Systematic reduction of sign errors in many-body calculations of atoms and molecules

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The self-healing diffusion Monte Carlo algorithm (SHDMC) [Phys. Rev. B 79, 195117 (2009), ibid. 80, 125110 (2009)] is shown to be an accurate and robust method for calculating the ground-state of atoms and molecules. By direct comparison with accurate configuration interaction results for the oxygen atom we show that SHDMC converges systematically towards the ground-state wave function. We present results for the challenging N2 molecule, where the binding energies obtained via both energy minimization and SHDMC are near chemical accuracy (1 kcal/mol). Moreover, we demonstrate that SHDMC is robust enough to find the nodal surface for systems at least as large as C20 starting from random coefficients. SHDMC is a linear-scaling method, in the degrees of freedom of the nodes, that systematically reduces the fermion sign problem.

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Since electrons are fermions, their many-body wave functions must change sign when the coordinates of any pair are interchanged. In contrast, the sign of a bosonic wave functions is unchanged for any coordinate interchange. Due to this misleadingly small difference, the ground-state energy of bosons can be determined by quantum Monte Carlo (QMC) methods1,2 with an accuracy limited only by computing time, while QMC calculations of fermions are either exponentially difficult, or are stabilized by imposing a systematic error, a direct consequence of our lack of knowledge of the fermionic nodal surface. Therefore, one of the most important problems in many-body electronic structure theory is to accurately find representations of the fermion nodes3,4, the locations where the fermionic wave function changes sign, the so-called “fermion sign problem”.

The sign problem limits (i) the number of physical systems where ab initio QMC can be applied and (ii) our ability to improve approximations of density functional theory (DFT) using QMC results5. More importantly, it limits our overall understanding of the effects of interactions in fermionic systems. Therefore, a method to circumvent the sign problem with reduced computational cost could transform Condensed Matter Theory, Quantum Chemistry and Nuclear Physics among other fields.

Arguably the most accurate technique for calculating the ground-state of a many-body system with more than 20 fermions is diffusion Monte Carlo (DMC). The standard DMC algorithm5 finds the lowest energy of all wave functions that share the nodal surface \( S_T(\mathbf{R}) \) imposed by a trial wave function \( \Psi_T(\mathbf{R}) \). This is the fixed-node approximation where the resultant energy \( E_{DMC} \) is a rigorous upper bound of the exact ground-state energy \( E_T \). The exact ground-state energy is obtained only when \( \Psi_T(\mathbf{R}) \) has the same nodal surface as the exact ground-state wave function.

If the exact nodes are not provided, the implicit fixed-node ground-state wave function \( \Psi_{FN}(\mathbf{R}) \) will exhibit discontinuities in its gradient \( \delta \) (i.e. kinks) on some parts of \( S_T(\mathbf{R}) \). We recently proved \( \delta \) that by locally smoothing these discontinuities in \( \Psi_{FN}(\mathbf{R}) \), a new trial wave function can be obtained with improved nodes. This proof enables an algorithm that systematically moves the nodal surface \( S_T(\mathbf{R}) \) towards the one of an eigen-state. If the form of trial wave function is sufficiently flexible, and given sufficient statistics, this process leads to an exact eigen-state wave function \( \delta \delta \). We named the method self-healing DMC (SHDMC), since the trial wave function is self-corrected in DMC and can recover even from a poor starting point.

In this Letter, we report the first applications of SHDMC to real atoms and molecules (O, N2, C20). SHDMC energies are within error bars of DMC calculations using the current state of the art approach \( \delta \delta \). Tests of SHDMC for C20 demonstrate our method can be applied at the nanoscale. Its cost scales linearly with the number of independent degrees of freedom of the nodes with an accuracy limited only by the achievable statistics and choice of representation of the nodes.

Brief review of SHDMC — SHDMC is fundamentally different from optimization methods used in variational Monte Carlo (VMC): \( \delta \delta \) (i) the wave function is directly optimized based on a property of the nodal surface and not on the local energy or its variance, and (ii) the nodes are optimized at the DMC level (as opposed to a VMC based algorithm).

