Temperature-dependent behavior of confined many-electron systems in the Hartree-Fock approximation

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Many-electron systems confined at substantial finite temperatures and densities present a major challenge to density functional theory. In particular, there is comparatively little systematic knowledge about the behavior of free-energy density functionals for temperatures and pressures of interest, for example, in the study of warm dense matter (WDM). As with ground-state functionals, development of approximate free-energy functionals is faced with significant needs for reliable assessment and calibration data. Here we address, in part, this need for detailed results on well-characterized systems. We present results on a comparatively simple, well-defined, computationally feasible but previously unexplored model, the thermal Hartree-Fock approximation. We discuss the main technical tasks (defining a suitable basis and evaluation of the required matrix elements) and give an illustrative initial application that probes both the content of the model and the solution techniques: a system of eight one-electron atoms with nuclei at fixed, arbitrary positions in a hard-walled box. Even this simple system produces physical behavior different from that produced by simple ground-state density functionals used at finite temperature (a common approximation in the study of WDM).

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I. INTRODUCTION AND MOTIVATION

Warm dense matter (WDM) is encountered in systems as diverse as the interiors of giant planets^{1,2} and in the pathway to inertial confinement fusion.^{3,4} WDM is challenging to theory and simulation because it occurs inconveniently, for theory, between the comparatively well studied plasma and condensed-matter regimes. Both the Coulomb coupling parameter $\Gamma := Q^2/(r_s k_B T)$ and electron-degeneracy parameter $\Theta := k_B T/\varepsilon_F$ are approximately unity for WDM. (Q is relevant charge, r_s is Wigner radius, ε_F is electron Fermi energy, T is temperature, and k_B is the Boltzmann constant.) A nonperturbative treatment therefore is required.

Contemporary computations on WDM^{5–18} are dominated by the use of the Kohn-Sham (KS) realization of thermal density functional theory (DFT)^{19–25} to generate a potential surface for ionic motion (treated classically). The majority of such calculations use approximate ground-state exchange-correlation (XC) functionals E_{xc} with the temperature dependence of the XC free energy picked up implicitly from the T dependence of the density $n(\mathbf{r},T)$. Though fruitful, this approach is not without potential difficulties, as is illustrated in Fig. 3 of Ref. 7. Three issues are germane here.

First, there is little systematic knowledge of the implicit T dependence of ground-state approximate E_{xc} functionals (especially beyond the local density approximation, LDA), whether they are constraint based or empirical. Compared to the ground-state situation, there are relatively few studies on explicitly T-dependent functionals, that is, XC free-energy functionals, and essentially, all of those studies are at the level of the LDA. $^{26-42}$

Second, there is the computational burden of solving for the Kohn-Sham orbitals and eigenvalues. Since the computational load from the eigenvalue problem scales, in general, as order N_{orbitals}^3 , the growth in the number of non-negligibly occupied KS orbitals with increasing temperature is a clear computational bottleneck. See the remarks, for example, in

Sec. 4 of Ref. 18. For complicated systems, the same bottleneck is encountered in ground-state simulations that use the DFT Born-Oppenheimer energy surface to drive the ionic dynamics. One result has been the emergence of active research on orbital-free DFT (OFDFT), that is, approximate functionals for the ingredients of the KS free energy, namely, the KS kinetic energy (KE) T_s , entropy S_s , and XC free energy \mathcal{F}_{xc} or their ground-state counterparts. Almost all of this effort has been for ground-state OFKE functionals. (Note that most of the OFKE literature invokes the KS separation of the KE in order to use existing E_{xc} approximations consistently.)

Third, the finite-temperature OFDFT work is dominated by variants on Thomas–Fermi–von Weizsäcker (TFvW) theory (see, for example, Ref. 41 and references therein). That type of theory, however, is known (on both fundamental and computational grounds) to be no more than qualitatively accurate in many circumstances relevant to WDM (e.g., chemical binding). Compared to the data-rich context for development of zero-temperature functionals, there is little to guide development and assessment of finite-T functionals beyond TFvW. Similarly, compared to the T=0 K situation (or the very high T situation), not much is known about the accuracy of approximate finite-T OFDFT functionals beyond TFvW.

The primary aim of the present study is to provide reference data for both development and appraisal of free-energy density functionals. Such reference data must come from a combination of a well-defined physical system with a well-defined approximation and its implementation. In the case of ground-state density functionals, three classes of reference data have been particularly influential: high-precision Monte Carlo results for the homogeneous electron gas⁴⁷ and small systems, ⁴⁸ Hartree-Fock (HF) calculations on myriad molecules (the literature is too vast to cite in detail, but see the textbooks by Jensen⁴⁹ and by Szabo and Ostlund⁵⁰), and high-precision (e.g., coupled cluster and

configuration interaction) calculations on atoms and small molecules (again, the literature is too large to cite fairly, but see, for one example, Ref. 51). Though the Hartree-Fock approximation has accuracy limitations with respect to the precision of thermochemistry (e.g., atomization energies), it has had at least two large impacts on the development of ground-state DFT. First, it provides the framework for all basic electronic structure concepts. Second, because of the many rigorous results known about the HF approximation (see, for only two examples, Refs. 52 and 53), understanding the distinction between HF and Kohn-Sham DFT has sharpened the understanding of DFT itself.

There are multiple reasons, therefore, that finite-temperature Hartree-Fock^{23,54–56} results will be valuable in constructing and appraising free-energy functionals. Note that we do not propose finite-temperature Hartree-Fock studies as competitors or replacements for DFT-based molecular dynamics simulations of WDM. Though we intend, later, to undertake a small number of HF-based molecular dynamics (MD) simulations, the predominance of DFT-based MD at zero temperature makes clear that HF-based studies cannot be truly competitive. This, in itself, is another way of focusing on finite-temperature HF for development of better functionals.

II. SYSTEM AND METHODOLOGY

We treat a neutral system of N_{ion} atoms with nuclei fixed at arbitrary positions in a hard-walled three-dimensional rectangular box. The confined system allows systematic treatment of pressure effects at stipulated finite temperature and hence is a small, treatable sample of WDM. For specificity and comparative simplicity in this first-stage study, we chose H atoms and $N_{\rm ion} \leq 8$. Other than commensurateness with cubic symmetry, there is nothing special about this value. Note the corresponding use of a small number of atoms in Ref. 18. Also note the considerable literature on spherically confined systems at T = 0 K.^{57–62} Other than DFT calculations with ground-state functionals⁶³ (which, though interesting, are not relevant to our goal of providing calibration and assessment data for development of better functionals), we have not found any work on lower-symmetry confinement of multiatom systems at nonzero T. At nonzero T there is a large amount of literature on average-atom methods (for example, Refs. 64 and 18 and many others), but such methods also do not provide the fiduciary data needed for functional development.

Many-electron problems require clearly defined approximations. We choose the finite-temperature Hartree-Fock (FTHF) scheme, $^{23,54-56}$ with issues of electron correlation to be addressed in the future. In addition to being well defined in the grand ensemble, FTHF provides the advantage that its T=0 K limit is, as noted above, the *lingua franca* of molecular electronic structure interpretation. Use of FTHF therefore also provides a semiquantitative (at least) framework for understanding chemical processes in WDM.

