_{\mathbf{i}}$ can be positive or negative. However, the total number of walkers N_{w} is defined to be the sum over the absolute values of the $N_{\mathbf{i}}$:

$$N_w = \sum_{\mathbf{i}} |N_{\mathbf{i}}| \tag{14}$$

and is always positive.

III. A POPULATION DYNAMICS ALGORITHM

We introduce a population dynamics algorithm which simulates the set of coupled differential equations in Eq. (11). The algorithm consists of three steps performed at each time step whose length is $\delta\tau$:

(i) The spawning step: For each walker α (located on $D_{\mathbf{i}_{\alpha}}$), we select a (coupled) determinant $D_{\mathbf{j}}$ with normalized probability $p_{\text{gen}}(\mathbf{j}|\mathbf{i}_{\alpha})$ and attempt to spawn a child there with probability

$$p_s(\mathbf{j}|\mathbf{i}_{\alpha}) = \frac{\delta \tau |K_{\mathbf{i}_{\alpha}\mathbf{j}}|}{p_{\text{gen}}(\mathbf{j}|\mathbf{i}_{\alpha})}.$$
 (15)

If a spawning event is successful (i.e., if p_s exceeds a uniformly chosen random number between 0 and 1), then the sign of the child is determined by the sign of $K_{\mathbf{i}_{\alpha}\mathbf{j}}$ and the sign of the parent: It is the same sign as the parent if $K_{\mathbf{i}_{\alpha}\mathbf{j}} < 0$, and opposite to the parent otherwise. Our method to compute the generation probabilities p_{gen} is given in Appendix B. If $p_s > 1$, then multiple copies of walkers are spawned on \mathbf{j} (namely with probability $1, [p_s]$ walkers are spawned, and with probability $p_s - [p_s]$ an additional walker is spawned). In general, the number of newly spawned walkers (N_s) is much smaller than number of parent walkers (N_w) , since the time step $\delta \tau$ is such that the probability to spawn is quite low. Typically we find $N_s \sim 10^{-4}N_w$.

(ii) The diagonal death/cloning step: For each (parent) walker compute

$$p_d(\mathbf{i}_{\alpha}) = \delta \tau(K_{\mathbf{i}_{\alpha}\mathbf{i}_{\alpha}} - S). \tag{16}$$

If $p_d > 0$, the walker dies with probability p_d , and if $p_d < 0$ the walker is cloned with probability $|p_d|$. The death event happens immediately, and such a parent does not participate in the following (annihilation) step to be described shortly. Cloning events are quite rare, and only occur for S > 0, and even then only on determinants for which $\langle D_i | K | D_i \rangle < S$. In simulations where we desire to grow the number of walkers rapidly, a positive value of S is adopted, and this can lead to cloning events. However, more often, the value of S is negative (as it tries to match the correlation energy), and in such cases there can be no cloning events at all.

Both the spawning step and the diagonal death step can be done without reference to other walkers. Therefore, these two steps are "embarrassingly parallel," and can be performed without communication overhead on a parallel machine.

(iii) The annihilation step: In this (final) part of the algorithm, we run over all (newly spawned, cloned, and surviving parent) walkers, and annihilate pairs of walkers of opposite sign which are found to be on the same determinant. Each time an annihilation event occurs, the corresponding pair is removed from the list of walkers, and the total number of walkers N_w reduced by two. By keeping sorted lists of walkers, it is possible to do the search for possible annihilation events using binary searches, with the result that the annihilation step can be done with $\mathcal{O}[N_s \ln(N_s N_w)]$

At the end of the annihilation step, the lists of surviving newly spawned walkers and parents are merged. The merged list remains sorted, and becomes the main list of walkers for the next time step.²⁸ The annihilation can achieved in a memory efficient manner, keeping only one copy of the main list of walkers. In a one-electron basis with 2M orbitals, ²⁹ each walker needs [2M/32]+1 4-byte integers of storage³⁰ to encode the occupation number information of the determinant it lives on, as well as the sign of the walker. For a simulation with N_w walkers, therefore, the main walker list requires $4N_w([2M/32]+1)$ bytes of RAM. As will be seen in the applications, the number of walkers required to achieve convergence is typically smaller than the size of the FCI space, often by a significant factor. Therefore the memory requirements of this algorithm are less severe than that of a conventional FCI calculation.

It should be noted that with the above algorithm all (symmetry-allowed) determinants in the FCI space are accessible, since no restrictions are placed on the spawning step as to which determinants can be generated: Given a walker on some determinant, any determinant connected to it can be chosen to be spawned upon. It follows that the entire space of (symmetry-related) determinants can eventually be reached starting from any one determinant. The fact that the entire space is accessible enables the algorithm to converge onto the FCI wave function, which, in general, has nonzero amplitudes on all such determinants. Of course, it is also a straightforward matter to impose a truncation in the CI space, for example, by excitation level, by simply not accepting any spawning events at determinants beyond the specified truncation level. In this way, one can perform calculations equivalent to truncated CI, such as CISDTQ, etc. In large calculations, this approach can be used to prepare equilibrated or near-equilibrated ensembles of walkers at truncated excitation levels, before attempting a simulation in the full CI space.

