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Published in: Journal of Chemical Physics

DOI: 10.1063/1.5000908

Published: 21/12/2017

Please cite the original version:
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Citation: The Journal of Chemical Physics 147, 234102 (2017);
View online: https://doi.org/10.1063/1.5000908
View Table of Contents: http://aip.scitation.org/toc/jcp/147/23
Published by the American Institute of Physics

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Self-consistent assessment of Englert-Schwinger model on atomic properties

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(Received 19 August 2017; accepted 16 November 2017; published online 18 December 2017)

Our manuscript investigates a self-consistent solution of the statistical atom model proposed by Berthold-Georg Englert and Julian Schwinger (the ES model) and benchmarks it against atomic Kohn-Sham and two orbital-free models of the Thomas-Fermi-Dirac (TFD)-vW family. Results show that the ES model generally offers the same accuracy as the well-known TFD-\(\frac{1}{2}\)vW model; however, the ES model corrects the failure in the Pauli potential near-nucleus region. We also point to the inability of describing low-Z atoms as the foremost concern in improving the present model. © 2017 Author(s).

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I. INTRODUCTION

Orbital-free density functional theory (OFDFT) is a lucrative path to linear scaling DFT methods. The modern OFDFT theory is mostly investigated in the setting of non-local or generalized gradient approximation (GGA)-style kinetic functionals. Potential functionals, such as those used in Berthold-Georg Englert and Julian Schwinger’s statistical atom model,1–3 have fallen out of favor. Yet, it has been shown that at least GGA-style orbital-free functionals usually suffer from a theoretical flaw: the atomic nucleus is not well described, as evidenced by the singular Pauli potential.4–6 The Pauli potential describes a positive effective repulsion arising from the Pauli exclusion principle, and its violation implies a fundamentally wrong solution of the electronic problem. Potential functionals allow linear-scaling DFT to sidestep this flaw, and our intention here is to quantify the potential functionals’ self-consistent accuracy by benchmarking Englert and Schwinger’s model2,3 to more well-known OFDFT models.

A flurry of recent activity uses potential as a variable instead of density. Much of the work is focused especially on the Pauli potentials’ role and especially in atoms.5,7,8 The potential variable has even been combined with machine learning,9 which also has had some applications for the search of kinetic energy functionals.10,11

Pseudopotentials5,12 have been useful in conventional approaches to extended systems with OFDFT, where an active effort to improve local pseudopotentials’ transferability is ongoing.13,14 While this solution works remarkably well,12 a few questions remain for systematic improvement. In general, how much of the overall accuracy is because of the better ion-energy interaction description or the better electronic kinetic energy functional? Even more importantly, in the case of fitted pseudopotentials, when is a pseudopotential overfitted compared to the kinetic energy functional? One approach is to investigate the full potential solution to OFDFT equations, as we did by using the projector-augmented-wave method,15 but one of the problems here is the near-nucleus singularity of the Pauli potential, which brings the solution’s plausibility into question.

The problem near the atomic nucleus is not a new one, and it motivated the development of the statistical atom model.1 Englert and Schwinger’s approach, however, is based on the effective potential instead of the density, which allows natural energy scale separation of the problematic core electrons from the bulk. In particular, we are interested in the model proposed in two earlier papers2,3 and later refined in Englert’s book, Semiclassical Theory of Atoms.16

The perturbative results from the statistical atom are of amazing accuracy, but because our focus is on extended systems, self-consistent solutions of atoms are more interesting to us. We therefore explore a self-consistent solution to Englert and Schwinger’s model and assess its quality by comparing it to two other representative orbital-free models that belong to the Thomas-Fermi-Dirac (TFD)-vW family of orbital-free models. Results show that the Berthold-Georg Englert and Julian Schwinger (ES) model generally offers the same accuracy as the well-known TFD-\(\frac{1}{2}\)vW model, but it works better for few quantities, especially for the Pauli potential. We also discerned that the model’s current limitation is the inability to model atoms with a low atomic number (<12).

II. THE MODEL

We briefly present the potential functional formalism in DFT and show how it can be used to correct the Thomas-Fermi description of atoms.
A. Formalism

We start with the usual DFT formalism, where energy is separated as

$$E[n] = E_{\text{kin}}[n] + E_{\text{int}}[n] + \int d\mathbf{r} V_{\text{ext}}(\mathbf{r}) n(\mathbf{r}),$$

where $E_{\text{int}}$ is the functional containing electron–electron interaction. We enforce the particle number $N$ restriction via Lagrange multiplier $\zeta$ and apply the Legendre transformation to the kinetic energy

$$E_i[V + \zeta] = E_{\text{kin}}[n] + \int d\mathbf{r} (V(\mathbf{r}) + \zeta) n(\mathbf{r}),$$

where we introduced an effective single particle potential $V$.