The FTHF approximation is defined in the grand-canonical ensemble by restricting the relevant traces to states that are single Slater determinants. 23,54,55 The result is an upper bound to the free energy $\mathcal{F}_{\text{FTHF}} \geqslant \mathcal{F}$. Standard thermodynamic

relationships for the grand ensemble follow. The FTHF Euler equations to be solved (in unrestricted form) are⁵⁵

$$\varepsilon_{i}\varphi_{i}(\mathbf{r}) = \left(-\frac{1}{2}\nabla^{2} + U_{\text{ion}}(\mathbf{r})\right)\varphi_{i}(\mathbf{r})$$

$$+ \sum_{j} f_{j} \int d\mathbf{r}' \frac{|\varphi_{j}(\mathbf{r}')|^{2}}{|\mathbf{r} - \mathbf{r}'|}\varphi_{i}(\mathbf{r})$$

$$- \sum_{i} \delta_{\sigma_{i}\sigma_{j}} f_{j} \int d\mathbf{r}' \frac{\varphi_{j}^{*}(\mathbf{r}')\varphi_{i}(\mathbf{r}')}{|\mathbf{r} - \mathbf{r}'|}\varphi_{j}(\mathbf{r}), \quad (1)$$

where $U_{\rm ion}$ is the ion-electron interaction potential and σ_i is the spin label; the sums are over all spin orbitals. Unless indicated otherwise, we use hartree atomic units ($\hbar=m_e=e=1$; energy is then in hartrees, 1 hartree = 27.2116 eV, and lengths are in bohrs, 1 bohr = 0.52918 angstrom). The spin orbitals (eigenstates φ_i) have Fermi-Dirac thermal occupations

$$f_i = (1 + e^{\beta(\varepsilon_i - \mu)})^{-1}, \quad N = \sum_i f_i,$$
 (2)

where $\beta = 1/k_BT$ and μ is the electron chemical potential. For a specified value of N, which is a grand-ensemble average, μ must be determined. These equations, along with specification of the nuclear sites and imposition of hard-wall boundary conditions, completely describe the problem.

The FTHF free energy and entropy are given by

$$\mathcal{F}_{\text{FTHF}} = \sum_{i} f_{i} \varepsilon_{i} - \frac{1}{2} \sum_{i,j} f_{i} f_{j} (J_{i,j} - K_{i,j}) - T \mathcal{S}_{\text{FTHF}},$$
(3)

$$S_{\text{FTHF}} = -k_B \sum_{i} f_i \ln(f_i) + (1 - f_i) \ln(1 - f_i), \quad (4)$$

with conventional definitions of J and K:

$$J_{i,j} := \int d\mathbf{r} d\mathbf{r}' \frac{|\varphi_i(\mathbf{r})|^2 |\varphi_j(\mathbf{r}')|^2}{|\mathbf{r} - \mathbf{r}'|}, \tag{5}$$

$$K_{i,j} := \delta_{\sigma_i \sigma_j} \int d\mathbf{r} d\mathbf{r}' \frac{\varphi_i(\mathbf{r}) \varphi_j(\mathbf{r}) \varphi_i^*(\mathbf{r}') \varphi_j^*(\mathbf{r}')}{|\mathbf{r} - \mathbf{r}'|}.$$
 (6)

Equations (3) and (4) clearly reduce to conventional groundstate Hartree-Fock expressions in the zero-temperature limit.

A. Basis set design

Solution of self-consistent field equations such as (1) via Gaussian-type-orbital (GTO) basis methods is the standard procedure in modern computational codes for molecules. First introduced by Boys, 65–69 such basis sets automatically satisfy the free-molecule boundary condition that the orbitals vanish at infinity. For a hard-wall confined system, the basis functions must vanish at the boundary, so standard molecular GTO matrix-element expressions are inapplicable. This simple distinction underlies the most critical implementation issue, namely, to find a basis that satisfies the boundary conditions yet allows for an efficient enough evaluation of the two-electron integrals to be computationally tractable on reasonable resources. A second, closely related, technical issue is that the high temperature also dictates what is "efficient enough," in that the basis must be large enough and flexible

enough to represent a sufficient number of thermally occupied higher-energy orbitals of the system, hence to represent the density and free energy accurately.

Those considerations eliminate several seemingly plausible options for a basis. For example, a real space finite difference or finite element scheme, while suitable for a DFT calculation or a Hartree-Fock calculation on a free diatomic molecule (for which curvilinear coordinates can be exploited 70), is far too expensive for the present case because of the number of matrix elements to be calculated. Another example is sine functions. They also satisfy the hard-wall boundary condition, but an adequate description of the rapidly varying electronic distribution near the nuclei requires prohibitively many matrix elements in our multicenter problems. So we chose an adaptation of standard GTO methods, which uses modified Gaussians that meet the boundary condition yet retain enough efficiency to complete the calculation.

The various ways to force a GTO to zero at the bounding planes of a rectangular box can have great impact upon the efficiency of the matrix-element calculations. Compared with the familiar practice for free molecules, in general, the confined case requires more primitive GTOs for each contracted one. More importantly, the finite integration volume makes it impossible to achieve completely analytic calculation of the two-electron integrals, which is, of course, precisely the category in which computational efficiency is most needed. We have addressed this issue by using truncated Gaussians, as described next.

B. Truncated Gaussians

The rectangular box makes Cartesian GTOs a convenient choice because each primitive function then is separable into Cartesian factors, which are simple one-dimensional (1D) functions. Consider the Cartesian factor

$$g^{n}(x) = (x - x_{c})^{n} e^{-\alpha(x - x_{c})^{2}}.$$
 (7)

To force this function to zero at the box boundaries x = 0 and $x = L_x$, we subtract a constant equal to the function value at each end. When x_c is not at the box center, the value to be subtracted differs for the two ends, so we split the function into two pieces, make the two subtractions, and scale the two pieces such that the resulting function is continuous. Each unnormalized Cartesian factor becomes

$$g_{\text{box}}^{n}(x) = a_0[g^{n}(x) - \Delta_0] \quad 0 \leqslant x \leqslant x_c,$$

$$= a_L[g^{n}(x) - \Delta_{L_x}] \quad x_c \leqslant x \leqslant L_x,$$
(8)

with $\Delta_0 = g^n(0)$, $\Delta_{L_x} = g^n(L_x)$. We call this the truncated GTO (TGTO) basis.

Two technical issues remain. The TGTO functions may not have continuous derivatives, so proper evaluation of the kinetic-energy matrix elements requires attention. Appendix A shows that nothing untoward happens and that the kinetic energy is simply a sum of piecewise contributions, except for *p*-type functions, which have a simple correction term. Second is the matter of evaluating two-electron matrix elements. In Appendix B, we show that this task reduces to computing finite-range integrals of products of Gaussians and error functions. At this juncture, we are doing those via Gauss-Legendre quadrature. We also note that the TGTO basis is simpler than the smoothly cutoff floating spherical GTO basis of Ref. 61 in the sense that the latter introduces a mixing of two symmetry types (e.g., *s* and *d*) in each basis function.