Using a short-time many-body propagator, SHDMC samples the coefficients of an improved wave function removing the artificial derivative discontinuities of \( \Psi_{FN}(\mathbf{R}) \) arising from the inexact nodes. Repeated application of this method results in the best nodal surface for a given basis. For wave functions expanded in a complete basis it can be shown that the final accuracy is limited only by the statistics \( \delta \delta \).

In SHDMC (see Refs. \( \delta \delta \) for details), the weighted walker distribution is

\[
f(\mathbf{R}, \tau' + \tau) = \Psi_T^*(\mathbf{R}, \tau') \left[ e^{-\tau(\hat{N}_{FN} - E_T)} \Psi_T(\mathbf{R}, \tau') \right]
\]

where

\[
f(\mathbf{R}, \tau' + \tau) = \lim_{{N_c \to \infty}} \frac{1}{N_c} \sum_{{i=1}}^{{N_c}} W_i^j(k)\delta \left( \mathbf{R} - \mathbf{R}_i^j \right),
\]
with

$$\Psi_T(R, \tau') = e^{J(R)} \sum_n \lambda_n(\tau') \Phi_n(R)$$  \hspace{1cm} (2)$$

is a trial function where $\sum_n$ represents a truncated sum, \{\(\Phi_n(R)\)\} forms a complete orthonormal basis of the antisymmetric Hilbert space and $e^{J(R)}$ is a symmetric Jastrow factor. In Eq. (1), $H_{FN}$ is the fixed-node Hamiltonian [$H_{FN}$ is the many-body Hamiltonian with an infinite potential at the nodes of $\Psi_T(R, \tau')$] and $E_T$ is an energy reference. Next, $R_i^j$ corresponds to the position of the walker $i$ at step $j$ of $N_c$ equilibrated configurations. The weights $W_i^j(k)$ are given by

$$W_i^j(k) = e^{-[E_i^k(\tau') - E_T]^2}$$ with $E_i^k(\tau') = \sum_{\ell=0}^{k-1} E_L(R_i^{\ell})$, \hspace{1cm} (3)

where $E_T$ in Eq. (3) is periodically adjusted so that $\sum_i W_i^j(k) \approx N_c$ and $\tau$ is $k\delta \tau$ (with $k$ being a number of steps and $\delta \tau$ a standard DMC time step).

From Eq. (1), one can formally obtain

$$\Psi_T(R, \tau' + \tau) = f(R, \tau', \tau)/\Psi_T(R, \tau') \hspace{1cm} (4)$$

We now define the local smoothing function to be

$$\bar{\delta}(R', R) = \sum_n e^{-J(R')} \Phi_n(R') \Phi_n^*(R) e^{-J(R)}.$$  \hspace{1cm} (5)

Applying Eq. (5) to both sides of Eq. (4), using Eq. (1), and integrating over $R$ we obtain

$$\Psi_T(R, \tau' + \tau) = e^{J(R)} \sum_n \lambda_n(\tau' + \tau) \Phi_n(R),$$  \hspace{1cm} (6)

with

$$\lambda_n(\tau' + \tau) = \lim_{N_c \to \infty} \frac{1}{N_c} \sum_{i=1}^{N_c} W_i^j(k) e^{-2J(R_i^j)} \frac{\Phi_n^*(R_i^j)}{\Phi_n^*(R_i^j, \tau')}$$  \hspace{1cm} (7)

where $N_c = \sum_{i=1}^{N_c} e^{-2J(R_i^j)}$ normalizes the Jastrow factor. These new $\lambda_n(\tau' + \tau)$ [Eq. (7)] are used to construct a new trial wave function [Eq. (4)] recursively within DMC (therefore the name self-healing DMC). The weights in Eq. (5) can be evaluated within (i) a branching algorithm \[8\] for $\tau' \to \infty$ or (ii) a fixed population scheme for small $\tau'$ \[9\]. The former method is more robust, but the latter improves final convergence. Equation (7) can be related to the maximum-overlap method used for bosonic wave functions \[10\].

Since SHDMC is targeted for large systems we report validations using pseudopotentials.