A second point to note is that in the current algorithm each walker attempts to spawn only once per time step. It is possible to construct a modified algorithm in which each walker at each time step systematically attempts to spawn at all connected determinants (but with a different spawning probability, given by $p_s(\mathbf{j}|\mathbf{i}_{\alpha}) = \delta \tau |K_{\mathbf{i}_{\alpha}\mathbf{j}}|$). Obviously, with this modification, the computational cost per time step is much greater, although it might be expected that a faster convergence is achieved. In tests, however, this version of the spawning algorithm proved to be less efficient overall than the one we outlined above.

A further point to note concerns time-step error. The above algorithm results in changes in populations on each determinant proportional to $\delta \tau$. The underlying propagator is $G=1-\delta\tau(H-S)$. As has been pointed out in Ref. 29, in discrete spaces (with a finite spread of eigenvalues), repeated application of such propagators starting from any initial state converges onto the exact ground state as long as δau $\leq 2/(E_{\text{max}}-S)$, where E_{max} is the largest eigenvalue of H and $S \approx E_0$. Longer time steps lead to propagators in which excited-state contributions do not decay with time. In a nonstochastic implementation of this propagator, convergence to the ground-state is therefore guaranteed as long as the time step does not exceed this upper bound value. It is possible to estimate E_{max} as the energy of the most highly excited determinant available in a given one-electron basis, with E_{max} increasing with the size of the basis.

In practice, for the present stochastic algorithm, we find that there is an additional constraint on the time step which makes it to be smaller than the above value, namely that is it desirable that p_s should not greatly exceed unity, since this results in large walker blooms (i.e., multiple copies of walkers being suddenly spawned on a determinant), which can lead to inefficient sampling. On the other hand, a small time step leads to slow evolution and requires proportionately longer simulations, which is also inefficient. In practice, for the systems under consideration in this study, we find that that values for $\delta \tau$ in the range $10^{-4}-10^{-3}$ a.u. (with the smaller values being used for larger basis sets), provides a reasonable balance between these two sources of inefficiency, and that we have explicitly verified that the correlation energies we compute are insensitive of the value of the time step as long as they are in this range. In the remainder of this section we discuss some additional features of our simulation methodology.

A. Constant S and constant N_w simulations

If the shift S is kept constant at a value above E_{corr} , then, in general, the number of walkers increases exponentially. It is therefore desirable to be able to do "constant" N_w simulation. This can be achieved by periodically (every A steps) adjusting the shift S according to the following simple prescription

$$S(\tau) = S(\tau - A \delta \tau) - \frac{\zeta}{A \delta \tau} \ln \frac{N_w(\tau)}{N_w(\tau - A \delta \tau)},$$
(17)

where $N_w(\tau)$ is the total number of walkers on time step τ , and ζ is a damping parameter, i.e., if N_w has grown over the current update cycle, the shift is reduced (made more negative), while if N_w has decreased, the shift is increased (made more positive). Thus in "constant" N_w mode, the shift S varies to attempt to keep walker number constant. The damping coefficient ζ , prevents undesirable large fluctuations in S. In constant S mode the number of walkers varies, typically increasing if $S > E_{corr}$. If the distribution of walkers is correct (i.e., distributed according to the ground-state amplitude), then the value of S which maintains the number of walkers is equal to the correlation energy of the problem. Conversely, in constant N_w mode, the shift will fluctuate about the correct correlation energy when the distribution of walkers is correct. In the examples we report below, we chose A=5-10, and $\zeta = 0.05 - 0.1$.

B. Simulation procedure and the projected energy

The simulations are started by placing a single walker on the HF determinant. The simulation is then run for a start-up period, in which the shift *S* is kept at a constant value (of zero or small positive number). This results in the number of walkers initially growing very rapidly. When a desired number of walkers is reached, we start to adjust the shift *S*, as given by Eq. (17). It is observed that the value of the shift initially falls, and then stabilizes around a value about which it oscillates. If the number of walkers is above a critical number (to be discussed later), this value is the correlation energy of the system. It is desirable to have another measure of the correlation energy. This is provided by the following projection:

$$E(\tau) = \frac{\langle D_0 | He^{-\tau H} | D_0 \rangle}{\langle D_0 | e^{-\tau H} | D_0 \rangle} \tag{18}$$