By incorporating this into functional (1), we arrive at a joint functional where the variables $n$, $V$, and $\zeta$ are treated as independent variables,

$$E[V, n, \zeta] = E_i[V + \zeta] - \int (d\mathbf{r}) V(\mathbf{r}) n(\mathbf{r}) + E_{\text{int}}[n] - \zeta N.$$

The variations of this functional lead us to the relations

$$n(\mathbf{r}) = \frac{\delta}{\delta V} E_i[V + \zeta],$$

$$N = \frac{\delta}{\delta \zeta} E_i[V + \zeta] = \int d\mathbf{r} V(\mathbf{r}),$$

$$V = V_{\text{ext}} + \frac{\delta}{\delta n} E_{\text{int}}[n] = V_{\text{ext}} + V_{\text{int}},$$

where $V_{\text{int}}$ is the single particle interaction potential. Here we use the binding energy $\zeta$ instead of the more common chemical potential $\mu = -\zeta$. The energy $E_i[V + \zeta]$ is possible to approximate semiclassically.\footnote{We emphasize here that working with this joint functional is still working within the DFT framework given by the Hohenberg–Kohn theorems, although in this formalism, we acknowledge the effective potential and chemical potential as independent variables. For simple cases such as the Thomas–Fermi theory, it is possible to reduce the potential functional version of $E_i[V + \zeta]$ to density functionals and vice versa.\footnote{Because we are working with a single particle potential and $E_i$ will be approximated with the help of a single particle potential, the kinetic energy in this model is non-interacting kinetic energy, which Kohn–Sham and other OFDFT schemes also use.}}

B. Semiclassical $E_1$

Englert and Schwinger derived a semiclassical Airy-average expression for $E_i$; this can be understood as a Thomas–Fermi expression that contains quantum corrections. The semiclassical expression is given in terms of Airy-average functions $F_m$, which are closely related to Airy functions.

The derivation is based on the approximation that each electron moves in a local harmonic potential. Then the time propagator of the harmonic oscillator is used to obtain the quantum corrected Thomas–Fermi expression.\footnote{The main quantity in the quantum-corrected theory is}

$$y = 2(V + \zeta)/2\nabla V|^{-2/3},$$

and the Airy-average functions are functions of this variable

$$F_0(y) = [\text{Ai}(y)]^2,$$

$$F_1(y) = -y \text{Ai}(y) + [\text{Ai}'(y)]^2,$$

$$F_{-1}(y) = -2\text{Ai}(y)\text{Ai}'(y).$$

We can obtain the remaining Airy-average functions recursively with the rule

$$(m - \frac{1}{2})F_m(y) = \frac{1}{4}F_{m-3}(y) - yF_{m-1}(y).$$

The quantum-corrected Thomas–Fermi expression for $E_i[V + \zeta]$ in terms of Airy-average functions is

$$E_i[V + \zeta] = -\frac{1}{4\pi} \int d\mathbf{r} \left[ |2\nabla V|^{3/2} F_2(y) \right.$$

$$\left. - \frac{1}{3} \nabla V |2\nabla V|^{1/2} F_1(y) \right],$$

with approximations to the second order in potential $O(\nabla^2 V)$. Reducing this to a kinetic energy functional\footnote{This is the main motivation for the joint functional treatment, as similar treatment is unavailable in terms of density functionals.} results in a Thomas–Fermi kinetic energy functional plus one-ninth of the von Weizsäcker kinetic energy functional, which is more widely known as the gradient-corrected Thomas–Fermi.

C. Partition of $E_i[V + \zeta]$

We note that the semiclassical evaluation is not valid near the atomic nucleus, which is why it is reasonable to split the evaluation of energy into two parts: Electrons described well by the Thomas–Fermi theory and strongly bound electrons (SBEs) that are better described as hydrogenic states.\footnote{Energy $E_1$ is the trace over the non-interacting atomic Hamiltonian}

$$E_i[V + \zeta] = \text{Tr}(H + \zeta)\eta(-H - \zeta),$$

where the Hamiltonian is $H = \frac{1}{2}\nabla^2 + V(\mathbf{r})$ and $\eta$ is the Heaviside function. We can split this into two parts using an energy scale given by $\zeta_s$, so that the electrons with energy levels above $\zeta_s$ are treated semiclassically and electrons with energy below $\zeta_s$ are treated with discrete quantum states. We accomplish this by adding an intelligent zero based on $\zeta_s$. This results in

$$E_1 = \frac{E_1[V + \zeta] - E_1[V + \zeta_s] + (\zeta - \zeta_s)N[V + \zeta_s] + ES}{E_{\zeta_s}},$$

where the functional $E_{\zeta_s}$ is approximated semiclassically and $E_S$ is evaluated with some other quantum mechanical method. Formally this can be thought of as the evaluation of the semiclassical approximation from which we subtract semiclassical evaluation on energy scales below $\zeta_s$ and finally adding the correct treatment for electrons below energy $\zeta_s$.

The critical part of the ES model is to treat the exact part $E_S$ as hydrogenic states. It is assumed that the potential near the nucleus is quite close to the Coulombic potential $-\frac{Z}{r}$ so that perturbative evaluation of the states is accurate enough,

$$E_S = Z^2 n_s + \int d\mathbf{r} (V + \frac{Z}{r}) n_{\text{SBE}},$$

where $n_s$ is the uppermost electron shell treated with hydrogenic states and $n_{\text{SBE}}$ is the density of the electrons treated...
with hydrogenic states. By shell, we mean the collection of all hydrogenic states with same energy; thus the shells are tabulated by the quantum number \( n \). The density \( n_{\text{SBE}} \) is obtained from spherically averaged hydrogenic states,

\[
n_{\text{SBE}} = \sum_{i \leq n} |\psi_i|_{\text{av}}^2,
\]

where \( |\psi_i|_{\text{av}} \) is the spherically averaged wavefunction of \( i \)th hydrogenic shell. For justification and details, see Refs. 1 and 19. Originally, Englert and Schwinger investigated corrections to the Thomas-Fermi model—and as they pointed out, the cutoff energy \( \zeta \) has certain ambiguity because we are trying to patch a continuous semiclassical model with a discrete quantum model. The most consistent way to achieve this for the Thomas-Fermi model is to average over two electronic shells with equal weights on an energy scale. This is also a good choice for the Thomas-Fermi model with corrections used here.

