Density scaling has a rich history in density functional theory, providing exact conditions for use in the construction of ever more accurate approximations to the unknown exchange-correlation functional. We define a conjugate potential scaling for each density scaling. This provides exact relations on various important density functionals, in particular, relating functionals evaluated on exact densities of different potentials. We illustrate these conditions on the two- and four-electron ion series.

Recently, a different form of density scaling was used in the development of the PBEsol functional[19]. Here, both the coordinate and the particle number are scaled, leading to new insights into the XC functional. We refer to this as charge-neutral scaling[20], as it is equivalent to simultaneously changing the charges on atoms and the number of electrons, so as to keep overall neutrality.

In this paper, we extend the use of density scaling as a tool in DFT. Most importantly we introduce the concept that any form of density scaling defines a related form of potential scaling. This leads to more exact conditions on the various DFT quantities as functionals of densities of different particle number. Yang and others[21] have emphasized the duality of the potential with the density, but have not related scaling of one to the other.

II. BACKGROUND

In DFT, the total ground-state energy for electrons in a given external potential, \( v(\mathbf{r}) \) is given as a functional of the ground-state density \( n(\mathbf{r}) \),

\[
E_e[n] = F[n] + V[n],
\]

where the external potential energy as a functional of the density is

\[
V[n] = \int d^3r \ n(\mathbf{r}) v(\mathbf{r}),
\]

and \( F[n] \) is the universal functional which may be defined via a constrained search[22] over all wavefunctions \( \Psi \) that yield density \( n(\mathbf{r}) \)

\[
F[n] = \min_{\Psi \rightarrow n} \langle \Psi | \hat{T} + \hat{V}_{ee} | \Psi \rangle,
\]

\[
= T[n] + V_{ee}[n].
\]

In the Kohn-Sham method, a reference system of non-interacting electrons with the same density is solved, so it is useful to write

\[
F[n] = T[n] + U[n] + E_{xc}[n],
\]

I. INTRODUCTION

Density functional theory (DFT) has become the method of choice for many electronic structure calculations in both computational chemistry and condensed-matter physics[1]. It balances the competing demands of accuracy and computational efficiency. The foundation of DFT is the Hohenberg-Kohn theorem[2] which states that the total ground-state energy of a system can be written as a functional of the ground-state density. This allows the construction of the Kohn-Sham[3] system of non-interacting fermions in a unique one-body potential, which is defined to yield the same density as the real interacting system. Since the ground-state energy is a density functional, this then gives the ground-state energy. However, this functional is not known exactly and in practice, the unknown exchange-correlation (XC) contribution must be approximated.

The plethora of XC functionals in current use in DFT has a rich history in density functional theory, providing exact conditions for functional development[4]. In this regard, the more exact constraints that can be discovered, the more tools available for developers to use in their quest for more accurate approximations. Any useful approximation in DFT should try to satisfy these constraints, or be tested against them.

Density scaling has been a particularly useful tool for the analysis and development of DFT. A singular example is uniform coordinate scaling[5], where the coordinates of a given density are linearly scaled, but normalization is preserved. This has led to fundamental exact conditions on the exchange-correlation (XC) energy functional[5–8]. For example, the form of the local approximation to the exchange energy can be deduced from this scaling. The adiabatic-connection formulation[9–12], much studied and used in DFT development, is essentially an integral over the uniform coordinate scaling parameter[5, 13, 14]. Here, the electron-electron interaction is scaled by a constant while the density is kept fixed, linking the non-interacting Kohn-Sham and the fully interacting systems, and leads to many more conditions. For example, the adiabatic connection formula is behind rationalizing the hybrid approach[15–18].
where \( T_0 \) is the kinetic energy of non-interacting electrons of density \( n(r) \), \( U \) is the Hartree energy and \( E_{xc} \) is the exchange-correlation energy, defined by Eq. (5). The Hartree energy (sometimes denoted by \( E_H \)) is defined as

\[
U[n] = \frac{1}{2} \int d^3r \int d^3r' \frac{n(r)n(r')}{|r-r'|}. \tag{6}
\]

In Thomas-Fermi (TF) theory\([23, 24]\), the universal functional is approximated as

\[
F[n] \approx F^{TF}[n] = T_s^{(0)}[n] + U[n], \tag{7}
\]

where \( T_s^{(0)} \) is the local kinetic energy functional,

\[
T_s^{(0)}[n] = A_s \int d^3r \, n^{5/3}(r), \tag{8}
\]

with \( A_s = (3/10)(3\pi^2)^{2/3} \).