$$=E_{\mathrm{HF}} + \sum_{\mathbf{j} \neq \mathbf{0}} \langle D_{\mathbf{j}} | H | D_{\mathbf{0}} \rangle \frac{C_{\mathbf{j}}(\tau)}{C_{\mathbf{0}}(\tau)}$$
(19)

$$=E_{\mathrm{HF}} + \sum_{\mathbf{j} \neq \mathbf{0}} \langle D_{\mathbf{j}} | H | D_{\mathbf{0}} \rangle \frac{N_{\mathbf{j}}(\tau)}{N_{\mathbf{0}}(\tau)}, \tag{20}$$

where $N_{\mathbf{j}}(\tau)$ is the number of walkers [defined by Eq. (13)] on $D_{\mathbf{j}}$ at time step τ , and N_0 is this number on $D_{\mathbf{0}}$ (which we assume to be the Hartree–Fock determinant). In addition, only the singles and double excitations are connected to $D_{\mathbf{0}}$ (the singles being connected only in case of open-shell systems), it is only the populations at these excitations that need to be evaluated for the purposes of computing the energy, i.e.,

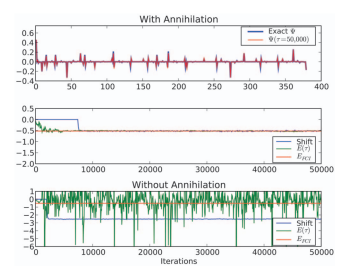


FIG. 1. Comparison of sampling of determinants compared to full diagonalization result for a stretched N_2 molecule in a space of 379 determinants. Ψ indicates the normalized walker number on each of the determinants after 50 000 iterations, comparing it to the exact wave function. The shift and E_0 values for each iteration are shown in the lower plot.

$$E(\tau) = E_{\rm HF} + \sum_{\mathbf{j} \in \{\text{singles,doubles}\}} \langle D_{\mathbf{j}} | H | D_{\mathbf{0}} \rangle \frac{N_{\mathbf{j}}(\tau)}{N_{\mathbf{0}}(\tau)}. \tag{21}$$

In the long-time limit, it is evident that $E(\tau)$ converges onto the exact ground-state energy:

$$E_0 = \lim_{\tau \to \infty} E(\tau) \tag{22}$$

as long as the ratio N_j/N_0 equals C_j/C_0 of the FCI wave function.

The two measures of the correlation energy, i.e., S and $E(\tau)$, are to a large extent independent of each other, since S depends on the total number of walkers, whereas $E(\tau)$ depends only on the population of walkers on the Hartree–Fock determinant and the singles and double excitations, which typically constitute only a small fraction of the total walker population. In the limit of large walker populations, the variations in S and $E(\tau)$ become largely uncorrelated.

IV. AN ILLUSTRATIVE EXAMPLE

Let us consider stretched N₂, which is a strongly multiconfigurational system. For the purposes of illustration of how the exact wave function emerges during the course of the simulation, we consider here a very small basis, with 379 Slater determinants, which allows a convenient visualization of the evolution of the walker distribution as the simulation proceeds. The behavior of the shift and of the projected energy are shown in Fig. 1. The instantaneous energy for each time step [Eq. (20)] converges on the FCI energy within 10 000 steps. In this simulation, we started varying the shift at step 10 000, and the shift very rapidly decreases from zero toward the FCI energy, and stabilizes at the correct value. The two independent measures of the correlation energy agree to better than 1 mhartree. In real applications, where the exact FCI energy is not known, the coincidence of the these two measures provides a strong degree of confidence in the obtained correlation energies. Equally remarkable is the

convergence of the walker distribution onto the exact wave function: An animation of the time evolution of this simulation (as well as other systems) is also provided as a movie held on our website.³¹ In the third panel of Fig. 1 we also show the behavior of the simulation if the annihilation step is removed. In this case, the shift does not stabilize at the correct value, and the instantaneous energy fluctuates wildly. Without annihilation, the walker distribution cannot converge onto to correct ground-state distribution; there is no "interaction" between the positive and negative walkers, and, in fact, the distribution of the positive walkers among the determinants increasingly matches the distribution of negative walkers as the simulation proceeds. In other words, in common with the classic sign problem, the signal to noise ratio decreases exponentially quickly. In addition, because the shift goes to the wrong (and much too negative) value, the rate of death processes becomes very large, leading to indiscriminate death of walkers, essentially irrespective of which determinant they occupy. The walker distribution is not able to build a signal on the determinants with significant weight. This example shows the crucial role played by the annihilation step. In fact the annihilation step allows a symmetry breaking to occur, in which one of the two allowed solutions, either $+\Psi$ or $-\Psi$, is settled upon. Without providing any information regarding the sign structure of the FCI wave function, our algorithm is capable of settling down onto the correct wave function. In a real sense, the manybody wave function has emerged during the course of the simulation. The breaking of the $\pm \Psi$ symmetry is a necessary condition for this emergence. An analogy can be made with phase transitions in classical spin systems. The ordered phase (e.g., ferromagnetic or antiferromagnetic) of a spin system is degenerate with respect to overall sign change in all spins. For the system to acquire net magnetization (or staggered magnetization) below the critical point, the symmetry must be spontaneously broken, and the ordered phase, which has a symmetry lower than that present in the Hamiltonian can then emerge.