Note that, so far, we have implemented only the restricted Hartree-Fock approximation (RHF; non-spin polarized in DFT language and closed shell or spin compensated in quantum chemistry language).

III. RESULTS

A. Zero temperature

Two simple ground-state test cases, the H atom and the H₂ molecule, illustrate the system behavior with increasing confinement (decreasing volume) as well as the correctness of

TABLE I. Optimized exponents and orbital energies for an H atom at the center of a cube of edge length L. 2p refers to the triply degenerate p_x , p_y , and p_z states. For L < 2, the 2s level could not be optimized beyond the first six exponents.

L	Fixed	1 <i>s</i>	2s	2p	ε_{1s}	$arepsilon_{2s}$	$arepsilon_{2p}$
1.0	1.6, 3.2, 6.4, 12.8, 25.6	179.1	244 (NA)	4.01	10.518	50.5898	26.683
1.5	0.8, 1.6, 3.2, 6.4, 12.8	92.15	244 (NA)	1.88	3.6738	21.3545	11.154
2	0.4, 0.8, 1.6, 3.4, 6.8	48	250	1.115	1.48471	11.3649	5.8753
3	0.2, 0.4, 0.8, 1.6, 3.4	24.5	175	0.545	0.11385	4.47073	2.25356
4	0.2, 0.4, 0.8, 1.6, 3.4	21.2	7.0	0.34	-0.268848	2.18313	1.06314
5	0.1, 0.2, 0.4, 0.8, 1.6	11.2	3.7	0.235	-0.40474	1.18372	0.547955
6	0.1, 0.2, 0.4, 0.8, 1.6	10.4	2.5	0.18	-0.458898	0.675591	0.288794
7	0.1, 0.2, 0.4, 0.8, 1.6	17.7	2.25	0.15	-0.481704	0.389716	0.145105
8	0.1, 0.2, 0.4, 0.8, 1.6	10.2	0.195	0.122	-0.491112	0.217062	0.0591604
9	0.1, 0.2, 0.4, 0.8, 1.6	10.2	0.07	0.103	-0.495291	0.10651	0.00450443
10	0.1, 0.2, 0.4, 0.8, 1.6	10.2	0.0365	0.088	-0.497104	0.0327616	-0.0321549
11	0.1, 0.2, 0.4, 0.8, 1.6	10.1	0.023	0.076	-0.497889	-0.0179536	-0.0578267
12	0.1, 0.2, 0.4, 0.8, 1.6	10.1	0.018	0.0665	-0.498227	-0.0535872	-0.0764093
13	0.1, 0.2, 0.4, 0.8, 1.6	10.1	0.015	0.059	-0.498372	-0.0789901	-0.0901715
14	0.1, 0.2, 0.4, 0.8, 1.6	10.05	0.014	0.053	-0.498435	-0.0972631	-0.100500
15	0.1, 0.2, 0.4, 0.8, 1.6	10.1	0.014	0.048	-0.498461	-0.108291	-0.108291

our implementation. The confined-system energies should be above the ground-state energies (in the basis selected) of the corresponding free systems and approach those free-system energies (and bond length for the molecule) in the limit of large box volume.

For the hydrogen atom in the center of a cubical box, Fig. 1(a) shows the ground-state energy for confinements $L\leqslant 10$ bohrs. (The basis exponents are shown in Table I; their selection is described in the next section.) The calculations were carried out to L=30 bohrs. Beyond 10 bohrs there was negligible difference in the energy with respect to the free-atom energy in the same basis, precisely as expected. At L=30 bohrs the ground-state energy is identical to that of the free-atom GTO calculation using the same basis, namely, -0.498476 hartree, validating our overlap-, kinetic-, and nuclear-energy integral calculations. A fit to the energies as a function of cube edge L,

$$E(L) = a/L^2 + b/L + c$$
, $0 \le L \le 10$ bohrs, (9)

yields a=14.5733 hartree bohr, b=-3.82369 hartree bohr, c=-0.238258 hartree. Though this fit does not have the correct infinite-size limit, it is the best fit of this simple form for $0 \le L \le 10$ bohrs. From this fit the pressure, p=-dE/dV ($V=L^3$) can be calculated; see Fig. 1(b).

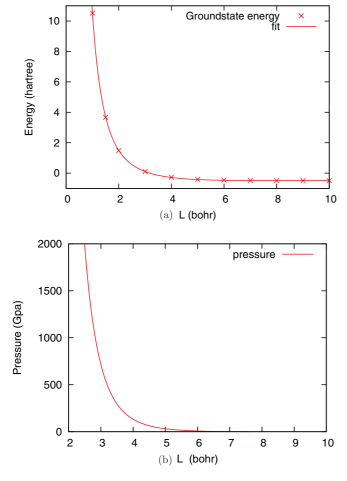


FIG. 1. (Color online) (a) Ground-state energy of an H atom in the center of a cube with edge length L. Fit is for function $f(L)=a/L^2+b/L+c$. (b) Pressure calculated from the energy fit.

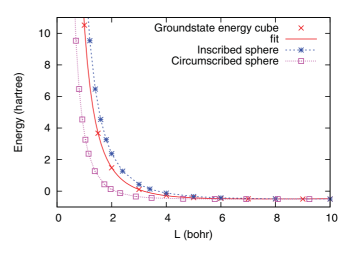


FIG. 2. (Color online) Comparison of ground-state energy of an H atom in a cube with its energy within two bounding spheres.

The effects of spherical versus cubic confinement can be assessed easily by comparison of the two confinement types both for bounding volumes and for equal volumes. For bounding volumes, the circumscribed-sphere $R = \sqrt{3}L/2$ and inscribed-sphere R = L/2 results from Ref. 71 (also see Ref. 72) can be compared to our cube results. Figure 2 shows

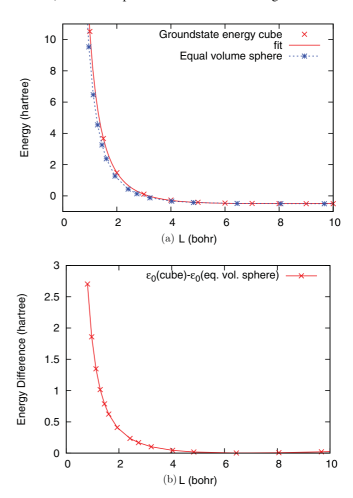


FIG. 3. (Color online) (a) Comparison of energy of an H atom in a cube with that in a sphere of equal volume. (b) The difference of the two ground-state energies.

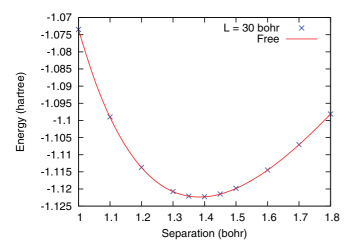


FIG. 4. (Color online) Energy vs atomic separation for the H_2 molecule centered in a 30-bohr cube. The ground state agrees exactly with the free GTO calculation using the same primitives as a basis.

that the energy of the cubically confined system is bounded by that of the two spherical systems, as expected. Shape effects of confinement are shown in Fig. 3. Though the spherical system has a lower energy than that of the equal-volume cubical system, they are quite close until the cube is smaller than $L \approx 4$ bohrs, where a significant indication of shape dependence begins. At L=1 bohr the difference in energy is over 25%, as shown in Fig. 3(b). A simpler basis of six s-type TGTOs with exponents $[0.15,0.3,0.6,1.2,2.4,4.8 \text{ bohr}^{-2}]$ reproduces the ground-state energies for $L \geqslant 1.5 \text{ bohrs}$.