### III. POTENTIAL SCALING

Consider a density \( n(r) \) that is the ground-state density of some interacting problem with potential \( v(r) \). Now, introduce some positive parameter, \( 0 < \gamma < \infty \), which produces a family of densities, \( n_\gamma(r) \), with \( \gamma \) defined so that \( \gamma \to \infty \) corresponds to the high-density limit. A simple example is the uniform coordinate scaling of Levy and Perdew\([5]\):

\[
n_\gamma(r) = \gamma^3 n(\gamma r), \quad 0 < \gamma < \infty, \tag{9}
\]

where the prefactor was chosen to keep the density normalized. For example, under uniform coordinate scaling with \( \gamma > 1 \), the density of He is squeezed into a smaller volume, and looks like a distorted version of a two-electron ion\([25]\). This scaling has become a mainstay of DFT and leads to many important results. Most importantly, when particles interact, the coordinate-scaled wavefunction is not the ground-state wavefunction of the scaled density. Considering such a wavefunction as a trial state in the Rayleigh-Ritz principle yields useful inequalities for the various density functionals\([5]\):

\[
T[n_\gamma] \leq \gamma^2 T[n], \quad \gamma \geq 1, \tag{10}
\]

\[
V_{ee}[n_\gamma] \geq \gamma V_{ee}[n], \quad \gamma \geq 1, \tag{11}
\]

and a similar condition applies for the correlation energy \( E_{xc}[n] \) itself.

A second example that we focus on here is what we call charge-neutral (CN) scaling, in which

\[
n_\zeta(r) = \zeta^2 n(\zeta^{1/3} r), \quad 0 < \zeta < \infty \tag{12}
\]

and so \( N_\zeta = \zeta N \). We use \( \zeta \) as the scaling parameter to distinguish from coordinate scaling. This choice both scales the coordinate and changes the particle number.

For Coulomb-interacting matter, this ensures neutrality as a function of \( \zeta \). For example, for single atoms, it simply implies \( Z_\zeta = \zeta Z \) and the atom remains neutral. Lieb and Simon\([28]\) showed that Thomas-Fermi (TF) theory becomes exact for neutral atoms as \( \zeta \to \infty \), and Lieb\([29]\) later generalized the proof to all Coulomb-interacting matter. In Fig 1, we illustrate this scaling on the He atom density.

In both coordinate and CN scaling, as the scaling parameter is taken to \( \infty \), the solution simplifies. Under uniform coordinate scaling to the high-density limit, the system becomes effectively non-interacting. Under CN scaling to the high-density limit, Thomas-Fermi theory becomes relatively exact. In either case, we can ask how the potential changes when the density is scaled. We define this as the potential scaling conjugate to the given density scaling, but consider it for all values of the scaling parameter, not just in the high-density limit.

Under coordinate scaling, in the large \( \gamma \) limit,

\[
v^\gamma(r) = \gamma^2 v(\gamma r). \tag{13}
\]

We therefore define our potential scaling by this equation, applied for all \( \gamma \). We use a superscript to indicate that the potential has been scaled, not the density. This is simply how the external potential would change when the density is scaled, if the particles were non-interacting particles. For example, for a neutral atom, this changes the nuclear charge by \( \gamma \), keeping the particle number fixed. As \( \gamma \to \infty \), the repulsion between electrons becomes negligible relative to the nuclear attraction, and the density becomes that of the non-interacting limit, scaled by \( \gamma \).

