A Simple Generalized Gradient Approximation for the Non-interacting Kinetic Energy Density Functional

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A simple, novel, non-empirical, constraint-based orbital-free generalized gradient approximation (GGA) non-interacting kinetic energy density functional is presented along with illustrative applications. The innovation is adaptation of constraint-based construction to the essential properties of pseudo-densities from the pseudo-potentials that are essential in plane-wave-basis ab initio molecular dynamics. This contrasts with constraining to the qualitatively different Kato-cusp-condition densities.

Introduction. Hohenberg-Kohn density functional theory (DFT) [1, 2] has come to prominence mainly in Kohn-Sham (KS) orbital form [3]. However, driving ab initio molecular dynamics (AIMD) [4–7] with KS DFT exposes a computational cost-scaling burden. The KS computational cost scales no better than \(N^3\) with \(N\), the number of electrons or number of thermally occupied bands. Additionally there is reciprocal space sampling cost or equivalent costs from large real-space unit cells used with Γ-point sampling. In contrast, orbital-free DFT (OF-DFT) offers linear scaling with system size [8, 9] for use of AIMD on arbitrarily large systems.

The long-standing barrier to widespread use of OF-DFT has been the lack of reliable non-empirical approximate kinetic energy density functionals (KEDFs). In terms of the KS orbitals \(\varphi_j\), the reference, positive definite KS kinetic energy (KE) density is

\[
t_s = t_s^{orb} = \sum_{j=1}^{N_e} |\nabla \varphi_j|^2 ,
\]

in Hartree atomic units with \(n(r)\) the electron number density (and unit occupation for simplicity). Two types of approximate KEDFs have been explored, semi-local (one-point)

\[
T_s[n] = \int d^3r t_s[n(r), \nabla n(r), \ldots]
\]

and two-point with a non-local term

\[
T_{NL}[n] = -c_{TF} \int d^3rd^3r' n^\alpha(r) K[n(r), n(r'), r, r'] n^\beta(r')
\]

with \(c_{TF} = c_{TF} = \frac{3}{16} (3\pi^2)\frac{3}{2}\). For a dimensionless \(K\), \(\alpha + \beta = 8/3\). Most approximate Ks are parametrized; see Refs. [9–16] for details as well as brief discussion below. In this communication, we propose a novel non-empirical one-point KEDF and show that it is competitive with current two-point KEDFs, generally better than other one-point functionals, more transferable, and notably faster.

Generalized Gradient Approximations. The simplest one-point functionals are Thomas-Fermi [17–19]

\[
T_{TF}[n] := \int d^3r t_{TF}(r), \quad t_{TF}(r) := c_{TF} n^{\frac{5}{2}}(r),
\]

and von-Weizsäcker [20]

\[
T_W[n] := \frac{1}{8} \int d^3r \frac{|\nabla n(r)|^2}{n(r)} \equiv \int d^3r t_W(r).
\]

Neither is satisfactory as a general KEDF. As with approximate exchange-correlation (XC) functionals [21], the gradient expansion of the weakly inhomogeneous electron gas KE leads to consideration of generalized gradient approximations (GGA) for \(T_s\),

\[
T_s^{GGA}[n] = \int d^3r t_{TF}(r) F_1(s(r)).
\]

Here \(F_1(s)\) is the GGA KE enhancement factor, a function of the dimensionless reduced density gradient \(s := |\nabla n| = \frac{1}{2(4\pi)^{3/2}} \frac{\int |\nabla n|}{\int n^{1/3}}\) familiar from GGA X functionals. GGA KEDFs so constructed automatically satisfy \(T_s\) uniform scaling requirements [22]. In GGA form the von Weizsäcker KE becomes \(F_W(s) = \frac{5}{3}s^2\).

From the Pauli term decomposition [8, 23, 24],

\[
T_s[n] = T_W[n] + T_P[n],
\]

three constraints follow[25],

\[
\begin{align*}
T_P[n] & \geq 0, \\
\nu_\theta(r) & \geq 0 \forall r, \\
\nu_\theta(r) & \geq \frac{\theta_0(r)}{\nu_\theta(r)} \forall r, \quad \theta_0 := t_s^{orb} - t_W,
\end{align*}
\]

with the Pauli potential defined as \(\nu_\theta(r) := \delta T_\theta[n]/\delta n(r)\) and the Pauli enhancement factor is \(F_\theta(s) = F_1(s) - F_W(s)\).