This example demonstrates that the method, in principle, works exactly. The question now is: It is capable of working for much larger FCI spaces? To answer this question, we have studied a number of molecular systems (Ne, H_2O , C_2 , N_2 , etc.) in a variety of basis sets, to which we now turn.

V. RESULTS

We next consider a range of molecules, treated in more realistic Dunning basis sets. ³² We also perform a benchmark study of the all-electron Ne atom in several families of basis sets, namely, cc-pVXZ, aug-cc-pVXZ, and cc-pCVXZ (X = D, T, Q). The size of spaces considered range from $\sim 10^6$ to $\sim 10^{14}$.

In spaces of such size, the pattern of walker growth generally changes. Keeping the shift fixed, initially the walker number grows exponentially. However, at a system specific value, this exponential growth attenuates, apparently suddenly, and the number of walkers hits a plateau. After a period of essentially zero growth in overall walker number, the growth picks up again, after which point we start adjusting

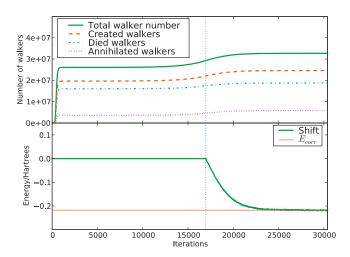


FIG. 2. A typical trend of walker growth for $\rm H_2O$ in a cc-pVDZ basis set in constant shift mode (S=0) until iteration 16 970, showing the appearance of a well-defined plateau in N_w , in this case at N_c =26×10⁶ walkers. For this system, $N_{\rm FCI}$ =451×10⁶. The number of walkers at the plateau is therefore about 6% of the FCI space. Also shown on the plot are the numbers of walkers created, died, and annihilated per A=10 iterations. It is evident that at the plateau, the combined rate of death and annihilation matches the birth rate from the spawning. The plateau gradually gives way to a growth phase, in walker number, which increases exponentially. Once in the growth phase, the shift is then allowed to vary according to Eq. (17), (in this example on iteration number 16 970). The number of walkers then rapidly stabilizes. The lower plot shows that the value for the shift which stabilizes the walker growth is exactly the correlation energy for the system.

the shift. An example of this is shown for the all-electron water molecule treated with a cc-pVDZ basis in Fig. 2.

During this plateau phase, the rate of walker birth is exactly matched by the combined rate of walker death and walker annihilation. Before reaching the plateau, the rate of annihilation is small and therefore the overall rate of walker growth is positive. At the plateau, sufficient numbers of walkers begin to annihilate each other, and this leads to the attenuation of walker number growth. Very significantly, owing to walker annihilation, we observe that during the plateau phase, the sign structure of the FCI wave function is converged upon, in every case we have studied. As long as we have a larger number of walkers in the system than the number defined by the plateau position, N_c , then when we allow the shift to vary (in constant N_w mode), it converges onto the correlation energy. This N_c value is therefore an important, system-dependent parameter which indicates the required sampling of the space. If this sampling can be achieved, then the correct correlation energy for the system can be confidently converged upon to the desired accuracy. This N_c value in all cases is smaller than the size of the complete space, often substantially so. We can therefore describe a parameter f_c which gives us the relative number of walkers required, compared to the number of determinants in the full space $(f_c = N_c/N_{FCI})$. f_c is one measure of the difficulty the method has in achieving convergence. Where f_c is small (or even zero), convergence can be achieved with a relatively small number of walkers.

It turns out that the value of N_c (for a given system) is remarkably insensitive to the initial conditions, and method of equilibration, of the walkers. Thus, we have found that we need the same number of walkers to attain the plateau if we

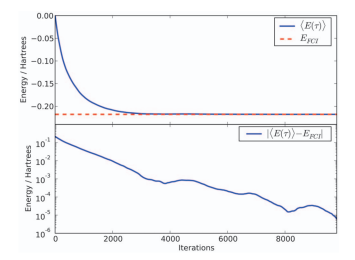


FIG. 3. Convergence of the energy estimator averaged over all previous iterations, showing an exponential convergence to the $E_{\rm FCI}$ for a cc-pVDZ water system at equilibrium geometry. The lower plot shows the absolute difference between the energy and the exact energy on a logarithmic scale. $\delta\tau=1\times10^{-3}$ a.u. In this example, the simulation was started by placing a single walker at the Hartree–Fock determinant and allowing the population to grow throughout the entire space until the postplateau growth phase has been reached.

start the simulation with one walker and let the number grow rapidly throughout the entire FCI space, or if we gradually enlarge the space, e.g., via excitation level. This indicates that N_c is an intrinsic parameter which characterizes the system, for a given one-electron basis.