Similarly, under CN scaling with \( \zeta \to \infty \), the TF equations become relatively exact\([30]\), and

\[
v^\zeta(r) = \zeta^{4/3} v(\zeta^{1/3} r), \quad N_\zeta = \zeta N. \tag{14}
\]

Again, the conjugate potential scaling is defined by this, applied to all values of \( \zeta \). Analogously, if self-consistent
IV. UNIFORM COORDINATE SCALING

In the old work[5], Levy and Perdew compared two different wavefunctions with the same density, whereas we compare two different wavefunctions in the same potential. To do this, begin from a given potential \( v(r) \) with ground-state density \( n(r) \). Define \( n^\gamma(r) \) as the ground-state density of \( v^\gamma(r) \), given by Eq. (13). Then \( n^\gamma(r) \) is a useful trial density for the original problem. It is found by first scaling the potential, solving the problem, and then scaling backwards to the original problem. (In Fig. 1, the dashed line corresponds \( n^\gamma(r) \) for the He density, with \( \gamma = 2 \).) This is exactly what was done (but with an approximate scale factor) in Ref. 25.

If \( n^\gamma_{1/\gamma}(r) \) is used as trial density for \( v(r) \), the variational principle states that

\[
F[n^\gamma_{1/\gamma}] + \gamma^{-2}V^\gamma[n^\gamma] \geq F[n] + \gamma^{-2}V^\gamma[n],
\]

which may be rearranged as

\[
F[n^\gamma_{1/\gamma}] - F[n] \geq \gamma^{-2}(V^\gamma[n] - V^\gamma[n^\gamma]).
\]

Conversely, \( n_{\gamma}(r) \) may be used as a trial density for \( v^\gamma(r) \), yielding

\[
E_{v^\gamma}[n_{\gamma}] = F[n_{\gamma}] + V^\gamma[n_{\gamma}] \geq F[n^\gamma] + V^\gamma[n^\gamma],
\]

which can also be rearranged as

\[
F[n^\gamma] - F[n_{\gamma}] \leq V^\gamma[n_{\gamma}] - V^\gamma[n^\gamma].
\]

Combining the two inequalities yields a constraint on the universal functional \( F[n] \):

\[
F[n^\gamma_{1/\gamma}] - \frac{F[n^\gamma]}{\gamma^2} \geq F[n] - \frac{F[n_{\gamma}]}{\gamma^2},
\]

which may be written in a concise form, with \( \lambda = 1/\gamma \),

\[
\Delta F^\lambda[n^{1/\lambda}] \geq \Delta F^\lambda[n],
\]

where

\[
\Delta F^\lambda[n] = F[n] - \lambda^2 F[n_{1/\lambda}].
\]

Now, \( F[n] \) is typically dominated by the kinetic energy contribution, but this can be removed, because \( T_s[n_{\gamma}] = \gamma^2 T_s[n] \). Thus

\[
\Delta E^\lambda_{\text{HXC}}[n^{1/\lambda}] \geq \Delta E^\lambda_{\text{HXC}}[n],
\]

where \( E_{\text{HXC}} = U + E_{\text{XC}} \). This tells us that if we begin from, e.g., the lowest value of \( Z \) that binds a given \( N \) electrons, then \( \Delta E^\lambda_{\text{HXC}}[n^{1/\lambda}] \) is an increasing function of \( \lambda \).

Simple results can be extracted from this very general formula by taking \( \gamma \) to be very large. This makes \( n^\gamma(r) \) an essentially non-interacting density, because the external potential dominates. Thus

\[
n^{1/\lambda}(r) \rightarrow n_{\text{SI}}(r), \quad \lambda \rightarrow 0,
\]

where \( n_{\text{SI}}(r) \) is the density of the system with only an infinitesimal electron-electron repulsion. But \( \Delta E^\lambda_{\text{HXC}}[n] \) also simplifies as \( \lambda \rightarrow 0 \), because all terms scale less than quadratically. Thus

\[
\Delta E^\lambda_{\text{HXC}}[n] \rightarrow E_{\text{HXC}}[n], \quad \lambda \rightarrow 0,
\]

yielding the universal result that

\[
E_{\text{HXC}}[n_{\text{SI}}] \geq E_{\text{HXC}}[n],
\]

applying to all potentials. For \( \gamma < 1 \), Eq. (20) is less useful, as most systems of interest lose an electron when the external potential becomes too small. To further simplify Eq. (22), we note that both the Hartree and exchange energies scale linearly with \( \gamma \), i.e.,

\[
E_{\text{HX}}[n_{\gamma}] = \gamma E_{\text{H}}[n],
\]

so that

\[
\Delta E^\lambda_{\text{H}}[n] = (1 - \lambda) E_{\text{H}}[n].
\]

