Quantum Monte Carlo study of the three-dimensional spin-polarized homogeneous electron gas

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We have studied the spin-polarized three-dimensional homogeneous electron gas using the diffusion quantum Monte Carlo method, with trial wave functions including backflow and three-body correlations in the Jastrow factor, and we have used twist averaging to reduce finite-size effects. Calculations of the pair-correlation function, including the on-top pair density, as well as the structure factor and the total energy, are reported for systems of 118 electrons in the density range $r_s = 0.5$ –20 a.u., and for spin polarizations of 0, 0.34, 0.66, and 1. We consider the spin resolution of the pair-correlation function and structure factor, and the energy of spin polarization. We show that a control variate method can reduce the variance when twist averaging, and we have achieved higher accuracy and lower noise than earlier quantum Monte Carlo studies.

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I. INTRODUCTION

The simplicity of the three-dimensional (3D) homogeneous electron gas (HEG) model,¹ consisting of electrons interacting via the Coulomb potential in a uniform, positive background for charge neutrality, allows the study of important features of the many-electron problem without the complication of a lattice potential. Early pioneers of quantum mechanics discovered much of the phenomenology of the HEG, including Wigner's celebrated insight^{2,3} that the fluid will crystallize at low density. More recently, the continuing research effort to understand the behavior of the HEG is motivated by the model's application in density functional theory (DFT).

Quantum Monte Carlo (QMC) methods can provide accurate estimates of the static properties of interacting many-body systems such as the HEG. The ground state of the 3D HEG for collinear spins is accepted to be an unpolarized Fermi fluid at high densities. As the density decreases, QMC calculations indicate an apparently continuous transition to a spin-polarized fluid occurs, starting at a density of about $r_s = 50 \pm 2$ a.u.⁴ The transition to a Wigner crystal is calculated to occur at about $r_{\rm s} = 106 \pm 1$ a.u.⁵ QMC calculations have furnished data used within DFT, most notably the energies of the HEG calculated by Ceperley and Alder,³ which are used to parametrize the local-density-approximation exchange-correlation functional. Semilocal⁶ and nonlocal^{7,8} exchange-correlation functionals may use additional properties of the HEG such as the paircorrelation function (PCF) and, especially, the on-top pair density.9

The PCF is a measure of the spatial correlations of electron positions that arise from the Coulomb repulsion and Pauli exclusion. The spin-resolved PCF of the HEG, $g_{\alpha\beta}(r)$, is defined such that $n_{\beta}g_{\alpha\beta}(r)4\pi r^2 dr$ is the expected number of spin- β electrons in an infinitesimal shell of radius r when a spin- α electron is found at the origin and n_{β} is the number density of spin- β electrons. Defining the spin polarization ζ as the ratio $\zeta = (N_{\uparrow} - N_{\downarrow})/(N_{\uparrow} + N_{\downarrow})$, the total PCF g is a weighted average of the spin-resolved components:

$$g = \left(\frac{1+\zeta}{2}\right)^2 g_{\uparrow\uparrow} + \left(\frac{1-\zeta}{2}\right)^2 g_{\downarrow\downarrow} + \left(\frac{1-\zeta^2}{2}\right) g_{\uparrow\downarrow}.$$
 (1)

The on-top pair density g(0) is the value of the PCF at contact. The region around r = 0 is, however, the most challenging to sample in a stochastic simulation, particularly at low densities, as the electrons are rarely found close to one another. Toulouse *et al.*¹⁰ have developed a possible solution to this difficulty, extending the zero-variance zero-bias estimators of Assaraf and Caffarel.¹¹ It has recently been found by Fantoni,¹² however, that the zero-bias correction required in diffusion Monte Carlo (DMC) calculations significantly increases the variance. We therefore opted to use the traditional histogram estimator of the PCF in this work; our simulations gathered enough data to ensure good precision in the reported g(0) values.

In light of the above challenges, while QMC methods have been very successful in calculating many quantities of interest in the HEG, such as the energy and the PCF at intermediate distances, the short-range behavior of the PCF has usually been obtained by fitting QMC data to some analytical function, which can be constrained to obey exact results such as the electron-electron cusp conditions. Even then, however, there is some disagreement between the PCFs obtained using different approaches. Gori-Giorgi and Perdew developed an analytical model based on exact constraints and QMC energy data.^{13,14} The same authors¹⁵ extended an approach originally due to Overhauser¹⁶ using a screened Coulomb potential and two-electron wave functions. An alternative starting point is afforded by ladder theory, which gave rise to an early and widely used result for g(0) due to Yasuhara.¹⁷ Nagy *et al.*¹⁸ offer insight into the good agreement between these results. More recently, Qian¹⁹ was able to relax an approximation made in the earlier ladder theory calculations and obtained a markedly different result. Although several QMC calculations of PCFs for the unpolarized 3D HEG have been reported, less attention has been paid to fully and, especially, partially spin-polarized systems. 12,20-25