To date, perhaps the best constraint-based GGA KEDF is VT84F (evaluated at T=0 K of course)[26].
It is successful in finite-T AIMD simulations [27] and is the only non-empirical GGA KEDF that yields reasonable binding in simple solids. It was constrained to satisfy Eqs. (8) and (9) for physical atom densities, i.e., those that obey the Kato cusp condition [28]. VT84F also was constrained to respect \( \lim_{s \to \infty} F_\theta(s)/F_W(s) = 0 \). This comes from the one-electron tail region of a many-electron atom[29] where \( t_\theta/t_W \) must vanish, hence \( t_s \to t_W \) [30].

In terms of the universal Hohenberg-Kohn-Levy density functional, such a physically motivated constraint is non-universal: the Kato cusp condition is specific to an external Coulomb potential. Such non-universality is rational for material and molecular property calculations. But the ubiquitous use of pseudo-potential plane-wave basis methods in AIMD simulations means that it is not the optimal non-universality for them. OF-DFT calculations in fact require a local pseudo-potential (LPP). The OF-DFT Euler equation then implies that \( \nu_\theta \) is closely related to the LPP \( \delta_{\text{pseudo}} \) and that \( \nu_\theta \) is evaluated with the corresponding pseudo-density. Thus any constraint based on density characteristics should be specific to a particular type or class of pseudo-potential.

Ref. [31] explored some elementary consequences for constraint satisfaction (or violation) with non-Kato densities. Difficulties with simpler one-point KEDFs (linear combinations of \( T_{TF} \) and \( T_W \) ) used with orbital-free projector augmented-wave pseudo-densities also have been reported [32]. So far as we know, no approximate KEDF has been constructed by explicit satisfaction of the foregoing constraints, Eqs. (8)-(9), for a specified type of pseudo-densities. Nor has Eq. (10) been used.

**New GGA KEDF** We resolve this pseudopotential AIMD deficiency by devising a GGA KEDF constrained to satisfy Eqs. (8) and (9) for pseudo-densities of a particular kind and show that in most spatial regions its \( \nu_\theta \) satisfies Eq. (10) as well. The new GGA KEDF enhancement factor is

\[
F_{t}^{LKT}(s) = \frac{1}{\cosh(\alpha s)} + \frac{5}{3} s^2
\]

with parameter \( \alpha > 0 \). Fig. 1 compares \( F_{t}^{LKT} \) with the VT84F and APBEK [33] enhancement factors. It satisfies the obvious homogeneous electron gas constraint \( \lim_{s \to 0} F_{t}/F(s) = 1 \) and obeys \( 0 \leq F_{t}^{LKT} \leq 1 \) so as to satisfy the bound conjectured by Lieb [34, 35]

\[
T_s \leq T_{TF} + T_W
\]

(12)

\( F_{t}^{LKT} \) also satisfies [25, 29, 36] \( \nu_\theta([n]; r) \geq 0 \ \forall r \), thus \( T_{\theta}^{LKT} \geq 0 \).

The sole parameter \( \alpha = 1.3 \) (which was used throughout all the subsequent calculation was determined as follows. A set of pseudo-densities was generated for the atoms H through Ne with a typical Hamann norm-conserving non-local pseudo-potential (NLPP) scheme [37] using default radii in the APE code [38] and the Perdew-Zunger (PZ) XC local density approximation (LDA) [39]. Then \( \alpha \) was found such that all the post-scf Pauli potentials from those pseudo-densities satisfied \( \nu_\theta \geq 0 \ \forall r \). Importantly, as long as an \( \alpha \) value gave \( \nu_\theta \geq 0 \) for the H atom, positivity also was met for all the heavier atoms. For Li \( \alpha < 1.4 \) is required, while for H, \( \alpha \leq 1.3 \) is needed to get a post-scf \( \nu_\theta \geq 0 \). For He, the \( \alpha \) value does not seem to matter within the range tested. While the \( \alpha \) value is non-universal, we expect reasonable transferability to those other pseudo-potential types for which the pseudo-densities are similar, specifically those with nearly flat pseudo-densities near the nucleus. The expectation is confirmed by post-scf and scf calculations for atoms.