In Fig. 3 we show the convergence of the averaged projected energy $\langle E(\tau) \rangle$ as the simulation proceeds, in the case of the water molecule. Also included, is the absolute error $|\langle E(\tau) \rangle - E_{\text{FCI}}|$ from the known FCI energy. It can be seen that there is an exponential convergence to an accuracy of under 0.1 mhartree of the energy. To give an idea of the computational cost of the method, the CPU time taken to converge all-electron cc-pVDZ water to this accuracy with our current code is approximately 3 hours using a quad-core (2.66 GHz) PC.

Apart from the water molecule, we have tested our algorithm in other systems for which FCI results are available for comparison, shown in Table I, namely, Ne atom, C_2 and N_2 at equilibrium and at stretched geometry. As can be seen, the computed energies are to within 1 mhartree of the published

FCI results for all systems considered, confirming the ability of the method to reproduce the FCI energies across a broad range of systems.

There is considerable variation in f_c among these systems. For the Ne atom, and all-electron H_2O , $f_c \le 0.1$, whereas for C_2 and N_2 , $f_c \sim 0.5$, i.e., comparable though still smaller than the FCI space. The behavior of the two N₂ systems is particularly instructive. As the N₂ bond is stretched from its equilibrium geometry, the description of the problem changes from one which is essentially single reference, to one which is highly multireference with significant contributions up to hextuple excitations. This makes an accurate description of the binding curve of nitrogen particularly challenging for many methods. Surprisingly, the results from these two systems show a similar f_c , indicating that the ease with which the method achieves convergence is not simply related to the dominance of the Hartree-Fock determinant, and that the number of significant determinants is not the key factor in its efficiency. This feature can hopefully be exploited in the study of multiconfigurational systems. The fact that in all cases f_c is less than one, sometimes substantially so, means that one needs fewer walkers than determinants to achieve converge the calculation. At any one time step, only a fraction of the space is occupied, but as long as f_c has been exceeded, averaged over time each determinant is correctly sampled according to its amplitude. We have exploited this property to study systems for which the FCI space is extremely large.

In Table II, we report the energies of several molecules, including CO, CH₄, O₂, and NaH—for which we have not been able to find published FCI energies, and therefore serve as predictions which could be verified by explicit FCI calculations. Included in this list are the open-shell systems $CN(^2\Sigma^+)$ and $O_2(^3\Sigma_{\rho}^-)$. As can be seen from the table, the size of FCI spaces for these systems range from a few million to over 10^{11} . Particularly striking are the variations in f_c . The worst case is for CH_4 , where we find an f_c =0.898, while for NaH $f_c = 3 \times 10^{-4}$. Remarkably, in this case, only 64 $\times 10^6$ walkers were needed to converge the energy for a space of 205×10^9 determinants. Results from coupled cluster with perturbative triple excitations, CCSD(T), have been included for comparison,³⁸ and as can be seen our correlation energies are generally slightly larger (with the notable exception of CH₄), although since CCSD(T) is not variational, not

TABLE I. Results for systems with FCI comparisons. The geometries for the N_2 molecule were eqm: $2.068a_0$, stretched: $4.2a_0$, and C_2 : 1.272.73 Å. The geometry for the water molecule was taken from Ref. 35. The working space includes all point group symmetry of the molecule from D_{2h} or the largest available subset thereof. All systems had core electrons frozen apart from H_2O . N_{FCI} is the size of the FCI space in the D_{2h} point group (C_{2v} for H_2O . The digit in italics for E_{total} , represents the first uncertain digit. N_c is the number of walkers required to achieve the plateau. $f_c = N_c/N_{FCI}$.

| System | (N,M) | $N_{\rm FCI}/10^6$ | $N_c/10^6$ | f_c | $E_{ m total}$ | $E_{ m FCI}$ | Reference |
|------------------------------------|---------|--------------------|------------|-------|----------------|--------------|-----------|
| Ne: aug-cc-pVDZ | (8,22) | 6.69 | 0.21 | 0.031 | -128.70949 | -128.709,476 | 33 |
| C ₂ : cc-pVDZ | (8,26) | 27.9 | 15.0 | 0.538 | -75.7299 | -75.729,853 | 34 |
| H ₂ O: cc-pVDZ | (10,24) | 451 | 26 | 0.058 | -76.24186 | -76.241,860 | 35 |
| N ₂ -eqm: cc-pVDZ | (10,26) | 541 | 270 | 0.499 | -109.27649 | -109.276,527 | 33 |
| N ₂ -stretched: cc-pVDZ | (10,26) | 541 | 345 | 0.637 | -108.9669 | -108.966,95 | 36 |