In this paper we present extensive results for PCFs, static structure factors (SSFs), and energies for HEGs containing 118 electrons over the density range relevant to DFT calculations, $r_s = 0.5$ –20 a.u. We consider four values of the spin polarization, $\zeta = 0$, 40/118, 78/118, and 1, although in the text we refer to the intermediate polarizations as 0.34

(40/118) and 0.66 (78/118). We use Slater-Jastrow wave functions incorporating backflow and three-body correlations, we investigate twist-averaged boundary conditions and finite-size effects, and we are able to gather sufficient data to ensure that the statistical uncertainty is relatively modest. A variance reduction method is found to improve significantly the precision of twist-averaged data without introducing bias. We show the spin resolution of the PCFs and the SSFs, and we consider, in particular, the on-top pair density and the energy of spin polarization as functions of r_s and ζ . We have performed spline fits to our PCF data, which appear to be more accurate than polynomial fits. A short FORTRAN 90 program is available to evaluate the spline fits to our PCF data.²⁶

Our paper is organized as follows. In Sec. II we discuss the QMC methods used in our calculations. Section III describes our PCF, SSF, and energy data and shows comparisons to results in the existing literature. We offer our conclusions in Sec. IV. Hartree atomic units are used throughout this paper, so that $\hbar = |e| = m_e = 4\pi\epsilon_0 = 1$.

II. METHODS

We have used the CASINO²⁷ code to perform variational and diffusion Monte Carlo²⁸ (VMC and DMC) calculations. In the VMC method, the Metropolis algorithm is used to generate a set of configurations distributed according to the square modulus of a trial wave function over which the local energy is averaged. In the DMC method, an initial wave function is evolved in imaginary time, which in principle projects out the ground state. For fermionic systems, the antisymmetry of the wave function is, however, imposed via the fixed-node approximation,²⁹ in which the nodal surface is constrained to remain unchanged during the evolution. Both the VMC and DMC methods give an upper bound to the ground-state energy of the system. The quality of the parametrization of the trial wave function and the VMC optimization procedure influence the accuracy of the results and the statistical noise.

Our wave functions consisted of Slater determinants of plane waves multiplied by a Jastrow factor and included a back-flow transformation. The Jastrow factor comprised polynomial and plane-wave expansions in the electron-electron separation, together with three-body terms.^{30,31} The electron coordinates in the Slater determinant were replaced by quasiparticle coordinates obtained by a backflow transformation represented by a polynomial in the electron-electron separation.^{32,33} The variable parameters in these wave functions were optimized using variance minimization^{34,35} and then linear least-squares energy minimization,³⁶ as this was found to give the most accurate trial wave functions.

The quality of our wave functions was such that, for unpolarized and fully polarized systems, the VMC and DMC methods produced essentially identical PCFs within the statistical precision we were able to obtain. For these systems we therefore report DMC expectation values rather than the extrapolated estimators often used in QMC studies. In partially polarized systems, however, our trial wave functions led to small but statistically significant differences between the VMC and DMC data for minority-spin electrons. In these cases we have used extrapolated estimation,³⁷ in which the PCF is

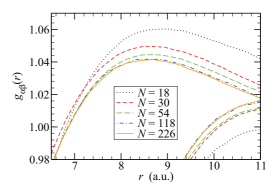


FIG. 1. (Color online) Finite-size effects in the spin-resolved PCF. The upper curves are for antiparallel spins and the lower-right curves are for parallel spins. The data are from VMC twist-averaged simulations at $r_s = 5$ a.u. and $\zeta = 0$, and at the various sizes shown. The region around the first peak in the antiparallel-spin PCF is shown, where the largest finite-size effects occur.

 $g = 2g_{\text{DMC}} - g_{\text{VMC}}$, where g_{DMC} and g_{VMC} are the DMC and VMC PCFs, respectively.

We studied 118-electron HEGs in face-centered-cubic simulation cells and imposed twisted boundary conditions³⁸ so that the wave function picks up a phase when an electron is translated by a simulation-cell lattice vector \mathbf{R}_s :

$$\Psi(\mathbf{r}_1,\ldots,\mathbf{r}_i+\mathbf{R}_s,\ldots,\mathbf{r}_N)=\exp(i\mathbf{k}_s\cdot\mathbf{R}_s)\Psi(\mathbf{r}_1,\ldots,\mathbf{r}_N).$$

The twist offset \mathbf{k}_s was allowed to vary with uniform probability over the first Brillouin zone of the simulation cell during our simulations. Averaging in this way has been shown to reduce single-particle finite-size effects in the energy.³⁹

We investigated the remaining variation with system size in the PCF using twist-averaged VMC simulations at $r_s = 5$ a.u. and $\zeta = 0$. We show the spin-resolved PCF in Fig. 1. This figure, like all others in this paper except Fig. 12, shows our raw QMC data, rather than a fit to the data. The region around the first peak in the antiparallel-spin PCF best shows the variation with system size. Both $g_{\uparrow\downarrow}(r)$ (upper curves) and $g_{\uparrow\uparrow}(r)$ (lower-right curves) exhibit systematic finite-size errors for small systems in this region, converging to the thermodynamic limit on the scale of the graph at approximately N = 100 electrons. (Results for 118- and 226-electron systems are almost indistinguishable on the scale of the graph.) Some cancellation of errors occurs when the spin-averaged PCF is calculated.