Though the reference atom set, H–Ne, encompasses 1–8 pseudo-electrons, equally good performance for other elements is not assured. Post-scf determination of \( \alpha \) also is distinct from self-consistent calculation, which might vitiate the supposedly constrained behavior. Atomic tests are the first line of investigating these issues. For a given pseudo-potential and XC approximation, self-consistent solution of the KS equation provides the reference KS \( t_\theta \) and the ingredients to construct the reference KS Pauli \( \nu_\theta \) (see Eq. (35) in Ref. [23]). Those are the standards against which to judge \( t_\theta \) and \( \nu_\theta \) from an approximate KEDF. In anticipation of the OF-DFT calculations on periodically bounded systems reported below, we focused upon the bulk-derived LPP (BLPS) [40, 41] for two atoms, Al and Li. Here we discuss Al because it was not in the \( \alpha \) calibration. Li discussion is in the Supplemental Material [42]. (The Li pseudo-atom is challenging because it is a one-orbital system (2s\(^1\)) for which \( T_\theta \) should vanish.) Again the XC functional is PZ.

Fig. 2 displays the reference \( t_\theta/n \) and \( \nu_\theta \) for the BLPS Al pseudo-atom in the 3s\(^2\) 3p\(^1\) configuration and the post-scf results with that pseudo-density for both VT84F and LKT. Note several features. Though VT84F was constructed to satisfy \( \nu_\theta^{VT84F} \geq 0 \) near a nucleus for Kato-
and the OF-DFT calculations used Profess LDA XC functional and BLPS were used. For comparison for AIMD requires accuracy tests on extended this constraint because doing so would require post-scf case. We did not insist on strict imposition of value that materially worsens results for solids.

convergences as well as cause other difficulties. Though \( v_\theta^{VT84F} \) decays similarly, its rapid oscillations in the dominant density region might slow scf convergence rates as well as cause other difficulties.

Self-consistent OF calculations for the BLPS Al pseudo-atom show that \( v_\theta^{LKT} \) stays positive, though it exhibits oscillations quite similar to those seen in the post-scf case; see Fig. 3. The LKT Pauli energy per particle is far from the KS value. However, the inequality Eq. (10) is violated only around \( r = 1.8 \) bohr as in the post-scf case. We did not insist on strict imposition of this constraint because doing so would require \( a \lesssim 0.8 \), a value that materially worsens results for solids.

**Performance on Solids.** Validation of the new functional for AIMD requires accuracy tests on extended systems. We therefore did KS-DFT and OF-DFT calculations on simple metals and semiconductors. Conventional KS calculations were done with ABINIT [44] and the OF-DFT calculations used PROFESS[45] and/or PROFESS@QUANTUM-Espresso [46]. Again the PZ LDA XC functional and BLPS were used. For comparison we included the Wang-Govind-Carter (WGC) [13], Huang-Carter (HC) [14], and Constantin et al. KGAP [16] two-point KEDFs and the one-point Constantin et al. SOF-CFD [47] meta-GGA (Laplacian-dependent) KEDF. Technical details and parameter values are in the Supplemental Material [42].

Note that WGC was parametrized for main-group metals and yields poor binding curves for semiconductors, while HC was parametrized for semiconductors. KGAP is parametrized to experimental direct band gaps. Results from the one-point functionals E00, APBEK, and PBE2 are omitted because of unrealistic binding curves for the former two and instability problems for the latter one. KGAP comparisons are from Tables I and II of Ref. [16]. SOF-CFD values are from Table I of Ref. [47]. Equilibrium volumes, energies, and bulk moduli for other functionals were generated by varying \( \pm 5\% \) around the equilibrium volume to obtain 11 energy-volume points, which then were fitted to the Birch-Murnaghan equation of state [48].

The metals were Li, Mg, and Al in the simple cubic, body-centered cubic, face-centered cubic, and hexagonal close-packed structures. Nine III-V semiconductors in zinc-blende structures were treated: AlP, AlAs, AlSb, GaP, GaAs, GaSb, InP, InAs, and InSb.