TABLE II. Predicted FCI results. The geometries of the molecules were (in Å): CN (1.1941), HF (0.91622), CH₄($r_{\rm CH}$ =1.087 728), CO (1.1448), H₂O($r_{\rm OH}$ =0.975 512, θ =110.565°) (Ref. 35), O₂ (1.2074), and NaH (1.885 977). CN and O₂ orbitals were constructed from a restricted open-shell HF calculation with a spin multiplicity of two and three, respectively. CN, CH₄, CO, and O₂ had frozen core electrons. The number in brackets represents the error in the previous digit, obtained through a Flyvbjerg–Petersen blocking analysis (Ref. 37) of $E(\tau)$.

| System | (N,M) | $N_{\rm FCI}/10^6$ | $N_c/10^6$ | f_c | $E_{ m total}$ | $E_{\text{CCSD(T)}}$ |
|----------------------------|---------|--------------------|------------|----------|----------------|----------------------|
| Be: cc-V5Z | (4,91) | 2.11 | 0 | 0 | -14.646 38(2) | -14.646 29 |
| CN: cc-pVDZ | (9,26) | 246 | 173 | 0.704 | -92.493 8(3) | -92.491 64 |
| HF: cc-pCVDZ | (10,23) | 283 | 0.998 | 0.0035 | -100.27098(3) | -100.27044 |
| CH ₄ : cc-pVDZ | (8,33) | 419 | 377 | 0.898 | -40.38752(1) | -40.38974 |
| CO: cc-pVDZ | (10,26) | 1080 | 777 | 0.719 | -113.05644(4) | -113.05497 |
| H ₂ O: cc-pCVDZ | (10,28) | 2410 | 47.4 | 0.0196 | -76.28091(3) | -76.28028 |
| O ₂ : cc-pVDZ | (12,26) | 5409 | 2651 | 0.490 | -149.9875(2) | -149.98562 |
| NaH: cc-pCVDZ | (12,32) | 205 300 | 63.8 | 0.000 31 | -162.609 0(1) | -162.609 01 |

too much can be read into this result. We do note that the agreement between the two is generally better for closed-shell systems than open-shell ones.

An important question with the present method is how the plateau varies for a given system as the basis set is improved, since we typically want to do calculations in substantial basis sets, which improve accuracy and allow extrapolation to the complete basis-set limit.³⁹ In Table III, we present a study of the Ne atom in various basis sets, to investigate the fraction f_c required to achieve the plateau. Several trends are discernible. Within a given type of basis set, the value of f_c is fairly constant, but there are marked variations between different families. Thus cc-pVXZ [denoted VXZ in Table III, where X=D,T,Q family have $f_c \approx 5$ $\times 10^{-4}$, whereas the aug-cc-pVXZ (AVXZ) family have a larger $f_c \approx 1.5 \times 10^{-3}$, while the cc-pCVXZ (CVXZ) have a very favorable $f_c \approx 2 \times 10^{-5}$. The latter are also give the most accurate energies. Clearly inclusion of core electrons with CVXZ basis sets is very favorable for the present method. Extrapolation to the complete basis-set limit³⁹ yields a correlation energy of 393.0 mhartree, which is within 2.5 mhartree of the "exact" nonrelativistic result. 40 Such extrapolated energies are of course not variational, and are known to yield errors of about 5 mhartree, consistent with what is observed here.

These results illustrate that the memory requirements of

the method are substantially lower than a conventional FCI calculation. For example, the all-electron Ne atom in a cc-CVQZ basis required N_c =2.2×10 9 walkers, which amounts to 62 Gbytes of RAM. By contrast, a FCI calculation of the same system would require a minimum of two vectors of length $N_{\rm FCI}$ =131×10 12 , i.e., over 2×10 6 Gbytes, a vastly greater amount of storage.

Since in general N_c scales linearly with $N_{\rm FCI}$, this implies that the present method has exponential scaling with N and M. The method, therefore, does not constitute a "solution to the sign problem," for which a polynomial scaling is required. It is perhaps best thought of as an alternative method to FCI, with a smaller prefactor (proportional to f_c), which in some cases is substantially so. It remains to be seen if variations on the present algorithm can be found in which either the prefactor can be further suppressed, or even better, if the scaling can be improved upon.