The variation with system size appears less pronounced and less systematic at smaller *r*. The on-top pair density was unaffected within statistical uncertainty. For systems with N = 54, 118, and 226 electrons, there was good agreement at small but nonzero distances: Typical differences were of order 1%–5%. In contrast, the long-range PCF is affected by unavoidable finite-size errors due to the finite size of the simulation cell. Twist averaging cannot remove the spurious correlation caused by the periodic boundary conditions. This shows up as a small but statistically significant variation in the long-range oscillations from about $r/r_s \gtrsim 2$ for 118 electrons that decreases in magnitude as the system size increases. Figure 1 is included as an XMGrace file in the Supplemental Material²⁶ accompanying this paper, so that other regions of the graph may be inspected. The energy is known to converge much more slowly with system size than the PCF.^{38,39} We have therefore applied analytical corrections to our energy data following the procedure developed in Refs. 39 and 40. Residual single-particle finite-size effects are removed by addition of the difference between the infinite-system and twist-averaged, finite-system Hartree-Fock kinetic energies. The leading-order correction to the Ewald energy, and leading- and next-to-leading-order corrections to the kinetic energy have previously²⁵ been found to be relatively insensitive to ζ , and so a cancellation of errors renders finite-size effects in the energies of spin polarization small even without any correction. The analytical arguments of Holzmann *et al.*⁴¹ lend further support to this conclusion.

We also examined possible time-step and populationcontrol biases and found them to be very small—less than one part in a thousand of the correlation energy in test cases—so we neglect them in what follows. All DMC calculations were performed using at least 1000 walkers and time steps for which the acceptance probability was generally greater than 99.7%.

We accumulate the spin-resolved PCF as an average over the set of configurations generated in the course of the simulation:

$$g_{\alpha\beta}(r) = \frac{\Omega}{4\pi r^2 N_{\alpha} N_{\beta}} \left\langle \sum_{\substack{j \in \alpha \\ l \neq j}} \sum_{\substack{l \in \beta \\ l \neq j}} \delta(|\mathbf{r}_j - \mathbf{r}_l| - r) \right\rangle,$$

where Ω is the simulation-cell volume and N_{α} is the number of electrons of spin α . The SSF is simply related to the Fourier transform of the PCF and is given by

$$S_{\alpha\beta}(\mathbf{k}) = \frac{1}{\sqrt{N_{\alpha}N_{\beta}}} \langle \rho_{\alpha}(\mathbf{k})\rho_{\beta}(-\mathbf{k})\rangle - \sqrt{N_{\alpha}N_{\beta}}\delta_{\mathbf{k}\mathbf{0}},$$

where $\rho_{\alpha}(\mathbf{k}) = \sum_{j \in \alpha} \exp(i\mathbf{k} \cdot \mathbf{r}_j)$. $S_{\alpha\beta}(\mathbf{k})$ is then spherically averaged in **k** space, because the SSF only depends on the magnitude of **k** in a homogeneous and isotropic system.

At high densities, the variation in the energy as the twist offset is changed can be much greater than other sources of noise in the simulation. Twist averaging can therefore be computationally expensive for high-density HEGs, because a large number of twist angles are required in order to obtain a precise energy and the simulation must be reequilibrated between twists, preventing rapid changes of twist offset in a DMC calculation. To circumvent this difficulty, we have used the Hartree-Fock energy (as a function of twist offset) as a control variate.⁴²

If E_{QMC} denotes the QMC energy, E_{HFKE} the Hartree-Fock kinetic energy, and E_{HFEX} the exchange energy as functions of twist angle, the standard way of getting the twist-averaged energy is $\langle E_{\text{QMC}} \rangle$, where the angular brackets denote an average over twists. Another estimator is

$$\theta = E_{\text{QMC}} + c_1 (E_{\text{HFKE}} - \langle E_{\text{HFKE}} \rangle) + c_2 (E_{\text{HFEX}} - \langle E_{\text{HFEX}} \rangle),$$

where the $\{c_i\}$ are coefficients that can be chosen to minimize the variance of θ . In order to account for the remaining sources of noise, we add the correction above to the energy data at each time step and, at the end of the simulation, reblock⁴³ the corrected data to obtain much smaller error bars. TABLE I. Energies from VMC simulations in a HEG at $r_s = 0.5$ a.u. containing two up-spin electrons and one down-spin electron. The raw VMC energies are compared to the same data after a correction has been applied using HF kinetic and exchange energies as described in the text. The VMC twist-averaged energy obtained in the standard way, using just over 21×10^6 twist angles, is 3.5484(3) a.u. per electron.