Next consider the confined H_2 molecule at zero temperature. Here we used the simpler six-TGTO basis just given. For a large cube, the new TGTO confined-box computations again should conform to known results for the integrals and produce essentially the energy vs bond length curve for the free molecule. With L=30 bohrs and the molecule centered in the cube and aligned along the body diagonal, we get energies shown as points in Fig. 4. These agree completely with the values of the free GTO calculation, which are shown in Fig. 4 as the curve labeled "free". The minimum is at 1.383 bohrs, satisfactorily close to the free-molecule RHF

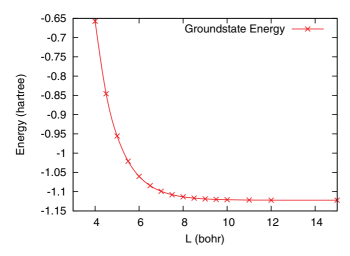


FIG. 5. (Color online) Ground-state energy for H_2 with a bond length of 1.4 bohrs confined in a cube of edge length L.

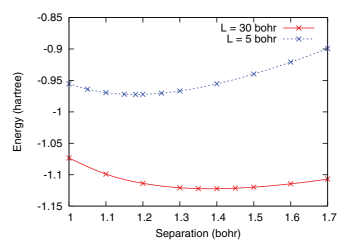


FIG. 6. (Color online) Energy vs bond length for the H_2 molecule centered in cubes L=30 and L=5 bohrs. The minima are at 1.383 and 1.178 bohrs, respectively.

value of 1.385 bohrs from a 6–31 G^{**} basis calculation.⁷³ Conversely, for fixed R (at 1.4 bohrs) and decreasing L (to 4 bohrs), the ground-state energy behavior is shown in Fig. 5. The onset of confinement effects becomes visible in the vicinity of $L \approx 11$ bohrs. Optimization of the bond length R at L = 5 bohrs is shown in Fig. 6. The total energy is, of course, higher than for the larger box, and the optimal R is shifted down to 1.178 bohrs from 1.383 bohrs.

Following this method, we obtain the optimized R and energy for decreasing L. A function fit to the energy similar to that used earlier yields R as a function of the pressure, as shown in Fig. 7.

B. Finite temperature

For finite-temperature calculations, at least a single p orbital needs to be included, and as many orbitals as feasible should be available to represent the fractionally occupied levels, which become increasingly important as T is increased. For all the following calculations, except where noted, the basis consists of seven s-type primitive GTOs and one p_x ,

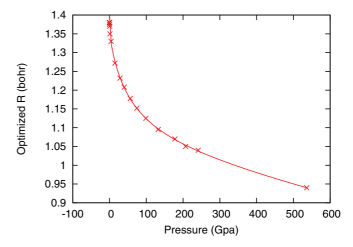


FIG. 7. (Color online) Optimized bond length R for the H_2 molecule vs pressure.

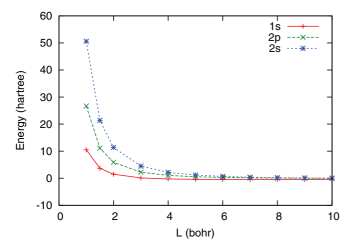


FIG. 8. (Color online) Lowest five eigenstate energies of H as a function of cube edge L. Energy values are from Table I.

one p_y , and one p_z GTO, with the p-GTO exponents equal. An elementary exponent optimization was done as follows. A set of five s-type exponents was picked and held fixed: $[0.1,0.2,0.4,0.8,1.6\,\mathrm{bohr}^{-2}]$. The sixth exponent was optimized to to minimize ε_{1s} for the single atom centered in a cube of specified L. With those six exponents fixed, the seventh s-GTO exponent was used to minimize ε_{2s} . With those seven fixed, the p exponent was used to minimize ε_{2p} . Additionally, for small L the smallest exponents produce orbitals that are so similarly flat that an approximate linear dependence exists and diagonalization fails. When this happens, the smallest exponent is replaced by extending the even-tempered exponent series to larger values. For example, at L=4 the 0.1 exponent is also replaced with 3.2, and at L=2 the 0.2 exponent is replaced with 6.4.

This procedure keeps the ratio of the effective length $(1/\sqrt{\alpha})$ of the most diffuse function to the edge length L at 0.75–0.79 for $L \leq 3$, with smaller ratios for larger L.

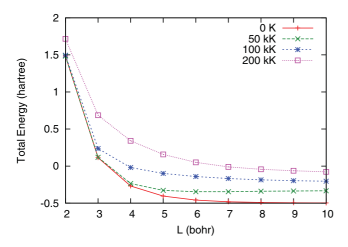
The optimization was done for each L. Table I shows the resulting exponents and energies for the orbitals that were optimized.

With this volume-dependent optimized basis, calculations were done for a single atom at the center of a cube with $1 \leqslant L \leqslant 15$ bohrs. The orbital energies of the five lowest states $(1s, 2p_x, 2p_y, 2p_z, 2s)$ are plotted in Fig. 8. Notice the inversion of ordering (2p below 2s) that is a result primarily of the confinement. To address finite temperature for this single-electron system, the one-electron levels were populated according to the Fermi-Dirac distribution. Observe that the one-electron Hamiltonian is independent of density, so the one-electron orbitals and eigenenergies are independent of occupancy, even though the density and total electronic energy are not. Figure 9(a) shows the resulting total energy as a function of L for four values of T, while Fig. 9(b) shows the free energy. The weak minimum in total energy in the vicinity of 6 bohrs at T = 50 kK appears to be a confinement effect. We have found a similar minimum at about the same volume by doing a Fermi-Dirac population of the high-precision eigenvalues of the spherically confined H atom given in Ref. 71.

Figure 10 shows the contributions to the free energy for the single atom as a function of T for four cube volumes (L=2, 3, 7, 15 bohrs). At L=7 bohrs, the KE is flat with respect to T at almost its T=0 K value. The T=50 kK nuclear-electron attraction $E_{\rm Ne}$, however, is much stronger for L=7 bohrs than for L=15 bohrs. By L=3 bohrs, the KE and $E_{\rm Ne}$ are roughly equal in magnitude.

Figure 10 also shows that the KE for the L=15 bohrs system falls with increasing temperature. Though this might seem odd, it is as it should be from virial theorem arguments for the free atom. The 2s KE is one-fourth the 1s value.⁷⁴ The finite-temperature population of the 2s and the depopulation of the 1s therefore reduces the KE with respect to its T=0 K value.