More concretely, this means

\[
E_{\zeta\zeta_i} \longrightarrow \sum_{i=1}^{2} \frac{1}{2} E_{\zeta_i},
\]

where \( \zeta_i \) is the corrected binding energy of \( i \)th electronic shell,

\[
\zeta_i = \frac{Z^2}{2n_i^2} - \int dr (V + \frac{Z}{r}) |\psi_i|_{\text{av}}^2.
\]

For practical purposes, these corrections can be embedded inside the Airy-average functions \( F_m \); thus they become corrected \( F_m \). The correction induces their own \( \zeta \) variables,

\[
y_i = 2(V + \zeta_i)|2\nabla V|^{-2/3},
\]

and by doing replacement,

\[
F_m \longrightarrow \sum_{i=1}^{2} \frac{1}{2} w_j \left[ F_m(y_i) - (y_j - y)F_{m-1} \right],
\]

we can use the energy expression (5). The corrections for strongly bound electrons are inside the functions \( F_m \).

Our numerical results indicate that for all atoms of the relevant size (\( Z < 100 \)), the only energetically believable correction is the one where only the lowest shell (i.e., 1s electrons) is treated exactly, and average occurs over shells 1 and 2. This approximation naturally breaks down when the number of total electrons is low, namely, \( Z \sim 12 \).

D. Semiclassical density

The density can be calculated via the relation \( \frac{\delta}{\delta V} E_1 \) (2). The resulting density splits straightforwardly into two contributions,

\[
n = \frac{\delta}{\delta V} E_1
\]

\[
= \left( \frac{\partial e_1}{\partial V} - \nabla^2 V \frac{\partial e_1}{\partial V} \nabla^2 V \right) \frac{\delta \zeta}{\delta V} \frac{\partial}{\partial \zeta} E_1 + n_{\text{SBE}},
\]

where \( e_1 \) is the energy density of \( E_1 \), \( \bar{n} \) is the semiclassical contribution, and the two last terms are the contribution of the strongly bound electrons. Note that the contribution from strongly bound electrons contains the derivative with respect to the core binding energy \( \zeta \), because it depends on the potential via (6). The derivative is

\[
\frac{\delta \zeta}{\delta V} \frac{\partial}{\partial \zeta} \left[ \frac{1}{2} \int d\mathbf{r} |2\nabla V|^{1/3} F_1(y_i) - \frac{1}{3} |2V|^2 V F_{-1} \right] |\psi_i|_{\text{av}}^2,
\]

so there will be contributions from the hydrogenic states that depend on the potential.

The semiclassical contribution \( \bar{n} \) in spherical symmetry is

\[
\bar{n} = \frac{1}{2\pi} \left[ 2\nabla|F_2| - \frac{1}{12\pi} |2\nabla|^{1/3} \nabla^2 V F_0 \right.
\]

\[
+ \frac{1}{r^2} \left[ \frac{1}{36\pi} |2\nabla|^{2/3} F_2 - \frac{1}{9\pi} |2\nabla|^{-2/3} \nabla^2 V F_{-3} \right]
\]

\[
- \frac{1}{108\pi} |2\nabla|^{-2/3} \nabla^2 V F_{-3}
\]

\[
+ \frac{1}{r^2} \left[ \frac{1}{36\pi} |2\nabla|^{1/3} F_2 - \frac{1}{108\pi} |2\nabla|^{1/3} F_{-3} \right],
\]

where the first term \( \frac{1}{2\pi}(2\nabla|F_2| \) contains the Thomas-Fermi limit. The expression is valid only for neutral atoms and positive ions as it assumes that the potential’s gradient is positive. Theoretically, using a simpler density expression is also a viable option, which is only the first two terms of (8). For our benchmarking, we use only the aforementioned density,

E. Interaction

So far, we have detailed how to calculate \( E_1 \) and electronic density, which are independent of any possible interaction. Now we detail the electronic interaction included via the effective potential \( V \).

The interaction term \( E_{\text{int}} \) is separated into an electrostatic term (Hartree) and into an exchange term,

\[
E_{\text{int}} = E_{\text{H}} + E_{\text{ex}}.
\]

The corresponding interaction potential is then

\[
\delta E_{\text{int}} = V_{\text{H}} + V_{\text{ex}}.
\]
The Hartree term is well known,

\[ V_H = \int d\mathbf{r}' \frac{n(\mathbf{r})}{|\mathbf{r} - \mathbf{r}'|}, \]

and its connection to density will be used to achieve a self-consistent solution. We do not add correlation effects to the model as the effect of correlation in atoms is below the accuracy of semiclassical \( E_1 \).

Before the model is complete, we must comment on how to add the exchange effects. The aim is to include local exchange effects, as described by the Dirac exchange functional,

\[ E_{\text{ex}} = -\frac{1}{4\pi^3} \int d\mathbf{r} (3\pi^2 n(\mathbf{r}))^{4/3}. \]  

(9)

To be consistent, we discard exchange effects on strongly bound electrons because we already approximated them to be non-interacting (although they will receive a marginal contribution via the potential). The exchange potential is necessary for self-consistency, given by

\[ V_{\text{ex}} = \delta \frac{\partial}{\partial n} E_{\text{ex}} = -n (3\pi^2 n(\mathbf{r}))^{1/3}. \]  

(10)

The most straightforward way to add exchange effects is simply by replacing \( n \rightarrow \bar{n} \) so that only the smooth part is treated with exchange. Testing showed us that while this method produced exchange effects of the correct magnitude, it is not the most accurate way to include the exchange (while discarding contributions of the strongly bound electrons).