No. twists	VMC energy (a.u./elec.)		
	Raw	Corr.	
10	3.73(40)	3.5480(1)	
20	3.57(39)	3.5480(1)	
100	3.42(10)	3.5480(1)	
441	3.51(6)	3.5480(1)	

To illustrate the reduction in variance obtained, we apply the method to VMC data for two systems in this section: first a HEG with two up-spin electrons and one down-spin electron at $r_s = 0.5$ a.u.; then a 118-electron HEG at $r_s = 0.5$ a.u. and $\zeta =$ 0. The VMC method allows one to simulate a large number of twist angles in parallel, so we are able to compare the corrected data to accurate twist averages performed the standard way. It can be seen in Tables I and II that the method successfully reduces the variance without introducing bias. Although we chose twist offsets randomly throughout the first Brillouin zone, the method might also improve convergence of twist averages using Monkhorst-Pack⁴⁴ grids.

III. RESULTS

A. Pair-correlation function

Spin-resolved PCFs are shown in Fig. 2 for the unpolarized HEG at different densities using twist averaging. Antiparallelspin PCFs are translated upwards by 0.2 units for greater visibility, and show much greater variation with density. For $\zeta = 0$ (and $\zeta = 1$, not shown) the VMC and DMC results are very similar, so in the plot we present only DMC data.

We show the variation of the PCFs with spin polarization at $r_s = 3$ a.u. in Fig. 3. In this figure, we translate the antiparallelspin PCFs upwards by 0.4 units and the parallel-majority-spin PCFs upwards by 0.2 units. For $\zeta = 0$ and $\zeta = 1$, we have used the DMC values. For $\zeta = 0.34$ and $\zeta = 0.66$, the extrapolated estimate is shown, because small but statistically significant

TABLE II. Energies from VMC simulations in a 118-electron HEG at $r_s = 0.5$ a.u. and $\zeta = 0$. The raw VMC energies are compared to the same data after a correction has been applied using HF kinetic and exchange energies as described in the text. The VMC twist-averaged energy obtained in the standard way, using 4.8×10^6 twist angles, is 3.41378(2) a.u. per electron.

	VMC energy (a.u./elec.)		
No. twists	Raw	Corr.	
10	3.424(4)	3.41377(4)	
20	3.424(4)	3.41375(4)	
50	3.414(5)	3.41380(3)	
156	3.416(2)	3.41377(2)	

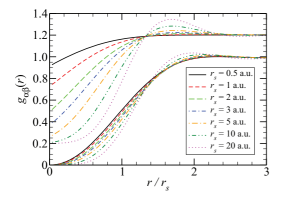


FIG. 2. (Color online) Spin-resolved PCFs for unpolarized HEGs calculated at the densities shown. The antiparallel-spin PCFs are translated upwards by 0.2 units. The data were obtained using twist-averaged DMC simulations.

differences emerged between the DMC and VMC data in the parallel-minority-spin PCFs. Compared with the parallel-spin PCFs, the antiparallel-spin PCF is relatively insensitive to the degree of spin polarization.

The PCF data discussed above are twist averaged to reduce single-particle finite-size effects. Twist averaging has a significant effect on the energy data, but the effect on the PCF data is much smaller. We show in Fig. 4 the difference $g_{TA} - g_{PBC}$ between imposing twist-averaged and periodic boundary conditions for parallel spins. Noise obscures any clear pattern for the antiparallel spins, although there is a tendency for the equivalent figure to show a small dip before rising to zero, so that g(0) appears to be largely unaffected.

We compare our results at $r_s = 5$ a.u. and $\zeta = 0$ with QMC data from the literature in Fig. 5. We find good agreement with the data of Gori-Giorgi *et al.*²³ for intermediate r/r_s . Our on-top pair densities are closer to those of Ortiz and Ballone²⁰ as shown in the inset, although at intermediate r/r_s modest (of order 5%) differences are obtained. The figure shows our raw

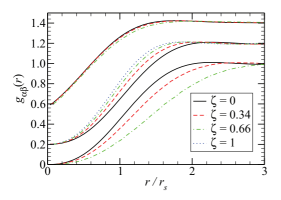
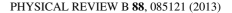


FIG. 3. (Color online) Variation in the PCF with spin polarization at $r_s = 3$ a.u. Antiparallel-spin PCFs are translated upwards by 0.4 units, parallel-majority-spin PCFs are translated upwards by 0.2 units, and the lower curves give the parallel-minority-spin PCFs. Note that the $\zeta = 0$ parallel-spin PCF is shown twice (in black) for comparison with both up-up and down-down spin PCFs in those cases where they differ. The DMC data are shown for unpolarized and fully polarized systems, while extrapolated estimates are used for intermediate polarizations. All of the data are twist averaged.