**Validation of SHDMC with configuration interaction (CI) calculations for the O atom** — In short, CI is the diagonalization of the many-body Hamiltonian in a truncated basis of Slater determinants. We chose to study the $^3P$ ground-state of the O atom because it has at least two valence electrons in both spin channels \[14\]. The single-particle orbitals were expanded in VTZ and V5Z Gaussian basis sets \[14\] using the GAMESS \[16\] code. To facilitate a direct comparison between SHDMC and CI, no Jastrow factor was employed.

Figure 1 shows a direct comparison of the first 250 converged coefficients $\lambda_n$ obtained using SHDMC with those from the largest CI calculation (see Table 1). The initial SHDMC trial wave function was the Hartree–Fock (HF) solution, and the final SHDMC coefficients resulted from sampling the 1481 most significant excitations in the CI. We used $\delta \tau = 0.01$ a.u., $\tau = 0.5$ a.u., and 16 iterations of trial wave function projection ($\approx 6 \times 10^7$ sampled configurations).

Figure 1 shows the excellent agreement between the coefficients $\lambda_n$ obtained independently by SHDMC and CI. A perfect agreement is guaranteed only in the limit of a complete basis and $N_c \to \infty$. The small differences in Fig. 1 are due to the truncation of the expansion and the stochastic error in $\lambda_n$. The inset shows the residual projection as a function of the total number $N_b$ of CSFs included in the expansion, normalized either using the entire CI expansion (circles) or using a $\Psi_{CI}$ that included only the $\lambda_n$ sampled in SHDMC (squares). The residual projection is much smaller for the truncated norm than the full norm illustrating that most of the error in $\Psi_{SHDMC}$ is from truncation and not limited statistics. Similar results were obtained for the C atom (not shown).

**Validation with Energy Minimization for $N_b$** — We also compared the VMC and DMC energies of wave functions optimized with energy minimization in VMC (EMVMC) \[10\], \[11\] and SHDMC using the QWALK \[18\] code. EMVMC can be briefly described as a generalized CI with an additional Jastrow factor (sampling the Hamiltonian stochastically and solving a generalized eigenvalue problem). Several basis sets were obtained from series of complete active space (CAS) and restricted active space (RAS) \[12\] multiconfiguration self-consistent field (MCSCF) calculations [distributing 10 electrons into $m$ active orbitals: CAS($10,m$)]. We retained the $N_b$ basis functions with coefficients of absolute value larger than a given cutoff. Subsequently, for each basis, we performed energy minimization of the Jastrow and the coefficients of trial wave function using a mixture of 95% of energy and 5% of variance. We also sampled these $N_b$ coefficients in SHDMC recursively starting from HF solution. For a clear comparison

| Method       | $N_b$ | $E$ [Ha] [%] | $N_b$ | $E$ [Ha] [%] |
|--------------|-------|-------------|-------|-------------|
| CI \[a\]    | 775182 | -15.88258(89.0) | 1762377 | -15.8957(95.7) |
| CCSD(T) \[b\] | -15.88204(88.8) | -15.90166(98.8) |
| SHDMC        | 539   | -15.9040(4)\{100.0(2)\} | 1481   | -15.9040(4)\{100.0(2)\} |

\[a\] full-CI in VTZ and CISDTQ in V5Z.
\[b\] from Ref. \[13\].
we used the same Jastrow in EMVMC and SHDMC.

We performed these calculations for the ground-state \( ^1\Sigma^+ \) of \( \text{N}_2 \) at the experimental geometry \((20)\). Figure 2 shows the resulting VMC and DMC energies obtained for wave functions optimized independently with EMVMC and SHDMC methods for the largest RAS(10,43) (2629447 CSFs yielding \( E=\text{-}19.921717 \)) Slater-Jastrow wave function (See also Table II). In EMVMC, as previously observed for \( \text{C}_2 \) and \( \text{Si}_2 \) \((11)\), we found a systematic reduction in the fixed-node errors, even when starting from the smallest CAS wave function (see Table II). When we compare with SHDMC optimized wave functions we find an excellent agreement in both VMC and DMC energies. Therefore, SHDMC improves the nodes systematically starting from the HF ground-state.

Since retaining all the determinants in the wave function would be costly, we performed calculations with different \( N_b \) to extrapolate (quadratically) the final energies as \( \sum_n (\lambda_n^{\text{MCSCF}})^2 \rightarrow 1 \) (see Fig. 2). The extrapolated DMC energies reached chemical accuracy (see also Table II).