With KS quantities as references, Table I shows the mean absolute relative error (MARE) percentages for equilibrium volume \( V_0 \), energy \( E_0 \) per atom (for metals) or per cell (for semiconductors), and bulk moduli \( B_0 \) from WGC, HC, KGAP, VT84F, SOF-CFD, and LKT. These are calculated as \(|Q_{KS} - Q_{KS}|/Q_{KS} \times 100/N_{systems} \), where \( Q \) is \( V_0 \), \( E_0 \), or \( B_0 \). (More detailed tabulations are in the Supplemental Material [42].) For \( V_0 \) and \( B_0 \), LKT is a significant improvement over VT84F. The \( V_0 \) and \( B_0 \) MAREs are reduced by 33% in metals. The reduction is more dramatic in the semi-conductors, a factor of 5 for \( V_0 \) and 13 for \( B_0 \). The semiconductor \( E_0 \) MARE is reduced by 22% but worsened slightly from 0.1% to 0.2% for the

![FIG. 3. Top: Al KS (solid, red) and LKT(dashed, blue) pseudo-densities as function of radial position. Bottom: KS vs. LKT \( \rho \) (solid red vs. dash-dotted blue, upper pair) and similarly \( t_s/n \) (dashed red vs. dotted blue, lower pair).](image-url)
TABLE I. KEDF performance on solid metals and semiconductors: MARE of equilibrium volumes $V_0$, energies $E_0$, and bulk moduli $B_0$, as percentages. See text for notation.

<table>
<thead>
<tr>
<th>KEDF</th>
<th>Metals</th>
<th>Semiconductors</th>
</tr>
</thead>
<tbody>
<tr>
<td>WGC</td>
<td>$V_0$</td>
<td>$E_0$</td>
</tr>
<tr>
<td>HC</td>
<td>0.7</td>
<td>0.0</td>
</tr>
<tr>
<td>KGAPa</td>
<td>5.5</td>
<td>0.6</td>
</tr>
<tr>
<td>VT84F</td>
<td>4.0</td>
<td>-</td>
</tr>
<tr>
<td>SOF-CFDa</td>
<td>5.2</td>
<td>0.6</td>
</tr>
<tr>
<td>LKT</td>
<td>4.0</td>
<td>0.2</td>
</tr>
</tbody>
</table>

*a Note: only metals with cubic symmetry were included and PBE XC was used.

metals. Except for performance on semi-conductor $E_0$, it also is clear that the LKT GGA is superior to the more-complicated non-empirical SOF-CFD meta-GGA KEDF. Despite noticeable discrepancies in absolute energies for semiconductors, it is important to note LKT OFDFT gives the same phase ordering as does conventional KS; see Table III in the Supplemental Material.

Regarding the two-point functionals, WGC outperforms all the other functionals on the metals but is inapplicable on semiconductors, recall above [13]. Conversely, HC with averaged parameters exhibits balanced error, with all three MAREs within 5% (except $B_0$ for metals). KGAP does well on volumes in both classes but not $B_0$. Remarkably LKT exhibits performance competitive with both HC and KGAP in prediction of equilibrium volumes for both material classes. Moreover, LKT outperforms HC for $B_0$ and is much more balanced than KGAP for $B_0$. (Comparison with the recent MGP two-point functional is of no avail, since its parametrization is tuned to match KS results for each system [15].)

For the case of AlP, we found that LKT converges for relatively smaller energy cutoff than needed with VT84F and HC. Typically LKT also requires fewer self-consistent iterations for solution to a given tolerance than needed by either HC or VT84F and each LKT iteration is typically about one-fifth the time of an HC iteration. Thus the one-point LKT is more useful as a broadly applicable functional than the highly parametrized two-point HC KEDF or the experimentally parametrized two-point KGAP KEDF yet is simpler, faster, and mostly better than the SOF-CFD one-point KEDF. LKT seems therefore to be currently the most promising candidate for general AIMD OF-DFT use or with small-box algorithms [49]. Though it remains to be tested, we anticipate the finite-T generalization [50] of LKT will be of value for warm dense matter simulations.

As to limitations, LKT does not yield a good value of $V_0$ for bcc Li with a 3-electron LPP. So far as we know, all GGA KEDFs developed so far share this limitation. The extent of transferability to another distinct class of pseudo-potential, along with the post-scf determination of $a$, remains to be examined.

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Kerala, India, 2009) p. 25.


[47] L.A. Constantin, E. Fabiano, and F. Della Sala, arXiv:1802.02889 [cond-mat.mtrl-sci]. This functional was denoted “SOF” (semi-local orbital-free) but that properly is the name of a class of functionals. For specificity we append the authors’ initials, “SOF-CFD”.