The second route taken by Englert and Schwinger is to start from the Thomas-Fermi theory to arrive at a potential description of the exchange. At the Thomas-Fermi level, the exchange potential can be calculated from \( V_{\text{ex}} = \pi \frac{\partial}{\partial V} n \), and the exchange energy density \( \epsilon_{\text{ex}} \) can be calculated from the relation \( \frac{\partial V_{\text{ex}}}{\partial V} = \frac{1}{2\pi} |2\nabla V|^{4/3} \). To include exchange effects on the Thomas-Fermi level, the obvious choice for \( n \) here is the Thomas-Fermi part of the density \( \frac{1}{2\pi} |2\nabla V| F_2 \), resulting in the potential

\[ V_{\text{ex}} = -|2\nabla V|^{1/3} F_1, \]  

(11)

and the exchange energy

\[ E_{\text{ex}} = \int d\mathbf{r} \frac{1}{2\pi} |2\nabla V|^{4/3} \left( \frac{1}{4} F_1 F_{-1} + \frac{1}{8} F_0^2 + \frac{1}{2} y F_1^2 \right). \]

We found the description of the exchange potential inadequate at larger distances in the atoms. Thus, we also use another version of the exchange potential, which uses the density expression \( \bar{n} \) where the Laplacian of the potential has been approximated out. The resulting exchange potential is

\[ V_{\text{ex}} = -|2\nabla V|^{1/3} (F_1 - \frac{1}{6} F_{-2}), \]  

(12)

which we will use in our implementation. We expect the previous exchange energy description to be accurate enough for this potential. Later we discuss different exchange approximations that are meaningful.

### III. IMPLEMENTATION

We obtained the self-consistent solution via the relation \( n = \frac{\partial}{\partial V} E_1 \) and the connection of a single particle potential \( V \) to the electrostatic potential \( V_H \). For a given effective potential \( V \), we find the corresponding binding energy \( \zeta \) with the relation \( N = \int d\mathbf{r} n[V(\mathbf{r})] + \zeta \). This yields the density \( n[V(\mathbf{r})] + \zeta \), which we can use to find a new Hartree potential \( V_H \) through a Poisson equation. The new effective potential is now obtained from this Hartree potential with \( V = V_H + V_{\text{ext}} + V_{\text{ex}}[n, V] \), where in \( V_{\text{ex}} \), we use the old potential and density. This is iterated until the change in the potential and binding energy is small enough. With a good initial guess, this procedure gives the self-consistent solution. Here the Thomas-Fermi potential is a sufficient initial guess.

We describe this idea in a bit more detail for spherically symmetrical atoms. With relation (4) and the connection of an electrostatic potential \( V_H \) with a Poisson equation, we arrive at

\[ -\frac{1}{4\pi} \nabla^2 \left( V + \frac{Z}{r} \right) = n(\mathbf{r}) - \nabla^2 V_{\text{ex}}, \]

with boundary conditions \( rV(\mathbf{r} \rightarrow 0) \rightarrow -Z \) and \( rV(\mathbf{r} \rightarrow \infty) \rightarrow -Z + N \), where zero potential has been assigned to infinitely far away from the nucleus. After we use the fact that our system is spherically symmetric and introduce the auxiliary quantity \( V(\mathbf{r}) = -\frac{Z}{r} \Phi(r) \), we have the differential equation

\[ Z \frac{\partial^2}{\partial r^2} \Phi(r) = 4\pi r n(r) - \frac{\partial^2}{\partial r^2} r V_{\text{ex}}, \]  

with boundary conditions

\[ \Phi(0) = 1, \]

\[ \Phi(\infty) = 1 - \frac{N}{Z}. \]

After obtaining the effective potential \( V \), we find the corresponding binding energy with Newton’s method from the relation \( N - \int d\mathbf{r} n(V + \zeta) = 0 \). We use a non-uniform grid that is denser near the nucleus, where the change in values is greater than in the tail (the grid is correspondingly sparser in the tail).

### A. Numerical method

Englert and Schwinger’s original paper uses the shooting method to solve the resulting differential equation. We solve the resulting differential with a simple 1D finite element method with linear elements. For the finite element method, we derive the weak form of the differential equation which is

\[ -Z \int dr \frac{\partial \Phi(r)}{\partial r} \cdot \frac{\partial v}{\partial r} = \int dr 4\pi r n(r) \cdot v + \frac{\partial}{\partial r} r V_{\text{ex}}(r) \cdot \frac{\partial v}{\partial r}, \]

where \( v \) is the test function. During testing, we noted that the exchange potential is the most sensitive quantity from a numerical point of view. In the numerical study, the solution for neutral atoms was not obtained because the effective potential’s long-range behavior is unknown.

We obtain the neutral atom solution simply by setting \( N = \int d\mathbf{r} n(\mathbf{r}) + \zeta \rightarrow 0 \), and then we converge the result with respect to grid size until the errors (due to the grid’s finite size) are below the error threshold.

The differential equation is a bit different than Englert and Schwinger’s book. We use the more straightforward expression derived from a Poisson equation than they mention in the...
book, where all the terms containing the Laplacian of the effective potential are moved to the left side of the equation.