VI. CONCLUSIONS

We have described a new QMC algorithm in Slater determinant space, which we show is able to converge on the FCI energy of the system under consideration without any *a priori* information, as long as a system-dependent critical number of walkers is exceeded. The exact FCI wave functions emerge spontaneously once this critical number has

TABLE III. Ne atom results in Dunning (Ref. 32) basis sets, giving correlation energies in hartrees. All electrons (10) were correlated over all excitation levels. The "exact" result is the nonrelativistic infinite-nuclear mass corrected experimental value, calculated in Ref. 40. The extrapolated result for comparison is obtained using the technique by Halkier *et al.* (Ref. 39) using the cc-pCVTZ and cc-pCVQZ correlation energies.

| Basis set | Orbitals | $N_{\rm FCI}/10^6$ | $N_c/10^6$ | $f_c/10^{-3}$ | $E_{ m corr}$ |
|---------------|----------|----------------------|------------|---------------|---------------|
| VDZ | 14 | 0.502 | 0 | 0 | 0.19211(4) |
| CVDZ | 18 | 9.19 | 0 | 0 | 0.23365(3) |
| AVDZ | 23 | 142 | 0.248 | 1.7 | 0.21510(3) |
| VTZ | 30 | 2540 | 0.506 | 0.199 | 0.28341(9) |
| CVTZ | 43 | 116 000 | 2.3 | 0.0198 | 0.33628(2) |
| AVTZ | 46 | 235 000 | 338 | 1.43 | 0.2925(4) |
| VQZ | 55 | 1.51×10^{6} | 681 | 0.451 | 0.3347(10) |
| CVQZ | 84 | 119×10^{6} | 2200 | 0.0185 | 0.3691(1) |
| Extrapolation | | | | | 0.3930 |
| Exact | | | | | 0.3905 |

been reached. Walker annihilation is shown to play a key role in this process. The computational requirements of the method are largely determined by the value of this critical number. If it can be reduced for a given system, e.g., by a judicious choice of orbitals obtained via orthonormal transformations of the canonical Hartree–Fock orbitals, then the computational effort required to compute the correlation energy will be similarly reduced. We are currently investigating such ideas.

Once this critical number of walkers has been reached, the correlation energy can be computed with confidence. Agreement between two estimators of the correlation energy provides additional support for the values obtained. We have reproduced existing FCI energies, as well as reporting new systems with very large FCI spaces. Favorable memory requirements, as well as ease of parallelization, are attributes of this algorithm which should enable yet larger systems to be tackled in the near future.

In common with the FCI method, the only systematic error in the calculation of the correlation energy arises through basis-set incompleteness, which can however be systematically improved via explicit calculation on larger basis sets, together with extrapolation to the complete basis-set limit. This is marked contrast with the uncontrolled approximation of fixed-node QMC where the error introduced is hard to reduce systematically.

The present method provides a synthesis of QMC and quantum chemistry. Single and multireference problems can both be tackled, and the difficulty of the procedure does not seem to be closely tied to this classification of systems. This suggests that future effort will be primarily focused on multireference systems which provide a sterner challenge to many other methods. In addition to this, several extensions of the present technique can be envisaged which should allow much larger systems to be treated. These include CASSCF methodologies, perturbation theory extensions, and the use of configuration state functions, which are all currently under investigation.

APPENDIX A: HAMILTONIAN MATRIX ELEMENTS

Two determinants $|D_{\bf i}\rangle$ and $|D_{\bf j}\rangle$ are said to be coupled if and only if $\langle D_{\bf i}|H|D_{\bf j}\rangle\neq 0$. In our applications, H is a molecular Hamiltonian with one-electron and two-electron terms:

$$H = \sum_{i}^{N} h_{i} + \sum_{i < j} \frac{1}{r_{ij}},$$
 (A1)

$$h_i = -\frac{1}{2}\nabla_i^2 + v_{\text{ext}}(\mathbf{r}_i). \tag{A2}$$

Determinants which differ by three or more (orthonormal) spin orbitals are therefore uncoupled, and we need only consider pairs of determinants which differ by two or fewer spin orbitals, i.e., determinants which are double or single excitations of each other. Such matrix elements can be computed using the Slater–Condon rules $^{41-43}$ as follows. In the case of double excitations, let $D_{\bf j}$ be the following double excitation of $D_{\bf i}$:

$$|D_{\mathbf{i}}\rangle = a_r^{\dagger} a_s^{\dagger} a_o a_p |D_{\mathbf{i}}\rangle. \tag{A3}$$

Then

$$\langle D_{\mathbf{i}}|H|D_{\mathbf{i}}\rangle = \langle rs||pq\rangle \equiv \langle rs|pq\rangle - \langle rs|qp\rangle,$$
 (A4)

where the (two-electron) four-index integrals are defined by

$$\langle rs|pq\rangle = \int d\mathbf{r}_1 d\mathbf{r}_2 \phi_r^*(\mathbf{r}_1) \phi_s^*(\mathbf{r}_2) \frac{1}{r_{12}} \phi_p(\mathbf{r}_1) \phi_q(\mathbf{r}_2).$$
 (A5)

Similarly, in the case of single excitations, let $|D_{\mathbf{j}}\rangle = a_r^{\dagger} a_p |D_{\mathbf{i}}\rangle$, then

$$\langle D_{\mathbf{i}}|H|D_{\mathbf{j}}\rangle = \langle r|h|q\rangle + \sum_{k}' \langle rk||qk\rangle,$$
 (A6)

where the sum over k extends over the N-1 spin orbitals common to D_i and D_i .