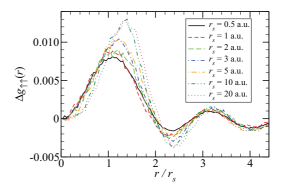


FIG. 4. (Color online) Changes in the PCF due to twist averaging. We plot parallel-spin PCFs with twist averaging minus the same quantity evaluated without twist averaging, for the densities shown and $\zeta = 0$. Only DMC results are shown, as the VMC results are very similar.

QMC data, whereas the other curves are fits to the QMC data of Ortiz, Harris, and Ballone,²² and Ortiz and Ballone²⁰ respectively. Both of these fits obey the Kimball cusp conditions and we are confident our QMC data are also consistent with these conditions. As mentioned below, our short-range PCFs are well described by a quadratic of the form $g_{\uparrow\downarrow}(r) = a + ar + br^2$, which satisfies the first Kimball condition.

We show in Fig. 6 our g(0) values for the unpolarized system in comparison to those in the literature. We have averaged our DMC and VMC data in this figure to reduce statistical uncertainty, because the $\zeta = 0$ PCFs produced by the two methods are identical within the statistical precision we were able to obtain. We used the above quadratic form to estimate g(0) from our PCF data at finite distances, g(r), and obtained good fits for all systems. Our results are in good agreement with the recent QMC data of Holzmann *et al.*,²⁴ and the results of Gori-Giorgi and Perdew¹³ who used various data including QMC results, and the results of Yasuhara.¹⁷ Our calculations give significantly smaller values of g(0) than those of Gori-Giorgi, Sacchetti, and Bachelet,²³ and those of Qian.¹⁹ This may suggest that Yasuhara's scheme successfully incorporates

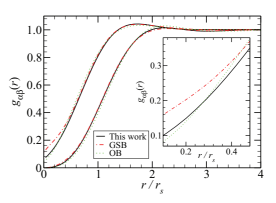


FIG. 5. (Color online) Spin-resolved PCFs for unpolarized HEGs at $r_s = 5$ a.u. We show our raw twist-averaged DMC data, together with the fit to DMC results from Ortiz and Ballone (Ref. 20) (labeled OB) given in their paper, and a fit due to Gori-Giorgi, Sacchetti, and Bachelet (Ref. 23) using the DMC data of Ortiz, Harris, and Ballone (Ref. 22) (labeled GSB). The inset shows the low-*r* region, where the differences are largest.

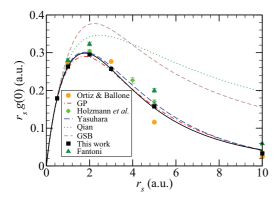


FIG. 6. (Color online) The on-top pair density g(0) (multiplied by r_s) as a function of r_s for unpolarized systems. The data shown are those of our averaged DMC and VMC results, together with the fit of Eq. (2) (solid line); Gori-Giorgi and Perdew (Ref. 13); Holzmann *et al.* (Ref. 24); Yasuhara (Ref. 17); Qian (Ref. 19); Gori-Giorgi *et al.* (Ref. 23); Ortiz and Ballone (Ref. 20); and Fantoni (Ref. 12). Error bars are shown for our data, those of Holzmann *et al.* (Ref. 24), and Fantoni (Ref. 12), but they are mostly smaller than the size of the symbols. Our values are slightly smaller than those of Holzmann *et al.* (Ref. 24).

screening effects into the ladder theory calculations. Nagy $et al.^{18}$ suggest that the description of screening used by Gori-Giorgi and Perdew is also an important factor in explaining the success of the latter authors' model.

We have also obtained on-top pair densities for partially polarized systems (see Fig. 7). In this case we used extrapolated estimation due to the small but statistically significant differences between our VMC and DMC results, although these differences are much less pronounced in the antiparallel-spin PCF than in the parallel-minority-spin PCF. This increases the statistical uncertainty in our results, an effect compounded by the increasing sparsity of sampling antiparallel spin coalescences as the polarization increases. Nevertheless, as expected from Fig. 3, the variation of g(0)with ζ arises largely from changes in the weights in Eq. (1)

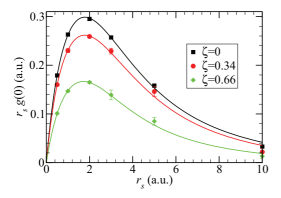


FIG. 7. (Color online) The on-top pair density (multiplied by r_s) as a function of r_s and ζ . Averaged DMC and VMC data are used for the unpolarized systems, and extrapolated estimates for the polarized systems. Twist averaging was used for all data shown. An error bar is plotted for each data point, although some of them are smaller than the size of the symbols. The partially polarized systems show a larger statistical uncertainty. The lines are fits as discussed in the text.