**IV. RESULTS**

We benchmark the numerical results against perturbative results calculated by Englert and Schwinger, the Kohn-Sham results, and the TFD-1/W model. First we evaluate our results against Englert and Schwinger’s and then compare the self-consistent results for the following methods:

- **ES**: The Englert-Schwinger model with exchange potential (12) and density expression (8).
- **KS-LDA**: Spherically symmetric Kohn-Sham atom, where the xc-functional is the Dirac exchange (9) and no correlation (KS-LDA stands for Kohn-Sham local density approximation).
- **TFD-1/W**: The OFDFT model, which contains Dirac exchange and Thomas-Fermi plus one-ninth of a von Weizsäcker term as a kinetic energy functional, where the von Weizsäcker factor is derived as a quantum correction to the Thomas-Fermi functional.
- **TFD-1/2/W**: The OFDFT model, which contains the Dirac exchange and Thomas-Fermi plus one-fifth of a von Weizsäcker term as the kinetic energy functional, where the von Weizsäcker factor is fitted rather than derived.

All of these models are based on DFT. In each model, we treat the exchange effect with the Dirac exchange (9). Thus we choose to benchmark against Kohn-Sham as the most accurate of the approximations. Originally, Englert and Schwinger opted to benchmark against Hartree–Fock data.

The exact difference between the orbital-free models is a bit more complex. Both orbital-free and ES models are (in a sense) approximating the energy $E_1$. The ES model approximates it directly, including both kinetic and potential terms, while TFD-1/W models approximate it via approximating the kinetic energy density functional only. From a formal point of view, the TFD-1/2/W and the ES models both expand the semiclassical trace to the same order in $\hbar$, but we must remember that the TFD-1/2/W model disregards the strongly bound electron correction completely. In model TFD-1/2/W, the von Weizsäcker fraction is fitted to produce best results for atoms.

The atomic Kohn-Sham solver used is available in a grid-based implementation of the projector-augmented waves (GPAW) DFT package. The kinetic energy density functional methods are implemented within this same Kohn-Sham solver. The implementation details are described elsewhere.

**A. Comparison to Englert-Schwinger reference numerical data**

We first study the model presented in the book *Semiclassical Theory of Atoms*. Thus, we use density expression (8) and exchange potential (11) and solve the resulting differential equation.

We compare the numerical results against the ones provided in Ref. 16 in Table I for the krypton electronic configuration. The numbers correspond to fair accuracy. The biggest error is the binding energy of $Z = 38$, charge = 2 where we have a deviation of 3.3%. The deviation in other values is below 2%. Core binding energies $\zeta_1$ and $\zeta_2$ show strong similarity, which is to be expected as the potential should have a $-\frac{Z}{r}$ shape near the hydrogenic states. The neutral atom results by Englert and Schwinger are the result of extrapolation, so they are omitted from the comparison.

**B. Effect of exchange potential**

The exchange potential and energy have few possible approximations, as indicated in Sec. II E. To choose the best one, we take a look at krypton to see the effects of different exchange functionals. We look at the total energy, binding energy $\zeta$, and averages over densities $\langle \frac{1}{r^2} \rangle$, $\langle r \rangle$, and $\bar{r}^2$. The last one is defined by

$$\bar{r}^2 = \frac{1}{N} \int d\mathbf{r} \, r^2 \rho(\mathbf{r}).$$

The results for different exchange expressions for neutral krypton are tabulated in Table II.

From the change $\bar{r}^2$, we can see that the choice of exchange potential has a strong effect near the atom’s edge, which is quite natural. As the exchange energy is negative and the corresponding potential is attractive, we expect the correct inclusion of exchange to make the atomic size smaller.

The energy difference is not that useful on a semiclassical scale if we compare it to a highly accurate Englert-Schwinger prediction for semiclassical energy, which is $-2747.64$ hartrees for krypton. The deviation from this value is 0.7%, 0.1%, and 0.2% for methods (10), (11), and (12), respectively. Our choice, then, should be based on the quality of density at the atom’s outer reaches. The experimental value for $\bar{r}^2$ can be obtained via diamagnetic susceptibilities, which are provided by the CRC Handbook of Chemistry and Physics. The experimental value of $\bar{r}^2$ for krypton is 1.010 bohrs$^2$. This indicates that all

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</tr>
</tbody>
</table>
methods are a bit insufficient but that (12) is clearly the best for $\tau^2$. Finally, we want to note that there is a deviation of 0.05–0.1 bohr$^2$ to the reported $\tau^2$ values so that the deviation to the experimental value might already be at the level of precision of the chosen methods.

We therefore determined that for the self-consistent ES atom model, the exchange potential (12) is preferred. Still, the model’s accuracy in the atom’s outer reaches is somewhat limited by the exchange approximation, and it remains unclear how the exchange effect should be treated for strongly bound electrons. We intend to address this point in future work.

In the following, we use the exchange potential (12).

C. Assessment of ES improvements in neutral atoms

We compute the energies and a few other descriptive quantities for the ES model and compare them to other models for neutral systems to assess the quality of the self-consistent solution. The total energy as a quantity is not so important, as the real predictive power lies in the energy differences but having this information is somewhat useful because it informs the quality of approximations made. As previously mentioned, though, here we consider Kohn-Sham as the “ground truth.”

Most of our quantities depend on the density to indicate the shape and quality. The near-nucleus area is probed by averaging $\langle 1/r \rangle$, which is related to the shielding of the nuclear magnetic moment.$^{16}$ The average over $r$ is related to electric polarizability.$^{16}$ Finally, as we mentioned in Sec. IV B, we measure $\tau^2$, which probes the density at atoms’ outer edges and is related to diamagnetic susceptibility.