Inserted into Eq. (22), we find

\[
E_{\text{H}}[n_{\gamma}] + \Delta E^\lambda_{\text{XC}}[n_{\gamma}] \geq E_{\text{H}}[n] + \Delta' E^\lambda_{\text{C}}[n],
\]

where

\[
\Delta' E^\lambda_{\text{C}}[n] = \Delta E^\lambda_{\text{C}}[n]/(1 - \lambda).
\]

The simplest way to test this result is by doing a Kohn-Sham calculation without any correlation (such as oep exact exchange). Then the correlation contributions vanish on both sides of Eq. (28), and so

\[
E_{\text{H}}[n] \leq E_{\text{H}}[n_{\gamma}]/\gamma \leq E_{\text{H}}[n_{\text{SI}}]
\]

This simplifies even further for the special case of two electrons in a spin singlet, where \( E_{\text{X}}[n] = -U[n]/2 \), so the inequality becomes a bound on the Hartree energy alone:

\[
U[n] \leq U[n_{\gamma}]/\gamma \leq U[n_{\text{SI}}]
\]
TABLE I: The Hartree energies, $U$, for the helium isoelectronic series as calculated with the oep exact-exchange method as implemented in the OPMKS code[32]. We also demonstrate how, for two values of atomic number $Z'$, the inequalities of Eq. (31) with $\gamma = Z'/Z$, are satisfied. Note that if $\gamma < 1$, the inequality is reversed. The values for bordering values of $Z$ bracket the value of $U$ at atomic number $Z$ and these bounds become tighter as $Z'$ increases.

| $Z$ | $U$ | $Z'=4$ | $Z'=20$ |
|-----|-----|------|--------|
| 1   | 0.790970 | 3.163880 | 15.819400 |
| 2   | 2.051538 | 4.103076 | 20.515380 |
| 3   | 3.303373 | 4.404497 | 22.022487 |
| 4   | 4.554137 | 4.554137 | 22.770685 |
| 6   | 7.054819 | 4.703213 | 23.516063 |
| 10  | 12.055315 | 4.822126 | 24.110630 |
| 20  | 24.555661 | 4.911132 | 24.555661 |

In Table I, we analyze the above inequality, Eq. (31), while in Fig 2, we plot $U[n\gamma]/\gamma$ as a function of $\gamma$ for exact-exchange calculations of the two-electron ion series, beginning with H$^-$. Indeed, the function increases toward the Bohr atom limit of $5/4$, found by inserting a doubly-occupied 1s Hydrogen atom orbital into the Hartree energy.

To test the exchange contribution in a non-trivial way, i.e., Eq. (28), we repeated the calculations for the four-electron ion series, this time beginning from Be. Again the inequality is satisfied, and the limiting value is found by evaluating the Hartree and exchange energies of doubly-occupied 1s and 2s Hydrogenic orbitals, as calculated in Appendix B. These values are reported in Table II and plotted in Fig 3.

Lastly, we can even include extremely accurate estimates of the correlation contributions for the two-electron series. We work from the data in Table I of Ref. [31]. Since the two-electron ions are generally weakly correlated, one can approximate the scaling of their correlation energies with a Taylor-series around the high-density limit:

$$E_c[n] = E_c^{(0)}[n] + \lambda E_c^{(1)}[n]$$  \hspace{1cm} (32)

where $E_c^{(p)}[n]$ are scale-invariant functionals. Since $T_c = -E_c + \partial E_c[n\gamma]/\partial \gamma(\gamma = 1)[5]$, and $T_c$ is reported in their table, one can solve for these two coefficients. This yields a value of -47.6 mH for $E_c^{(0)}$ for He, in excellent agreement with the value of 47.9 estimated in Ref. [25], and predicts a value of -56.1 mH for H$^-$. Using this approximate scaling, we can insert all terms into Eq. (22) explicitly.