TABLE III. Parameters (in a.u.) obtained from fits to the ontop pair density as a function of r_s using Eq. (2). The data for the unpolarized systems are averaged DMC and VMC results; extrapolated estimation is used for the others.

ζ	а	b	С	d
0	0.18315	-0.0784043	1.02232	0.0837741
0.34	0.284118	-0.110062	1.1618	0.0874753
0.66	0.0659538	-0.0590569	0.836458	0.0832258

rather than in the spin-resolved PCFs themselves. We fit our g(0) data to the following parametrized form in the density range $0.5 \le r_s \le 20$ a.u.:

$$g_{\uparrow\downarrow}(0;r_{\rm s}) = \frac{1 + a\sqrt{r_{\rm s}} + br_{\rm s}}{1 + cr_{\rm s} + dr_{\rm s}^3},\tag{2}$$

and we list the optimal fit parameters obtained in Table III. This functional form obeys the exact high-density limit and fits our data well up to $r_s = 20$ a.u., although at $r_s = 10$ a.u., it tends to give slightly higher values than our QMC calculations obtained, typically by about two standard deviations.

B. Static structure factor

SSFs with $\zeta = 0$ at three representative densities are shown in Fig. 8. The upper curves show $S_{\uparrow\uparrow} - S_{\uparrow\downarrow}$ and the lower curves, $S_{\uparrow\uparrow} + S_{\uparrow\downarrow}$. The differences between the DMC and VMC results are again smaller than the statistical noise, and so we plot DMC values. The parallel-spin structure factors change very slowly with density, and the r_s dependence shown arises almost entirely from the antiparallel-spin structure factor, which is (mostly) negative and of increasing magnitude as the density decreases.

Twist averaging was used to obtain the results shown in Fig. 8, and its effect was more pronounced than in the PCFs, as shown in Fig. 9, where we plot antiparallel-spin SSFs at $\zeta = 0$ with and without twist averaging for the same three densities. Shell-filling effects are clearly visible in the

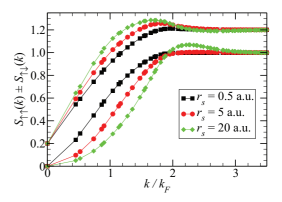


FIG. 8. (Color online) Structure factors at $\zeta = 0$ for three densities. The upper curves show $S_{\uparrow\uparrow} - S_{\uparrow\downarrow}$ and are translated upwards by 0.2 units; the lower curves show $S_{\uparrow\uparrow} + S_{\uparrow\downarrow}$. The lines are visual guides only. The statistical uncertainty is smaller than the size of the symbols. The DMC method with twist averaging was used, as the VMC data are again very similar. Intermediate densities lie between the curves shown.

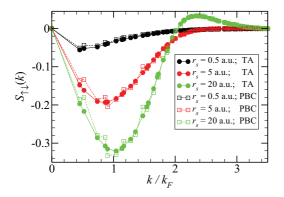


FIG. 9. (Color online) Effect of twist averaging on the antiparallel-spin SSF. We plot SSFs at $\zeta = 0$ for three densities, with twist averaging (TA) and without (PBC). Higher densities are towards the top of the graph; intermediate densities lie between the data shown and similar effects are seen in parallel-spin SSFs. Only DMC data are plotted as the VMC data are very similar.

non-twist-averaged data. The VMC and DMC results are very similar and so we can be confident that the differences do not arise from statistical noise. Small finite-size effects are visible in the low- $|\mathbf{k}|$ structure factor data. These correspond to the inevitable finite-size effects in the long-range part of the PCF mentioned above.

C. Energies

Twist-averaged DMC energies are reported in Table IV for $r_s = 0.5-20$ a.u. and for $\zeta = 0$, 0.34, 0.66, and 1. The data are for finite (small) time steps such that the DMC acceptance probability was almost always greater than 99.7%. We verified in sample cases that the time-step and population-control biases were small: typically less than one part in a thousand of the correlation energy. The finite-size effects in the energy are substantial for the 118-electron HEGs, so we have corrected for these, as discussed in Sec. II. The data are further corrected using the control variate method, also discussed in Sec. II, because twist averaging would otherwise lead to large error bars, especially at high densities.