For atoms, the energy difference is tested only by calculating the ionization potential and comparing the total energies of charged and neutral systems. As the semiclassical model incorporates no information of the valence electron shells, our primary interest is in the ionization potential of alkali metals, which have a closed-shell structure. In xenon ($Z = 54$), we see that the experimental value is closer to the semiclassical value than the KS-LDA. The ES model is actually in better agreement with the experimental values as it assigns smaller sizes for almost all atoms (see supplementary material). The effect is mostly due to the exchange term as seen from Table II. The origin of these shell oscillations in the Kohn-Sham model is obvious: some outer orbitals are more delocalized than others.

We can see next in the comparison of different atomic models the absolute value of the numbers presented as general trends in Figs. 1–4. We feel it is informative enough to focus on three representative closed-shell systems: argon

\begin{table}[h]
\centering
\caption{Krypton $Z = 36$ with different exchange potentials with full density expression (8). Classical radius is defined by $V(r_{\text{classical}}) + \zeta = 0$.}
\begin{tabular}{|c|c|c|c|c|c|}
\hline
Exchange potential $V_{\text{ex}}$ & $E$ & $\zeta$ & $\langle \frac{1}{r} \rangle$ & $\langle r \rangle$ & $\tau^2$ & $r_{\text{classical}}$ \\
\hline
(10) & $-\frac{1}{2}(3\pi^2/\hbar^2)^{1/3}$ & -2765.852 & 0.08867 & 186.618 & 29.0069 & 1.3718 & 4.7613 \\
(11) & $-2\nu V_1^{1/3}F_1$ & -2744.064 & 0.03152 & 187.351 & 28.6654 & 1.3650 & 4.9333 \\
(12) & $-2\nu V_1^{1/3}(F_1 - \frac{1}{2}F_{-2})$ & -2742.266 & 0.02609 & 187.744 & 27.3852 & 1.1655 & 4.1455 \\
\hline
\end{tabular}
\end{table}

The expectation value $\langle \frac{1}{r} \rangle$ in Fig. 2 is quite smooth for all models, as mainly strongly bound electrons contribute to the density near the nucleus. The ES model handles them explicitly, while the TFD-$\frac{1}{2}$ model seems to handle them implicitly with the von Weizsäcker term.

In Fig. 3, we start to see the so-called shell oscillation for $\langle r \rangle$. The main comment here is that we should not take this value too seriously when doing comparisons, as long as the semiclassical model has a decent average over the shell effects. As is obvious, both models satisfy this requirement.

Next we look at the general trend of $\tau^2$ in Fig. 4. The shell oscillations already present in the case of $\langle r \rangle$ are magnified, as we are probing even farther reaches of the atoms. We see again that both OFDFT models are reasonable.

Comparing the ES model with the KS-LDA and experimental numbers should only be taken seriously for inert atoms, which have a closed-shell structure. In xenon ($Z = 54$), we see that the experimental value is closer to the semiclassical value than the KS-LDA. The ES model is actually in better agreement with the experimental values as it assigns smaller sizes for almost all atoms (see supplementary material). The effect is mostly due to the exchange term as seen from Table II. The origin of these shell oscillations in the Kohn-Sham model is obvious: some outer orbitals are more delocalized than others.

We can see next in the comparison of different atomic models the absolute value of the numbers presented as general trends in Figs. 1–4. We feel it is informative enough to focus on three representative closed-shell systems: argon

\begin{figure}[h]
\centering
\includegraphics[width=0.5\textwidth]{fig1.png}
\caption{Total energy error compared to the Kohn-Sham energy as a function of $Z$.}
\end{figure}
(low-$Z$, where the strongly bound electron approximation is still valid), krypton (medium-$Z$), and xenon (high-$Z$).

From Figs. 1–4 and Tables III–V, we can draw some conclusions. First is the remarkable and well-known accuracy of the TFD+$\frac{1}{2}$vW model and the fact that the ES model is much better than the TFD+$\frac{1}{2}$vW model, which has a functional that is formally expanded to the same order in $\hbar$ but is missing the corrections for strongly bound electrons. We did not include the TFD+$\frac{1}{2}$vW model in Figs. 1–4 because the accuracy is substantially lower than that for other models. The deviation of the TFD+$\frac{1}{2}$vW model from the KS-LDAs for argon, krypton, and xenon is 7.1%, 5.4%, and 4.6%, which is substantially higher than for the TFD+$\frac{1}{2}$vW or ES model, as seen from Fig. 1. Overall, it seems that after $Z > 40$, the TFD+$\frac{1}{2}$vW and the ES models offer relatively similar accuracy, with the TFD+$\frac{1}{2}$vW model being slightly better.

For all models, the density-dependent quantities $\langle 1 \rangle$, $r^2$, and $\langle \frac{1}{r} \rangle$ are quite reasonable, surprisingly even for the TFD+$\frac{1}{2}$vW model. This reflects the fact that all the models have a reasonable density average over the shell effects in KS-LDA densities, which are shown in Fig. 5.

Near the nucleus, the TFD+$\frac{1}{2}$vW model has the best average description compared to the KS-LDA if we look at quantity $\langle 1 \rangle$. The worst deviation for the TFD+$\frac{1}{2}$vW model is for krypton with 0.6% from the KS-LDA, while for the TFD+$\frac{1}{2}$vW model and for the ES model, the worst case is argon, with deviations of 6.1% and 6.7%, respectively.

The first quantity to contain shell effects is $\langle r \rangle$. For these values, all the models give surprisingly similar results. The worst case for the ES model is argon, where we have a deviation of 3.2%, while the worst case for kinetic energy functionals is krypton, where the deviation is ~5.1%. Again we note that shell oscillation plays a role here and that the argon value for the ES model is probably particularly bad; in Fig. 3, we see a bump in the ES values for small $Z$. This is most likely a slight artifact caused by the averaging procedure.