![Fig. 2](image_url)  \hspace{1cm} FIG. 2: Using the Hartree energies from Table I, Eq. (31) is illustrated for $\gamma = Z'/Z$ and $Z = 1$. The trend is identical to that seen in Table I, however it is clear that the value is approaching it’s asymptote, 5/4. This is the Hartree energy for density consisting of the doubly occupied hydrogen 1s orbital.

![Fig. 3](image_url)  \hspace{1cm} FIG. 3: The Hartree-exchange energies reported in Table II are used to illustrate the inequalities of Eq. (30) with $\gamma = Z'/Z$ and $Z = 4$. Compared to Fig. 2, the value of $E_{\text{HX}}[n\gamma]/\gamma$ is not as fully converged to its asymptote, however the maximum value of $\gamma$ is 4 times smaller. The asymptotic value for this case is 586373/93312 = 6.284, which is found by doubly occupying both 1s and 2s hydrogenic orbitals with $Z = 4$ and calculating Hartree and exchange energies.
and find their behavior. The numerical corrections to our previous results are negligible.

V. CHARGE-NEUTRAL SCALING

In this section, we repeat all the logic of the previous section, but apply it now to CN scaling. After repeating similar steps (given in Appendix A), we arrive at the general result:

\[ \Delta F^\alpha[n_1^{1/\alpha}] \geq \Delta F^\alpha[n] \],

where

\[ \Delta F^\alpha[n] = F[n] - \alpha^{7/3} F[n_1/\alpha] \],

and \( \alpha = 1/\zeta \). Just as we did for coordinate scaling, we can refine our inequality substantially. By construction, \( \Delta F^\alpha[n] = 0 \) for \( F^{TF}[n] \), so we define the useful functional:

\[ F^{NT}[n] = F[n] - F^{TF}[n] \]

as the Non-Thomas-Fermi contribution to \( F[n] \). Our inequality then reads:

\[ \Delta F^{NT\alpha}[n_1^{1/\alpha}] \geq \Delta F^{NT\alpha}[n] \],

where

\[ \Delta F^{NT\alpha}[n] = F^{NT}[n] - \alpha^{7/3} F^{NT}[n_1/\alpha] \].

We find an interesting result in the limit \( \alpha \to 0 \), if we make the reasonable assumption that all non-Thomas-Fermi contributions scale less strongly than \( \zeta^{7/3} \):

\[ F^{NT}[n^{TF}] \geq F^{NT}[n] \],

as TF becomes relatively exact in the high \( \zeta \). This inequality is fiendishly hard to test, even in the large \( \zeta \) limit. Consider, e.g., the He atom. The corresponding TF density is well-known[33] but we would have to evaluate the exact interacting functional on it to find the non-TF contribution. All the above results also apply directly to non-interacting electrons in a potential, such as the Bohr atom[34], with \( F \) replaced by \( T_b \), and the TF contributions calculated with no Hartree term. But the same difficulties remain.

There is one case where we know enough already to test. For the hydrogen atom (or any one-electron system), \( F = T \) only, and is given by the von Weizacker functional. The TF density (with or without interaction) is well-known and singular at the origin, making the von Weizacker energy diverge. Thus, the formula is satisfied, but not very informative.

Lastly, we consider Thomas-Fermi-Dirac-Weizsäker theory[35] (TFDW). Here we add to TF the local exchange

\[ E_x^{(0)}[n] = A_x \int d^3 r \ n^{4/3}(r) \],

where \( A_x = -(3/4)(3/\pi)^{1/3} \), and the next order gradient correction to the kinetic energy,

\[ T^{(2)}_n[n] = \frac{1}{72} \int d^3 r \ \frac{\nabla n(r)^2}{n(r)}. \]

Both these terms scale the same way under CN density scaling, i.e.,

\[ F^{(2)}[n_\zeta] = T^{(2)}[n_\zeta] + E_x^{(0)}[n_\zeta] = \zeta^{5/3} (T^{(2)}[n] + E_x^{(0)}[n]). \]

Then can write the inequality as

\[ F^{(2)}[n_\zeta] \geq \zeta^{5/3} F^{(2)}[n], \]

where \( n(r) \) has been evaluated self-consistently within TFDW and \( \zeta \geq 1 \). Thus

\[ F^{(2)}[n^{TF}] \geq F^{(2)}[n^5]/\zeta^{5/3} \geq F^{(2)}[n], \]

where \( n^{TF}(r) \) is the Thomas-Fermi solution for the same potential as for \( n(r) \).