Next we turn to the system of eight H atoms. We examined a symmetric configuration in which the eight atoms were situated at the corners of a smaller cube, edge length of L/2, centered within the hard-wall cube, edge length of L. The basis used was ten s-type GTOs centered on each atom. Strict s-type symmetry is broken, of course, by enforcement of the hard-wall boundary conditions. An even-tempered set of exponents also was used in this



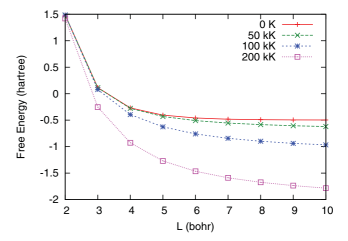


FIG. 9. (Color online) (a) Single H atom total energy, $E = \mathcal{F}_{\text{FTHF}} + T \mathcal{S}_{\text{FTHF}}$, and (b) free energy $\mathcal{F}_{\text{FTHF}}$ as a function of cube edge L, with the one-electron levels populated according to the Fermi distribution.

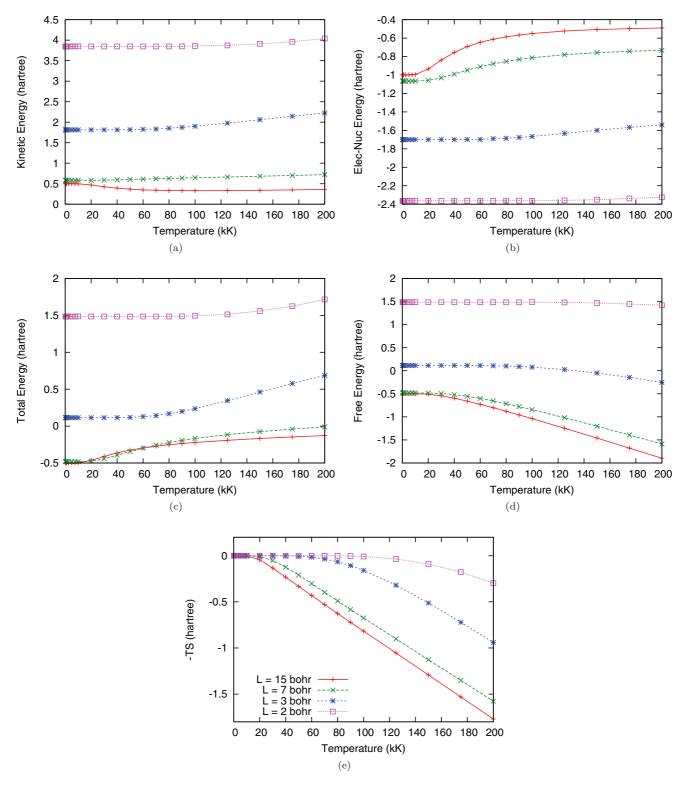
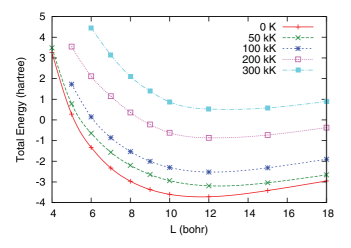


FIG. 10. (Color online) Energy components of a cubically confined single H atom vs temperature: (a) kinetic energy, (b) electron-nuclear energy, (c) total energy, (d) free energy, and (e) entropic energy [(d) minus (c)]. The key for all plots is shown in (e).

case: [0.2,0.4,0.8,1.6,3.2,6.4,12.8,25.2,50.4,100.8 bohr⁻²]. As a test, the calculations were redone with two fewer basis functions per atom; the exponents [50.4,100.8 bohr⁻²] were removed. The two calculations agreed to 2 millihartrees in total and component energies up to 200 kK. Matrix elements are calculated only once and stored, after which

fully self-consistent calculations may be done at many temperatures.

Figure 11 shows the total energy $E = \mathcal{F}_{\text{FTHF}} + T \mathcal{S}_{\text{FTHF}}$ as a function of L for various temperatures, as well as the free energy $\mathcal{F}_{\text{FTHF}}$ itself. The nuclear-nuclear repulsion energy is included (constant with respect to temperature, it varies



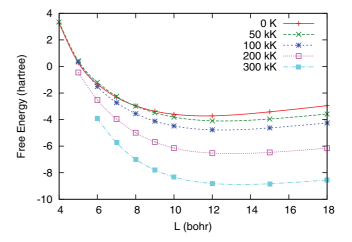


FIG. 11. (Color online) Zero- and finite-temperature total energy and free energy for eight H atoms in different sized cubes: (a) $E = \mathcal{F}_{\text{FTHF}} + T \mathcal{S}_{\text{FTHF}}$ and (b) $\mathcal{F}_{\text{FTHF}}$.

with L). Figure 12 shows various components of the free energy (electron-nuclear Coulomb energy, electron-electron Coulomb energy including exchange, kinetic energy, and entropic energy). Also shown are the total energy and free energy, with the difference of these two being the entropic energy. Again, the nuclear-nuclear repulsion energy is included in Figs. 11 and 12. It has values of 9.118, 7.598, 5.699, 4.559 hartrees for L = 5, 6, 8, 10 bohrs, respectively. The energies are shown as a function of temperature $0 \le T \le 250$ kK for four cube sizes, L = 5, 6, 8, and 10 bohrs. The comparatively flat plateau up to roughly T = 15 kK is a direct consequence of Fermi-Dirac level filling. Up to about 25 kK, the interval between the highest occupied molecular orbital (HOMO) at zero temperature and the lowest unoccupied molecular orbital (LUMO) is roughly constant at 0.5 hartree. Therefore the filling ratio of those two is roughly $\exp(-13/1.3) \approx 5 \times 10^{-5}$ or smaller up to about T = 15 kK. We return to this matter below.

C. Comparison with approximate functionals

Orbital-free treatment of WDM has been dominated, not surprisingly, by local density approximations for both the KE and exchange contributions to the free energy. For the KE, the choice is physically motivated by the fact that the high-pressure and/or high-temperature limit for a WDM system is Thomas-Fermi. This leads to finite-temperature Thomas-Fermi (FTTF)⁷⁵ $T_0 = \int \tau_0 d\mathbf{r}$ by itself or with some fraction of the von Weizsäcker contribution (in its zero-temperature form) $T_W = \int \tau_W d\mathbf{r}$, with

$$\tau_0 = \frac{\sqrt{2}}{\pi^2 \beta^{5/2}} I_{3/2}(\beta \mu),\tag{10}$$

$$n = \frac{\sqrt{2}}{\pi^2 \beta^{3/2}} I_{1/2}(\beta \mu),\tag{11}$$

$$\tau_W = \frac{|\nabla n|^2}{8n},\tag{12}$$

where I are Fermi integrals. Note a parametrized form²⁶ may be used to eliminate μ between τ_0 and n.

In Fig. 13 we compare the FTHF KE as a function of T with the FTTF KE alone and with it supplemented by both the full \mathcal{T}_W and $(1/9)\mathcal{T}_W$ for L=6 bohrs. The latter three functionals were evaluated with the FTHF density and hence are non-self-consistent. As is known at T=0, pure Thomas-Fermi underestimates the KE, while the addition of the full \mathcal{T}_W overestimates it. None of the three is close to quantitative agreement with $\mathcal{T}_{\text{FTHF}}$. Moreover, FTTF and FTTF augmented with $(1/9)\mathcal{T}_W$ can be ruled out from the T=0 K comparison since the exact (fully correlated) KE must be above $\mathcal{T}_{\text{FTHF}}$ (from a virial theorem argument).