The quantity $r^2$ is an interesting one. For argon, all the semiclassical models give similar results, which are significantly above the KS-LDA, up to ~10% deviation for the TFD+$\frac{1}{2}$vW model. The case for krypton is similar, except for the ES model, which is closer to the KS-LDA than kinetic energy functionals. Finally, for krypton, the ES model is below all the other models, which is a good thing if we look at the experimental values in Fig. 4.

For inert atoms, there is a trend: all semiclassical values start above KS-LDA values in argon and end up below the KS-LDA in xenon. However, we should not read too much into this as the quantity is quite dependent on the outer orbitals of the particular element, as Fig. 4 shows. The ES model clearly wins in the description of the atom’s outer reaches when considering experimental values.

### Table III. Energies and averages of different models for argon $Z = 18$.

<table>
<thead>
<tr>
<th>Quantity</th>
<th>KS-LDA</th>
<th>TFD+$\frac{1}{2}$vW</th>
<th>TFD+$\frac{1}{2}$vW</th>
<th>ES</th>
</tr>
</thead>
<tbody>
<tr>
<td>$E$</td>
<td>-524.51</td>
<td>-561.80</td>
<td>-524.75</td>
<td>-518.60</td>
</tr>
<tr>
<td>$\langle r \rangle$</td>
<td>16.205</td>
<td>16.202</td>
<td>16.267</td>
<td>16.729</td>
</tr>
<tr>
<td>$r^2$</td>
<td>1.489</td>
<td>1.61</td>
<td>1.630</td>
<td>1.611</td>
</tr>
<tr>
<td>$\langle \frac{1}{r} \rangle$</td>
<td>69.567</td>
<td>73.796</td>
<td>69.566</td>
<td>74.228</td>
</tr>
</tbody>
</table>
TABLE IV. Energies and averages of different models for krypton \(Z = 36\). Results are in atomic units.

<table>
<thead>
<tr>
<th>Quantity</th>
<th>KS-LDA</th>
<th>TFD+(\frac{1}{2})vW</th>
<th>TFD+(\frac{1}{2})vW</th>
<th>ES</th>
</tr>
</thead>
<tbody>
<tr>
<td>(E)</td>
<td>-2746.649</td>
<td>-2895.528</td>
<td>-2744.153</td>
<td>-2742.266</td>
</tr>
<tr>
<td>(\langle r \rangle)</td>
<td>26.37</td>
<td>27.69</td>
<td>27.72</td>
<td>27.81</td>
</tr>
<tr>
<td>(r^2)</td>
<td>1.120</td>
<td>1.274</td>
<td>1.277</td>
<td>1.166</td>
</tr>
<tr>
<td>(\langle 1/r \rangle)</td>
<td>182.63</td>
<td>190.14</td>
<td>181.63</td>
<td>186.34</td>
</tr>
</tbody>
</table>

TABLE V. Energies and averages of different models for xenon \(Z = 54\). Results are in atomic units.

<table>
<thead>
<tr>
<th>Quantity</th>
<th>KS-LDA</th>
<th>TFD+(\frac{1}{2})vW</th>
<th>TFD+(\frac{1}{2})vW</th>
<th>ES</th>
</tr>
</thead>
<tbody>
<tr>
<td>(E)</td>
<td>-7223.567</td>
<td>-7556.458</td>
<td>-7208.302</td>
<td>-7207.0235</td>
</tr>
<tr>
<td>(\langle r \rangle)</td>
<td>39.131</td>
<td>37.610</td>
<td>37.606</td>
<td>38.650</td>
</tr>
<tr>
<td>(r^2)</td>
<td>1.172</td>
<td>1.092</td>
<td>1.092</td>
<td>1.000</td>
</tr>
<tr>
<td>(\langle 1/r \rangle)</td>
<td>317.670</td>
<td>330.690</td>
<td>317.600</td>
<td>323.931</td>
</tr>
</tbody>
</table>

The ionization potential calculated with a difference of total energies \(E(N) - E(N-1)\) and the results are in Table VI. We only calculated it for alkali metals as the ES model will fail to provide reasonable potential for other elements because the shell effects are missing. A deviation in the ionization potential emerges (up to 9% deviation for cesium), but the trend is similar for both methods.

Atomic densities for closed-shell atoms are shown in Fig. 5. We also plot Kohn-Sham densities as a reference. We can see that the ES densities are not completely structureless; they do contain some structure due to averaging over the hydrogenic shells. Yet, obviously, they do not contain the shell structure of Kohn-Sham due to single particle states.

Previous results prove that the ES model does not lose to the density functional models and is even better than Kohn-Sham in some special cases. But what the ES model mostly supplies is theoretical clarity and rigor. One thorny issue with TFD+\(\lambda\)vW models and other GGA-based OFDFT models has been the Pauli potential \(\nu_{\Theta}\)’s negativity, which is defined as

\[
\nu_{\Theta} = \delta T_{\Theta}/\delta n - \delta T_{vW}/\delta n = V - \mu - \delta T_{vW}/\delta n,
\]

where \(T_\Theta\) is the non-interacting kinetic energy and \(T_{vW}[n]\) is the von Weizsäcker kinetic energy functional. It has been shown that \(\nu_{\Theta}\) should always be positive,\(^4\) but for TFD+\(\lambda\)vW and many other GGA kinetic functionals, it has been found to be negative near a nucleus.\(^5\) This emerges because the semiclassical evaluation is not valid near a nucleus; see Ref. 16 for an excellent discussion. The ES model does not have this problem because the problematic region is removed from the semiclassical evaluation. Figure 6 shows the self-consistent Pauli potentials. Clearly in the self-consistent calculations, the Pauli non-positivity problem is contained in a small region, but the Pauli potential is still qualitatively wrong. In comparison, the ES model’s Pauli potential is qualitatively better, although naturally missing the shell effects. The ES model has oscillations due to the averaging over shells, but this is more of an artifact than a result of the oscillations being in the wrong region.