VI. CONCLUSION

Potential scaling, conjugate to a given density scaling, promises to be a useful tool in density functional theory. It leads to many exact conditions that can be used in functional construction. We have applied it to two distinct types of scaling: uniform coordinate scaling and charge neutral scaling. In both cases, we have found several interesting bounds. Uniform coordinate scaling was useful for analyzing Kohn-Sham DFT, leading to inequalities involving the only unknown in DFT, the exchange-correlation functional. The limit of this inequality involves evaluating the Hartree-exchange-correlation energy of the density of non-interacting fermions in the external potential. This connection between the interacting and non-interacting systems resonates with standard approaches in many-body perturbation theory. We illustrate the bounds on the Hartree-exchange energy this inequality provides by performing OEP exact exchange calculations on helium and beryllium, showing the approach to their asymptote. On the other hand, charge-neutral scaling provides inequalities involving Thomas-Fermi quantities. The Thomas-Fermi approximation becomes relatively exact for all electronic systems[28, 29] and these relations link the corrections to Thomas-Fermi with the true system, including those of TFDW theory. However evaluating the TF density within these theories often leads to divergences[36, 37]. It is not clear how important these relations will be for functional development, but those derived from uniform coordinate scaling are siblings to those derived by Levy and Perdew[5], which proved very useful in constraining approximations in DFT.

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APPENDIX A: CHARGE-NEUTRAL SCALING INEQUALITY

We follow the steps in deriving Eq. (20) but applied to the charge neutral scaling defined in Eq. (14). Taking \( n_{1/\zeta}^c(r) \) as a trial density for the \( v(r) \) system, then the variational principle states:

\[
F[n_{1/\zeta}^c] + V[n_{1/\zeta}^c] \geq F[n] + V[n] ,
\]

(A1)

where \( n^c(r) \) is the exact density for the scaled potential \( v^c(r) \). Conversely, use \( n_{c}(r) \) as a trial density for the exact potential \( v^c(r) \) system:

\[
F[n_{c}] + \zeta^{7/3}V[n] \geq F[n^c] + \zeta^{7/3}V[n_{1/\zeta}^c] ,
\]

where we have used \( V^c[n] = \zeta^{7/3}V[n] \). Combining these inequalities gives:

\[
\frac{F[n_{c}] - F[n]}{\zeta^{7/3}} \geq \frac{F[n] - F[n_{1/\zeta}^c]}{\zeta^{7/3}} ,
\]

(A3)

which may be written as

\[
\Delta F^c[n_{1/\zeta}^c] \geq \Delta F^c[n] ,
\]

(A4)

with

\[
\Delta F^c[n] = F[n] - \frac{F[n_{c}]}{\zeta^{7/3}} .
\]

(A5)

APPENDIX B: EXCHANGE ENERGY FOR NON-INTERACTING BERyllIUM

The limit of the inequality, Eq. (28), is the Hartree-exchange functional evaluated on the density of the corresponding non-interacting system. Since the g.s. orbitals which sum to this density are known analytically (they are simply hydrogenic orbitals), we may calculate the exact Hartree-exchange value.

Written in spherical coordinates, \( r = |r| \), the 1s and 2s hydrogenic orbitals are:

\[
\phi_{1s}(r) = \left( \frac{Z^3}{\pi} \right)^{1/2} e^{-Zr} ,
\]

(B1)

\[
\phi_{2s}(r) = \left( \frac{Z^3}{32\pi} \right)^{1/2} (2 - Zr)e^{-Zr/2} .
\]

(B2)

For beryllium, both these orbitals are doubly occupied, giving the total density as

\[
n(r) = 2|\phi_{1s}(r)|^2 + 2|\phi_{2s}(r)|^2 .
\]

(B3)

The Hartree energy is defined by Eq. (6), however in the special case of spherical densities it may be written as:

\[
U[n] = \frac{1}{2} \int_0^\infty dr \ (f[n](r))^2 ,
\]