For the exchange contribution to the free energy, the use of ground-state functionals is common (recall the Introduction), with the LDA being dominant. Figure 14 shows the FTHF exchange contribution to the free energy $\mathcal{F}_{x,\text{FTHF}}$ in comparison with the exchange free energy generated by the ground-state LDA functional and with the Perrot and Dharmawardana parametrization for the temperature-dependent LDA functional.²⁸ Again, this is for the cubic symmetry, eight-H system at L = 6 bohrs. The other functionals are evaluated with the FTHF density. (Note that such "post-self-consistentfield" evaluation is fairly common in assessment of newly developed ground-state functionals. See, for example, Ref. 76.) Here one sees a marked difference: the ground-state functional fails completely, while the temperature-dependent functional has at least semiquantitatively correct temperature dependence.

We may also make some semiquantitative comparison with a more widely used model for extended systems at substantial T. In Fig. 15 we show the internal energy per atom of the eight hydrogen atoms in the cubic symmetry arrangement at L=7 bohrs (corresponding to average $r_s=2.17$), with that of the average-atom DFT calculation of Dharma-wardana and Perrot⁷⁷ at $r_s=2$, with $(4\pi/3)r_s^3=L^3/8$. Their system includes DFT exchange and correlation, whereas we have pure Hartree-Fock exchange. Additional energy differences are due to the different boundary conditions. Decomposing the near-parallel temperature dependence into those components, however, is a task outside the scope of the present work. We

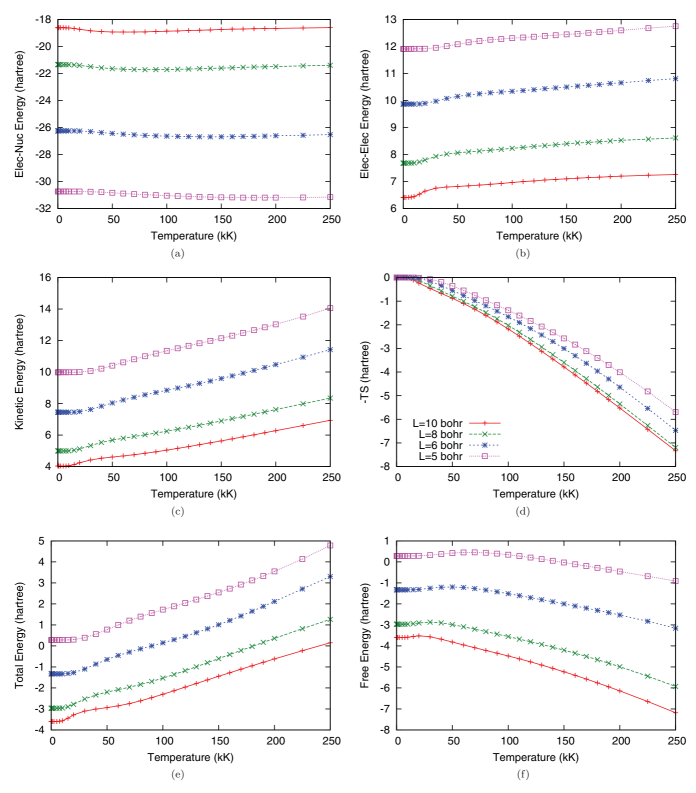


FIG. 12. (Color online) Self-consistent FTHF free-energy contributions for eight confined H atoms in a cubic array as a function of T for four different cube edge lengths L. The key for all plots is given in (d).

can see one aspect immediately. Though the kinetic energy is the major contributor to the change in total energy as a function of temperature (recall Fig. 12), the change due to electron-electron interaction, including exchange, is at least a third that of the kinetic energy.

IV. DISCUSSION AND CONCLUSIONS

Comparison, evaluation, and betterment of functionals for WDM simulations is the long-term motive of this work. As just shown, even at this initial state of development (Hartree-Fock, small particle number, no molecular dynamics), the approach

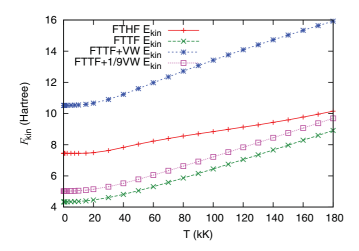


FIG. 13. (Color online) FTHF kinetic energy compared with finite-temperature Thomas-Fermi KE and two forms of von Weizsäcker augmentation of FTTF for the eight-H cubic system at L=6 bohrs. See text for details.

gives insight regarding that task. There are some specific issues worth discussion also.

A. One-particle spectrum effects

It is well known that zero-temperature HF calculations overestimate both band gaps in solids and the so-called HOMO-LUMO gaps of finite systems. This occurs because the occupied HF orbitals are free of self-interaction, but the unoccupied ones are not. As temperature increases in the FTHF scheme, however, levels unoccupied at T=0 K become increasingly occupied and shifted. In the self-consistent solution of the HF problem in a basis, those levels then contribute to the Hamiltonian matrix ("Fock matrix" in quantum chemistry terminology). Thus there are two questions to address. What is the extent to which FTHF exhibits HOMO-LUMO gap overestimation similar to that of ground-state HF? At what fractional occupancy is an energy level changed from an overestimated virtual to a more properly estimated partially occupied one?

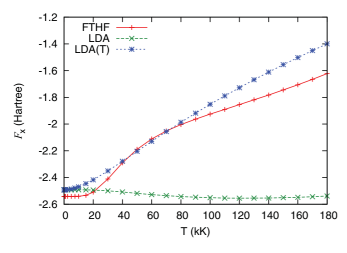


FIG. 14. (Color online) FTHF exchange free-energy ground-state LDA and Perrot and Dharma-wardana temperature-dependent LDA (eight-H cubic system, L=6 bohrs). See text for details.

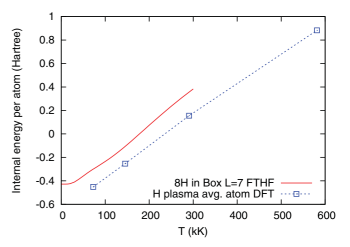


FIG. 15. (Color online) FTHF internal energy per atom for the eight-H cubic-symmetry system at L=7 bohrs compared with $r_s=2$ hydrogen plasma average atom DFT calculation.