Proof of principle in larger systems — Figures 1 and 2 show that SHDMC produces reliable and accurate results for small systems starting form the HF nodes. It is also important to demonstrate that SHDMC is a robust approach that can find the correct nodal surface topology of much larger systems even when starting from random nodal surfaces.

Figure 3 shows proof of principle results obtained for a \( \text{C}_{20} \) fullerene. These calculations used the branching SHDMC algorithm \((8)\) implemented by us in CASINO \((22)\). Two electrons were removed from the system to obtain a non-interacting DFT ground-state wave function invariant under any transformation belonging to the icosahedral group \((I_h)\) symmetry. The orbitals were obtained directly with the real space code PARSEC \((23)\) and classified according to their irreducible representations for \( I_h \) and its subgroup \( D_{2h} \). For this calculation 694 excitations (determinants) were sampled. No CI prefiltering of determinants is required; we only use the selection rules of both \( I_h \) and \( D_{2h} \) symmetries.

The \( \text{C}_{20}^+ \) system has a large DFT gap (5.53 eV) which is often associated with a dominant role of the non-interacting solution in the many-body wave function. The \( \lambda_0 \) coefficient is expected to dominate the final optimized trial wave function. All initial coefficients \( \lambda_n \) of \( \Psi_T(R) \) were set to random values, but for \( \lambda_0 \) which was set to zero. New \( \lambda_0 \) values were sampled with \( \sim 5094 \) walkers every 100 DMC steps. We found that when the quality of the wave function is poor, it is better (i) to update \( \lambda_n \) frequently (after only 4 samplings),
and (ii) to use the T-moves approximation \[24\] which limits persistent configurations. As the quality of the wave-function improved, we gradually increased the accumulation time (up to 96 samplings) and removed the T-moves approximation (which, in practice, hinders the final SHDMC convergence). Figure 3 shows that SHDMC can correct nodal errors as large as 0.5 Ha. The calculation was stopped when we obtained an energy of \(-112.487(2)\) Ha compared with the single determinant energy of \(-112.473(1)\) Ha. We have confidence that SHDMC can be applied to cases where the nodal structure of the ground-state is completely unknown since it is successful and converges to the expected result starting from random.

The SHDMC recursive runs required 220 hrs on 1024 processors (Cray XT4). This can be reduced to \(\sim 100\) hrs starting from the ground state determinant. Comparable EMVMC calculations with the same basis were unsuccessful, presumably due to the statistical errors in the Hessian and overlap matrices. The energy was not improved with EMVMC \((-112.488(3)\) Ha) even selecting a basis with the largest 104 coefficients of the 694 sampled in SHDMC. The estimated running time for EMVMC with CASINO 2.5 using \(N_b = 694\) and just 400 configurations \[24\] on 1024 processors is already \(\sim 100\) hrs, suggesting that for \(C_{20}\) SHDMC is faster than EMVMC. However, both methods can be improved for large \(N_b\) (e.g. as in Ref. \[26\]), by removing redundant IO etc.

**Summary** — We have shown that the SHDMC wave function converges to the ground-state of our best CI calculations and is systematically improved as the number of coefficients sampled increases and the statistics are improved. SHDMC presents equivalent accuracy to the EMVMC approach \[10, 11\] starting from random coefficients. SHDMC is numerically robust and can be automated.

The number of independent degrees of freedom of the nodes increases exponentially with the number of electrons. \[9\] Since EMVMC is based on VMC, the prefactor for its computational cost is much smaller than SHDMC. However, the number of quantities sampled in EMVMC is quadratic with respect to the number of degrees of freedom. In addition, EMVMC requires inverting a noisy matrix. These requirements cause EMVMC to scale at least quadratically. In contrast, SHDMC only requires one to sample a number of quantities linear in the number of optimized degrees of freedom. Therefore, a crossover between the methods is expected for systems of sufficient size or complexity. Tests on the large \(C_{20}\) fullerene system demonstrate that SHDMC is robust and that the nodes are systematically improved even starting from a random coefficients in the trial wave function. This shows that SHDMC can be used to find the nodes of unknown complex systems of unprecedented size.

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