(B4)

where

\[
f[n](r) = 4\pi \int_r^\infty dr' r'n(r') ,
\]

(B5)

and we use square brackets to indicate that it is a functional of the density. The exchange energy for a spin-unpolarized system is:

\[
E_x = -2\frac{1}{\zeta} \sum_{i,j} \int d^3r \int d^3r' \frac{\phi_i^*(r)\phi_j^*(r')\phi_j(r)\phi_i(r')}{|r - r'|} ,
\]

(B6)

where the factor 2 is due to spin, and the sum is over occupied orbitals only, in this case 1s and 2s. If we define a new quantity, \( \tilde{n}(r) \):

\[
\tilde{n}(r) = \phi_{1s}(r)\phi_{2s}(r) ,
\]

(B7)

then we may write

\[
E_x = -2 \left(U[n_{1s}] + 2U[\tilde{n}] + U[n_{2s}]\right) .
\]

(B8)

We can use Eq. (B5) and Eq. (B4) for each term separately and then combine to find the total exchange energy. The answer will be equivalent to solving Eq. (B6) using the orbitals, however this method avoids performing integrals involving \( 1/|r - r'| \) and, in this case, are easy to solve using integration by parts. The values for Hartree, exchange and their sum are:

\[
U[n] = \frac{5 \times 23 \times 431}{2^{63}4} = \frac{49565}{5184} = 9.561 ,
\]

\[
E_x = \frac{59 \times 71 \times 73}{2^{36}} = \frac{305797}{93312} = -3.277 ,
\]

\[
E_{nx} = \frac{383 \times 1531}{2^{36}} = \frac{586373}{93312} = 6.284 .
\]

[1] A Primer in Density Functional Theory, ed. C. Fiolhais, F. Nogueira, and M. Marques (Springer-Verlag, NY,
The surface energy of a bounded electron gas

Self-consistent equations including exchange and correlation effects, W. Kohn and L.J. Sham, Phys. Rev. 140, A1133 (1965).

Which functional should I choose?, D. Rappoport, N. Crawford, F. Furche, and K. Burke, to appear in Encyclopedia of Inorganic Chemistry: Computational Inorganic and Bioinorganic Chemistry, Wiley (2009).

Hellmann-Feynman, virial, and scaling requisites for the exact universal density functionals. Shape of the correlation potential and diamagnetic susceptibility for atoms, M. Levy and J.P. Perdew, Phys. Rev. A 32, 2010 (1985).

Density-functional exchange-correlation through coordinate scaling in adiabatic connection and correlation hole, M. Levy, Phys. Rev. A 43, 4637 (1991).

Tight bound and convexity constraint on the exchange-correlation-energy functional in the low-density limit, and other formal tests of generalized-gradient approximations, M. Levy and J.P. Perdew, Phys. Rev. B 48, 11638 (1993); ibid., 55, 13321 (1997)(E).

Nonuniform coordinate scaling requirements for exchange-correlation energy, M. Levy and H. Ou-Yang, Phys. Rev. A 42, 651 (1990).

Exchange-correlation energy of a metallic surface: Wavevector analysis, D.C. Langreth and J.P. Perdew, Phys. Rev. B 15, 2884 (1977).

Theory of nonuniform electronic systems. I. Analysis of the gradient approximation and a generalization that works, D.C. Langreth and J.P. Perdew, Phys. Rev. B 21, 5469 (1980).

The surface energy of a bounded electron gas, J. Harris and R.O. Jones, J. Phys. F 4, 1170 (1974).

Exchange and correlation in atoms, molecules, and solids by the spin-density-functional formalism, O. Gunnarsson and B.I. Lundqvist, Phys. Rev. B 13, 4274 (1976).

Some remarks on scaling relations in density functional theory, W. Yang, in Density Matrices and Density Functionals, eds. R. Erdahl and V. H. Smith, Jr. (D. Reidel Publishing Company, Dordrecht, The Netherlands, 1987), pp 499-506.

A new functional with homogeneous coordinate scaling in density functional theory: \( F[p,\lambda] \), M. Levy, W. Yang, and R.G. Parr, J. Chem. Phys. 83, 2334 (1985).