We find that the ground state is unpolarized for all densities studied here, in agreement with the most recent calculations.³⁸ This is also clearly evident in our VMC results, despite the VMC method suffering a bias towards polarized systems

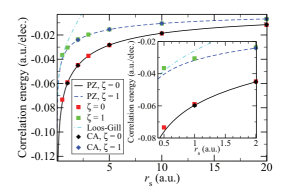


FIG. 10. (Color online) Correlation energy at $\zeta = 0$ and 1 against density parameter r_s . The present work is simply labeled " $\zeta = 0$ " and " $\zeta = 1$." Also shown are the Ceperley-Alder QMC data (Ref. 3); the Perdew-Zunger fit to the latter data (Ref. 45); and the recent high-density result of Loos and Gill (Ref. 46). Our data are corrected for finite-size effects. Time-step extrapolation was not performed. The statistical error bars are shown, but are usually smaller than the symbols.

due to the relative ease with which correlation effects in the wave function may be parametrized. This bias decreases with increased variational freedom in the trial wave function.

Our energies for unpolarized and fully polarized HEGs are shown in Fig. 10, where we also plot for comparison the Ceperley-Alder data³ and the fit to the latter obtained by Perdew and Zunger,⁴⁵ as used in the LSDA. In addition, a recent high-density (RPA) limit for the polarized system due to Loos and Gill⁴⁶ is shown. The present results follow the Perdew-Zunger fit closely, except at high density, where we find slightly smaller correlation energies, especially at full polarization. The finite-size corrections we have applied are significantly larger than the differences shown in Fig. 10, and it is possible that higher order finite-size corrections would account for the difference in energies obtained. It is worth noting that Ceperley and Alder did not perform QMC calculations at densities higher than $r_s = 2$ a.u. for $\zeta = 1$ and that the Loos-Gill result also gives smaller correlation energies in this region than the Perdew-Zunger fit.

To compare our intermediate- ζ energies with those in the literature, we used the procedure developed by Perdew and Zunger to interpolate between $\zeta = 0$ and $\zeta = 1$. We used our QMC data for the unpolarized and fully polarized systems

TABLE IV. Energies as a function of density parameter and spin polarization. The DMC data are twist averaged and include corrections for finite-size effects, as discussed in the text. Time-step extrapolation was not performed. Data include a control variate correction.

	Energy (a.u./elec.)			
<i>r</i> _s (a.u.)	$\zeta = 0$	$\zeta = 0.34$	$\zeta = 0.66$	$\zeta = 1$
0.5	3.43011(4)	3.69287(6)	4.44164(6)	5.82498(2)
1	0.58780(1)	0.64919(2)	0.82394(4)	1.14634(2)
2	0.002380(5)	0.016027(6)	0.05475(2)	0.12629(3)
3	-0.067075(4)	-0.061604(5)	-0.04608(2)	-0.017278(4)
5	-0.075881(1)	-0.074208(4)	-0.069548(4)	-0.060717(5)
10	-0.0535116(5)	-0.053214(2)	-0.052375(2)	-0.0507337(5)
20	-0.0317686(5)	-0.0317156(7)	-0.0315940(7)	-0.0313160(4

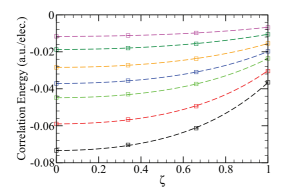


FIG. 11. (Color online) Correlation energies obtained in DMC calculations together with the fit of Eqs. (3), (4), and (5). Dashed lines show the fit; square symbols the QMC data. Lower densities appear towards the top of the graph; higher densities towards the bottom of the figure. From top-to-bottom, therefore, the densities shown are: $r_s = 20$, 10, 5, 3, 2, 1, and 0.5 a.u. Error bars on the QMC data are shown, but are smaller than the size of the symbols. Corrections have been applied for finite-size effects. Time-step extrapolation was not performed.

and interpolated using Eq. (C12) in their paper. These fits are included as part of the Supplemental Material accompanying this paper.²⁶ The interpolation scheme is very successful at low density, whereas our high-density data for intermediate polarizations are not reproduced with such high accuracy.

In Fig. 11 we show our correlation energies together with a quartic fit in ζ at each density:

$$E_c(r_s,\zeta) = f_0(r_s) + \Xi(r_s)\Delta f(r_s)\zeta^2 + [1 - \Xi(r_s)]\Delta f(r_s)\zeta^4,$$
(3)

where $\Delta f(r_s) = f_1(r_s) - f_0(r_s)$ and $f_{\zeta}(r_s)$ denotes the correlation energy at spin polarization ζ . We fit $f_{\zeta}(r_s)$ for $\zeta = 0$ and 1 over the density range $r_s = 0.5$ –20 a.u. using the functional form

$$f_i(r_s) = \frac{\gamma_i}{1 + \beta_1^i r_s^{\frac{1}{2}} + \beta_2^i r_s}$$
(4)

suggested by Ceperley,⁴⁷ and Perdew and Zunger.⁴⁵ We find the interpolation

$$\Xi(r_{\rm s}) = a + br_{\rm s} + cr_{\rm s}^2 \tag{5}$$

gives a good fit to the intermediate- ζ QMC data. The energies can be extrapolated to the high-density limit using the method of Perdew and Zunger,⁴⁵ by matching the correlation energy and potential arising from f_i to the high-density expansions at $r_s = 0.5$ a.u., although it is not clear if the interpolation between unpolarized and polarized systems used here will be reliable at higher densities.