This raises the question of how the TFD+\(\frac{1}{2}\)vW’s energetics can be so good (while having reasonable geometric properties, too) if it completely ignores the correction for strongly bound electrons. The answer must be related to the dual nature of the von Weizsäcker term because it has two roles as a density functional: it is a gradient correction to Thomas-Fermi, but it also is exact for up to two non-interacting particles. These two roles naturally differ by a constant factor, but the form is the same. Kinetic energy functionals of GGA type can be built around the dual nature by interpolating between these two extremes.\(^5,25,26\) Here we are inclined to support the view of Englert and Schwinger: it is more straightforward to exclude the strongly bound electrons from the semiclassical evaluation than to modify the functional to include them via the von Weizsäcker term. The near-nucleus Pauli potential singularity is also avoided by the use of pseudopotentials as the strongly bound electrons are excluded from the OFDFT calculation. The non-negativity constraint might be a non-issue for valence density.\(^27\) We also must remember that the \(\lambda\) value of \(\frac{1}{2}\) is obtained by fitting, while the strongly bound electron correction is better motivated. If the ES model is compared to the gradient expansion in atoms, then it is clearly preferable.
TABLE VI. Ionization potential in Hartrees calculated by the energy difference \( E(Z) - E(Z - 1) \) for ES and KS models. KS results are calculated with the GPAW DFT code,\(^{22}\) where we used Dirac exchange functional.

<table>
<thead>
<tr>
<th>Atom</th>
<th>Density</th>
<th>Energy</th>
<th>( \bar{r} )</th>
<th>( \langle \frac{1}{r} \rangle )</th>
</tr>
</thead>
<tbody>
<tr>
<td>K</td>
<td>0.1310</td>
<td>0.1364</td>
<td>218.591</td>
<td>74.228</td>
</tr>
<tr>
<td>Rb</td>
<td>0.1202</td>
<td>0.1319</td>
<td>274.230</td>
<td>187.744</td>
</tr>
<tr>
<td>Cs</td>
<td>0.1133</td>
<td>0.1223</td>
<td>7207.024</td>
<td>323.598</td>
</tr>
</tbody>
</table>

D. Accuracy of self-consistent density approximations

As mentioned earlier, it is not necessary to calculate the density with the full expression (8) to get the same semiclassical accuracy. Only the first line of (8) is necessary, as Englert argued.\(^{16}\) The rest of the expression contains full divergence—i.e., it integrates to zero, so it does not contribute to the number of electrons, only to the distribution. We are interested in the approximation for future use, if the method is extended to larger systems. We also want to see the effect on \( \bar{r}^2 \) to see how sensitive it is to density approximations because we already established the effect of exchange.

The simple density approximation is obtained simply by discarding the higher-order variations. The resulting expression is

\[
\hat{n}_{\text{approx}} \approx \frac{\partial e_1}{\partial V} = \frac{1}{2\pi} [2\nabla V F_2 - 1/6 \pi |2\nabla V|^{-1/3} \nabla^2 V F_0]. \tag{14}
\]

We compare the difference of two density expressions (14) and (8) in self-consistent calculations. We focus on krypton for simplicity, but the trends are reproduced over a range of Z. As both integrate to the electron number, the difference is purely just a redistribution of the density.

From Table VII, we can see that the simple density assigns more density near the nucleus while the full expression localizes the density more, which is seen in \( \bar{r}^2 \). Full density has a bit better energy when compared to the KS-LDA. For energy, the maximal deviation between density expressions is 0.4% percent in xenon, and for \( \bar{r}^2 \), the maximal deviation is 3% percent in argon. On a semiclassical scale, these differences are quite small. Overall the differences show that the full density (8) is better, as expected, but the simpler expression (14) is still valid.

V. CONCLUSION

Having benchmarked the self-consistent ES model, we found that it compares favorably against the simple TFD-\( \lambda vW \) model. The self-consistent model does not achieve the accuracy of Englert and Schwinger’s perturbative work, yet the performance is good and could lead to a viable means of obtaining physically plausible full solutions to OFDFT equations. An important point to remember here is that self-consistent atoms are not valuable themselves, as they are just a stepping stone toward larger systems. The problematic points of the self-consistent ES model are that it cannot be applied to light elements (according to our tests, the strongly bound electron approximation breaks down numerically at \( Z \sim 12 \)) and its non-ambiguous inclusion of exchange effects. Given this model’s potential, however, such issues are worth considering and investigating with rigor; these issues can and will be addressed in future work.

SUPPLEMENTARY MATERIAL

See supplementary material for more experimental data points for the quantity \( \bar{r}^2 \).

ACKNOWLEDGMENTS

We would like to thank Tom Gustafsson for help with the Finite Element Method solver. This work was supported by the Academy of Finland Projects 279240, 284621, and 312556. The authors are also grateful to CSC, the Finnish IT Center for Science Espoo, and the Applied Physics Department of Aalto for computations.

24 W. Haynes, CRC Handbook of Chemistry and Physics, 93rd ed. (Taylor & Francis, 2012).
27 Authors are not aware of any kinetic energy functional that breaks this condition for valence densities.