For specificity, we consider the eigenspectrum of a single, moderately compressed eight-atom cube with L=6 bohrs. Figure 16 shows the Fermi distribution of the single-particle energies for four temperatures. The continuous curves in Fig. 16 show the Fermi function with calculated chemical potential μ . The discrete points mark the input energies to the calculation of μ . Note that these distributions are for one spin. Also keep in mind that the cubical symmetry causes the lowest four one-electron orbitals to group as singly degenerate and triply degenerate (a_{1g}, t_{1u}) in crystal-field notation). At zero temperature, therefore, only two points are shown with unit occupancy, but the higher-energy point corresponds to three degenerate HOMO states (indexed as 2, 3, 4). The LUMO is degenerate states 5, 6, and 7, with the singly degenerate state 8 above them (again, as would be expected from a cubical crystal field). For simplicity of discussion, the HOMO and LUMO (at T=0) are labeled ε_4 and ε_5 , respectively. One can see that the spacing between ε_4 and ε_5 decreases with temperature. This difference is shown directly in Fig. 17 along with the occupation number for the ε_5 level. Then in Fig. 18,

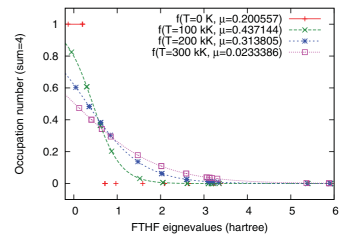


FIG. 16. (Color online) Fermi distribution for eight H atoms in a box, L=6 bohrs. Points may represent more than one state due to energy degeneracies.

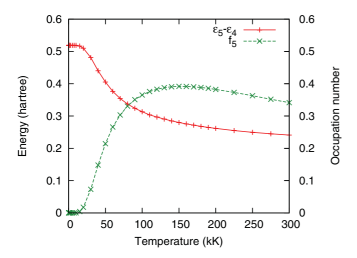


FIG. 17. (Color online) Energy difference between the fourth and fifth energy levels along with the occupation number of the fifth energy level. Eight H atoms, cubic box, L = 6 bohrs.

 μ is plotted as a function of T. Observe that the chemical potential is nearly midway between ε_4 and ε_5 up to just below 50 kK. This behavior is exact at zero temperature.

Those total-energy and eigenspectrum results together resolve the matter of the behavior of what would be virtual states at zero temperature in FTHF. First, examination of the kinetic- and total-energy plots for all box sizes shows that there is a change in the form of the temperature dependence at roughly 20 kK. That change is complemented by the change in the spacing between the ε_4 and ε_5 levels. They are essentially static for lower temperatures, then begin to change abruptly well below 50 kK, and then change more moderately at higher temperatures. Thus, above 50 kK, states corresponding to zero-temperature virtuals are sufficiently incorporated in the interaction terms to make a material modification of T=0behavior. However, as the temperature is decreased below roughly 50 kK, the FTHF Coulomb and exchange terms increasingly are dominated by the T=0 occupied levels, which therefore keeps the lightly occupied higher-energy

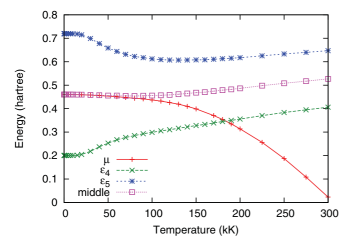


FIG. 18. (Color online) Chemical potential μ and the fourth and fifth energy levels for eight H atoms in a cubic box, L=6 bohrs. The curve labeled "middle" is half way between ε_4 and ε_5 .

levels artificially high. This is not a basis issue, but an issue with discrete eigenstates. In a solid such as jellium or a metal with a continuum of states, this should not be an issue, but for a system with energy gaps the issue remains.

B. Other findings and considerations

The preceding discussion about depopulation and repopulation of levels relative to the ground-state HF HOMO-LUMO gap illustrates a broader challenge for the construction of approximate functionals for the various contributions to the free energy. Whether explicit or implicit, such approximate functionals correspond to restricting the required traces to specific classes (or subsets of classes) of state functions. The consequence of such a restriction is to incorportate the spectral properties of that class into the approximate functionals. For example, in FTHF that class is single Slater determinants constructed with respect to ground-state HF minimization. Although we have not attempted its construction here, in principle, there is an FTHF free-energy functional. It would have exactly the same problem with a plateau in its T dependence as we have found here.

The small number of particles is another issue. For sufficiently large numbers, all standard ensembles (grand canonical, canonical, and microcanonical) give the same thermodynamics. Fluctuations characteristic of small particle counts can degrade that relationship. The main issue regarding particle count is computational cost. The problem is endemic to many computational studies, especially when a large temperature and pressure domain such as characterizes WDM is involved (see, for only one example, Ref. 78). At the least, we have an even-handed comparison of different methods (e.g., the comparison of functionals given above) for a given number of electrons and of ions.

Clearly, we have shown that the TGTO basis is feasible and effective. As is typical of GTO basis methods, the computational cost is essentially entirely in the CPU time for the calculation of the two-electron integrals. The eight-atom systems described are calculated with 64 or 80 total basis functions, making diagonalization trivial. A simple double array of all N^4 two-electron integrals only occupies 128 or 312.5 MB, respectively, storable in memory. In practice, we calculated N^4/λ integrals, with $\lambda = 7.60$, 7.68 and not 8 due to the looping procedure we used. While, for the cubic system discussed here, this calculation requirement could have been reduced further by exploiting symmetry, we need the capability to explore other, lower-symmetry geometries. Note, however, that if the box size or the atomic positions are changed, all integrals affected by the change must be recalculated.

On a modern desktop processor (Intel Core i5 650 at 3.2 Ghz) one two-electron integral can be calculated in about 29.1 ms. So the times to calculate all integrals for a 64- and an 80-orbital basis would be 17.84 and 43.13 h. The integrals are calculated independently, so they can be parallelized effectively. Calculations reported in this work were done on the University of Florida High Performance Computing Center Linux clusters.

Though these are quite acceptable costs for fixed geometries and small numbers of ions and electrons, the burden becomes formidable for direct application in Born-Oppenheimer molecular dynamics. We are currently working on ways to ameliorate that problem.

ACKNOWLEDGMENTS

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APPENDIX A: CORRECTION FOR PIECEWISE INTEGRALS FOR THE BASIS FUNCTIONS

From the definition of the basis functions in Eqs. (7) and (8), it follows that derivatives of the basis function may not be continuous at x_c , the center of the function. A discontinuity of the second derivative would, of course, be significant for the kinetic energy. The issue is whether the kinetic-energy matrix elements can be evaluated piecewise, as is the case with the overlap, nuclear-electron, and electron-electron integrals. We may examine this issue by writing the basis formally with Heaviside functions as follows:

$$\varphi = [\theta(x) - \theta(x - x_1)]\varphi_0 + [\theta(x - x_1) - \theta(x - L)]\varphi_L.$$
 (A1)

For the TGTO basis, the identification from Eq. (8) is

$$\varphi_0 = a_0[g^n(x) - \delta_0],
\varphi_L = a_L[g^n(x) - \delta_L],
g^n(x) = (x - x_1)^n e^{-\alpha(x - x_1)^2}, \quad x_1 = x_c.$$
(A2)

The derivatives are

$$\frac{\partial \varphi}{\partial x} = [\delta(x) - \delta(x - x_1)]\varphi_0 + [\theta(x) - \theta(x - x_1)]\varphi_0'$$

$$+ [\delta(x - x_1) - \delta(x - L)]\varphi_L$$

$$+ [\theta(x - x_1) - \theta(x - L)]\varphi_L'$$
(A3)

and

$$\frac{\partial^{2} \varphi}{\partial x^{2}} = [\delta'(x) - \delta'(x - x_{1})]\varphi_{0} + 2[\delta(x) - \delta(x - x_{1})]\varphi'_{0}
+ [\theta(x) - \theta(x - x_{1})]\varphi''_{0}
+ [\delta'(x - x_{1}) - \delta'(x - L)]\varphi_{L}
+ 2[\delta(x - x_{1}) - \delta(x - L)]\varphi'_{L}
+ [\theta(x - x_{1}) - \theta(x - L)]\varphi''_{L}.$$
(A4)