All data shown in Fig. 11 have been corrected for finite-size effects, as discussed above. A control variate correction has also been applied to reduce the statistical uncertainty arising from the finite sample of twist angles, as discussed in Sec. II. The optimum parameter values used in the fit of Eqs. (3), (4), and (5) are given in Table V.

TABLE V. Parameters (in a.u.) obtained from the fit to the correlation energy as a function of r_s and ζ using Eq. (3).

Parameter	Value
а	0.575073
b	0.0383567
С	-0.00144917
γ_0	-0.138971
γ_1	-0.0633399
β_1^0	1.04452
$\beta_2^{\dot{0}}$	0.311702
$\beta_1^{\overline{1}}$	0.872563
β_2^1	0.225783

D. Fits to PCFs

We have performed cubic-spline fits to our PCF data. The spin resolution across the density range $r_s = 0.5-20$ a.u. and spin polarizations $\zeta = 0, 0.34, 0.66, \text{ and } 1$ was well represented by cubic-spline fits with ten knots for each spin-resolved PCF. The boundary conditions used were those of natural splines: second derivatives set to zero at the first and last data points. An example of the fits obtained is given in Fig. 12, where we plot raw QMC data together with fits to the data. The splines reproduce our estimate for the on-top pair density at r = 0, and are accurate for $0 \le r/r_s \le 3$. For unpolarized and fully polarized systems, we fit the DMC data, whereas for partially polarized systems, we provide fits to the extrapolated estimates of the PCFs. Note that we used down spins as the majority spins in our simulations. We did not impose any exact results as constraints when performing the spline fits, in order to give the most accurate fit possible to our raw data. As discussed above, our OMC data are consistent with the Kimball cusp conditions, and the spline fits follow the data closely in all systems studied. On the other hand, our PCF data inevitably suffer finite-size effects at large r, so various sum rules requiring integration of the PCF over all space will not be satisfied exactly.

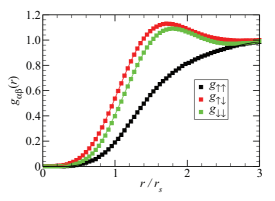


FIG. 12. (Color online) Cubic-spline fits to our twist-averaged extrapolated pair-correlation function data at $r_s = 20$ a.u. and $\zeta = 0.66$. Ten knots were used in the spline fits. Square symbols represent the raw QMC data points and lines represent our spline fits to those data. In all other figures, we have shown the raw QMC data rather than the spline fits displayed here.

These spline fits are available in the form of a small FORTRAN 90 program that we have written and which is included as part of the Supplemental Material accompanying this article.²⁶ Subroutines from the SLATEC Common Mathematical Library⁴⁸ were used to write this program. SSFs can be obtained from these data by Fourier transformation.

We also compared fitting schemes previously used to represent PCF data for these systems, including those proposed by Ortiz and Ballone,²⁰ and Gori-Giorgi, Sacchetti, and Bachelet.²³ The former scheme struggled to describe the region around the peak in the PCF with quantitative accuracy. The latter scheme was successful at high density (for $\zeta = 0$), but less reliable otherwise.²³

IV. CONCLUSIONS

We have investigated ground-state properties of the 3D HEG using the VMC and DMC methods. Our simulations cover the density range $r_s = 0.5-20$ a.u. and spin polarizations $\zeta = 0, 0.34, 0.66, \text{ and } 1$. We used highly accurate wave functions incorporating backflow and three-body correlations. Twist averaging was used to reduce finite-size effects, and we show that this can be performed efficiently using a control variate method.

The spin resolution of the PCF and SSF are reported at each spin polarization studied. At $\zeta = 0$, we obtain good agreement with previous QMC studies, except for $r/r_s \leq 0.5$, where our data are significantly smaller than those of Gori-Giorgi *et al.*²³ The effects of twist averaging on these quantities are shown. We report the r_s and ζ dependence of the energies for 118-electron HEGs. It is hoped that the higher accuracy and lower noise than earlier QMC studies that we have achieved, particularly of the PCFs, will be valuable in, for example, aiding the construction of nonlocal density functionals. A small FORTRAN 90 program is available to reproduce spline fits to our PCF and SSF data.

Note added in proof. Recently, we became aware of complementary, finite-temperature results for the warm-dense HEG in the paper by E. W. Brown *et al.*⁴⁹

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