Rationale for mixing exact exchange with density functional approximations, J.P. Perdew, M. Ernzerhof, and K. Burke, J. Chem. Phys. 105, 9982 (1996).

The adiabatic connection method: A non-empirical hybrid, K. Burke, M. Ernzerhof, and J.P. Perdew, Chem. Phys. Lett. 265, 115 (1997).

Mixing exact exchange with GGA: When to say when, K. Burke, J.P. Perdew, and M. Ernzerhof, in Electronic Density Functional Theory: Recent Progress and New Directions, eds. J.F. Dobson, G. Vignale, and M.P. Das (Plenum, NY, 1998), pp 57-68.

Construction of the adiabatic connection, M. Ernzerhof, Chem. Phys. Lett. 263, 499 (1996).

Generalized gradient approximation for solids and their surfaces, J. Perdew, A. Ruzsinszky, G.I. Csonka, O.A. Vydrov, G.E. Scuseria, L.A. Constantin, X. Zhou, and K. Burke, Phys. Rev. Lett. 100, 136406 (2008).

Relevance of the slowly-varying electron gas to atoms, molecules, and solids, J. P. Perdew, L. A. Constantin, E. Sagvolden, and K. Burke, Phys. Rev. Lett. 97, 223002 (2006).

Potential Functionals: Dual to Density Functionals and Solution to the \( v \)-Representability Problem, W. Yang, P.W. Ayers, and Q. Wu, Phys. Rev. Lett. 92, 146404 (2004).

Universal variational functionals of electron densities, first order density matrices, and natural spin-orbitals and solution of the \( v \)-representability problem, M. Levy, Proc. Natl. Acad. Sci. (U.S.A.) 76, 6062 (1979).

The calculation of atomic fields, L.H. Thomas, Proc. Camb. Phil. Soc. 23, 542 (1926).

A statistical method for the determination of some atomic properties and the application of this method to the theory of the periodic system of elements, E. Fermi, Zeit. Fur Physik 48, 73 (1928).

Adiabatic connection from accurate wavefunction calculations, D. Frydel, W. Terilla, and K. Burke, J. Chem. Phys. 112, 5292 (2000).

Comparison of approximate and exact density functionals: A quantum monte carlo study, C.J. Umrigar and X. Gonze, in High Performance Computing and its Application to the Physical Sciences, Proceedings of the Mardi Gras 1993 Conference, edited by D. A. Browne et al. (World Scientific, Singapore, 1993).

Accurate exchange-correlation potentials and total-energy components for the helium isoelectronic series, C.J. Umrigar and X. Gonze, Phys. Rev. A 50, 3827 (1994).

Thomas-Fermi Theory Revisited, E.H. Lieb and B. Simon, Phys. Rev. Lett. 31, 681 (1973).

Thomas-fermi and related theories of atoms and molecules, E.H. Lieb, Rev. Mod. Phys. 53, 603 (1981).

Semiclassical Origins of Density Functionals, P. Elliott, D. Lee, A. Cangi, and K. Burke, Phys. Rev. Lett. 100, 256406 (2008).

Local correlation energies of two-electron atoms and model systems, C.J. Huang and C.J. Umrigar, Phys. Rev. A 56, 290 (1997).

OPMKS: atomic DFT program, University of Frankfurt, Germany, E. Engel

Condition on the Kohn-Sham kinetic energy, and modern parametrization of the Thomas-Fermi density, D. Lee, L. A. Constantin, J. P. Perdew, K. Burke, J. Chem. Phys. 130, 034107 (2009).

Semiclassical theory of atoms, B.-G. Englert, Lect. Notes Phys. 300 (1988).

Quantum corrections to the Thomas-Fermi equation, D.A. Kirzhnits, Sov. Phys. JETP 5, 64 (1957).

Sixth-order term of the gradient expansion of the kinetic-energy density functional, D. R. Murphy, Phys. Rev. A 24, 1682 (1981).

Numerical test of the sixth-order gradient expansion for the kinetic energy: Application to the monovacancy in jellium, Z. D. Yan, J. P. Perdew, T. Korhonen, and P. Ziesche, Phys. Rev. A 55, 4601 (1997).