The diagonal matrix elements are given by

$$\langle D_{\mathbf{i}}|H|D_{\mathbf{i}}\rangle = \sum_{p \in \mathbf{i}} \langle p|h|p\rangle + \frac{1}{2} \sum_{p,q \in \mathbf{i}} \langle pq||pq\rangle. \tag{A7}$$

In the present work, all the necessary integrals (four-index and two-index) were generated from restricted Hartree–Fock orbitals using a modified version of QCHEM.³⁸

APPENDIX B: EXCITATION GENERATION ALGORITHM

In our algorithm, it is necessary to be able to generate all single or double excitations of any determinant in such a way that its generation probability, $p_{\rm gen}(\mathbf{j}|\mathbf{i})$ is computable, nonzero, and normalized. We employed the following simple strategy (which is probably not optimal from a sampling perspective; but this consideration is left for a future study).

Let us consider the generation of $|D_{\mathbf{j}}\rangle = a_r^{\dagger} a_s^{\dagger} a_q a_p |D_{\mathbf{i}}\rangle$, which involves the selection of the occupied pair (p,q) and the unoccupied pair (r,s) (with respect to $D_{\mathbf{i}}$). Then

$$p_{\text{gen}}(\mathbf{j}|\mathbf{i}) = p_{\text{gen}}(r,s|p,q)p_{\text{gen}}(p,q), \tag{B1}$$

where $p_{\text{gen}}(p,q)$ is the probability to select the orbital pair (p,q) in $D_{\mathbf{i}}$, and $p_{\text{gen}}(r,s|p,q)$ is the probability to select the orbitals pair (r,s) given that we have selected (p,q). For an N electron system, we select the occupied pair (p,q) with uniform probability, i.e.,

$$p_{\text{gen}}(p,q) = {N \choose 2}^{-1} = 2[N(N-1)]^{-1}.$$
 (B2)

We further write

$$p_{gen}(r,s|p,q) = p_{gen}(r|s,p,q) \cdot p_{gen}(s|p,q) + p_{gen}(s|r,p,q) \cdot p_{gen}(r|p,q).$$
(B3)

In other words, we select s with probability $p_{\rm gen}(s|p,q)$, and r with probability $p_{\rm gen}(r|s,p,q)$, before computing the probability that we *could* have picked the unoccupied orbital r first, $p_{\rm gen}(r|p,q)$, followed by s, to obtain the same excitation. Generally, $p_{\rm gen}(r|s,p,q)\neq p_{\rm gen}(s|r,p,q)$. The advantage of this approach is that we can combine spin and symmetry information, which is usually available in the form of the irreducible representation spanned by each spatial orbital. For example, in the generation of r, given (s,p,q), the irre-

ducible representation (Γ_r) of r is dictated to be the direct product of the irreps of (s, p, q):

$$\Gamma_r = \Gamma_s \otimes \Gamma_p \otimes \Gamma_a \tag{B4}$$

[In the case of Abelian groups this uniquely determines Γ_r . In this study of atoms and homonuclear diatomics, we used D_{2h} as such a group]. Having selected s, we then determine the irreducible representation of r, and then explicitly count how many unoccupied orbitals of this irreducible representation are available for selection (imposing, in addition, that the net change in spin polarization is zero). The reciprocal of this number gives $p_{\text{gen}}(r|s,p,q)$. This completes the generation algorithm for double excitations.

The same method can be used to compute the generation probability of a single excitation, although it is much simpler. Given $|D_{\bf i}\rangle = a_r^{\dagger} a_p |D_{\bf i}\rangle$, we need

$$p_{gen}(r,p) = p_{gen}(r|p) \cdot p_{gen}(p)$$
. (B5)

We select p uniformly out the occupied orbitals of $D_{\bf i}$, (i.e., $p_{\rm gen}(p) = N^{-1}$) and then select r such that $\Gamma_r = \Gamma_p$. We then count the number of unoccupied orbitals of this irreducible representation that are available with the same spin, and compute $p_{\rm gen}(r|p)$ as the reciprocal of this number. The probability of choosing to create a double excitation is given by P_d , where $0 < P_d < 1$ and hence the probability of choosing to generate a single excitation is $1 - P_d$. P_d is chosen to approximately reflect the relative number of double excitations compared to single excitations. To obtain the final $p_{\rm gen}({\bf j}|{\bf i})$, we need to multiply by P_d if it is a double excitation, or $1 - P_d$ if it is a single excitation to maintain normalization of the probabilities.

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