Now consider a generic kinetic-energy matrix element involving the foregoing function and another similar basis function χ with left and right constituents χ_0 , χ_L , centered at x_2 . Without loss of generality, take $x_1 < x_2$. Then

$$\int_0^L \chi \frac{\partial^2 \varphi}{\partial x^2} \, dx := I_A + I_B. \tag{A5}$$

The terms of the second derivative with the Heaviside functions contribute just the piecewise integration I_A , while the δ function and first derivative δ function terms contribute I_B . Of the terms in Eq. (A4) only those at x_1 contribute, as the

constituents χ_0 , χ_L , φ_0 , and φ_L go to zero at x=0 and x=L. Thus

$$I_B = \int \chi_0 [\delta'(x - x_1)(\varphi_L - \varphi_0) + 2\delta(x - x_1)(\varphi_L' - \varphi_0')] dx.$$
(A6)

From the definition of δ' this expression becomes

$$I_{B} = \int \chi_{0}[-\delta(x - x_{1})(\varphi'_{L} - \varphi'_{0}) + 2\delta(x - x_{1})(\varphi'_{L} - \varphi'_{0})] - \chi'_{0}\delta(x - x_{1})(\varphi_{L} - \varphi_{0}) dx,$$
(A7)

which reduces to

$$I_B = [\chi_0(\varphi_L' - \varphi_0') - \chi_0'(\varphi_L - \varphi_0)]|_{x = x_1}.$$
 (A8)

For the case of $x_1 = x_2$, that is, for diagonal terms or functions that have the same center, χ'_0 must be replaced in Eq. (A7) by the analog of Eq. (A3), with the result

$$I_{B} = \chi_{0}(\varphi'_{L} - \varphi'_{0})|_{x=x_{1}} - \int \delta(x - x_{1})(\varphi_{L} - \varphi_{0})\{[\delta(x) - \delta(x - x_{1})]\chi_{0} + [\theta(x) - \theta(x - x_{1})]\chi'_{0} + [\delta(x - x_{1}) - \delta(x - L)]\chi_{L} + [\theta(x - x_{1}) - \theta(x - L)]\chi'_{L}\} dx.$$
(A9)

Note χ_0 in the first term follows because the functions themselves are continuous, while the first and second derivatives may not be. For the same reason, it follows that the remaining integral in Eq. (A9) and the second term of Eq. (A8) vanish. Thus so long as the functions are continuous, the correction to the piecewise kinetic-energy integral is simply

$$I_B = \chi(\varphi_L' - \varphi_0')|_{x=x_1},$$
 (A10)

and so long as the first derivative is continuous, this reduces to zero.

With continuity of the basis functions enforced by construction, only the first derivative needs to be examined for a possible correction to simple piecewise integration. For the basis defined in Eq. (8), those corrections are

$$g^{0}(x) = e^{-\alpha(x-x_{1})^{2}} \quad I_{B} = 0,$$

$$g^{1}(x) = (x-x_{1})e^{-\alpha(x-x_{1})^{2}} \quad I_{B} = \chi(a_{L}-a_{0}), \quad (A11)$$

$$g^{2}(x) = (x-x_{1})^{2}e^{-\alpha(x-x_{1})^{2}} \quad I_{B} = 0.$$

Basis functions $g^n(x)$ with higher powers of the prefactor $(x - x_1)$ all have continuous first and second derivatives at x_1 . In fact, those derivatives are all zero. So only the p-type basis functions (n = 1) have a kinetic-energy matrix-element contribution beyond that given by piecewise integration.

APPENDIX B: FINITE-RANGE GAUSSIAN INTEGRALS

Following the methods of Boys, 65-69 we use the transform of the Coulomb potential to separate the Coulomb integrals

into one-dimensional Cartesian pieces:

$$\frac{1}{|\mathbf{r} - \mathbf{R_N}|} = \frac{1}{\sqrt{\pi}} \int_{-\infty}^{\infty} e^{-s^2(\mathbf{r} - \mathbf{R_N})^2} ds$$

$$= \frac{1}{\sqrt{\pi}} \int_{-\infty}^{\infty} e^{-s^2(x - X_N)^2} e^{-s^2(y - Y_N)^2} e^{-s^2(z - Z_N)^2} ds.$$
(B1)

Hence all Coulomb integrals require integration over the transform variable s, which is done by Gauss-Laguerre quadrature.

For 1D primitive TGTOs, we note the required finite range (a,b) integrals are of the form

$$I_n = \int_a^b x^n e^{-\alpha(x - x_c)^2} dx.$$
 (B2)

This simply transforms to

$$I_n = \int_{a-x_c}^{b-x_c} (x' + x_c)^n e^{-\alpha x^2} dx'.$$
 (B3)

This result leaves us needing to compute

$$J_n = \int_{a'}^{b'} x^n e^{-\alpha x^2} dx. \tag{B4}$$

Integrating by parts, we find

$$J_n = -x^{n-1} \frac{e^{-\alpha x^2}}{2\alpha} \bigg|_{a'}^{b'} + \frac{n-1}{2\alpha} J_{n-2}.$$
 (B5)

So with the initial two integrals, we may find the higher-order integrals by recursion:

$$J_0 = \int_{a'}^{b'} e^{-\alpha x^2} dx = \frac{\sqrt{\pi}}{2\sqrt{\alpha}} \operatorname{erf}(\sqrt{\alpha}x) \Big|_{a'}^{b'},$$
 (B6)

$$J_1 = \int_{a'}^{b'} x e^{-\alpha x^2} dx = -\frac{e^{-\alpha x^2}}{2\alpha} \bigg|_{a'}^{b'}.$$
 (B7)

All two-center (overlap, nuclear-electron, kinetic) integrals reduce to expressions in terms of I_n . After two applications of the Gaussian product rule, four-center (two-electron) integrals reduce to terms of the form

$$\int_{a_2}^{b_2} \int_{a_1}^{b_1} x_1^m x_2^m e^{-\alpha_1(x_1 - x_i)^2} e^{-\alpha_2(x_2 - x_j)^2} e^{-s^2(x_1 - x_2)^2} dx_1 dx_2.$$
(B8)

Two further applications of the product rule bring us to the form

$$\int_{a_2}^{b_2} \int_{a_1}^{b_1} x_1^n x_2^m e^{-\kappa_1 [x_1 - R_1(x_2)]^2} e^{-\kappa_2 (x_2 - R_2)^2} dx_1 dx_2.$$
 (B9)

Here the integral over x_1 may be evaluated as I_n , so we are left with

$$\int_{a_2}^{b_2} I_n(x_2) \Big|_{x_1 = a_1}^{x_1 = b_1} x_2^m e^{-\kappa_2(x_2 - R_2)^2} dx_2, \tag{B10}$$

which we evaluate by Gauss-Legendre quadrature.

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