

Orbital-Free Density Functional Theory: An Attractive Electronic Structure Method for Large-Scale First-Principles Simulations

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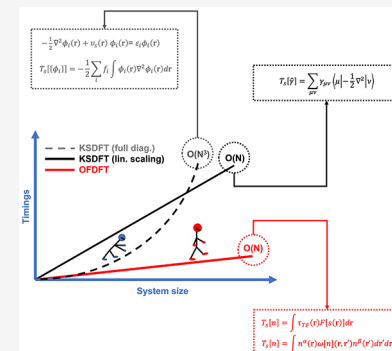

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ABSTRACT: Kohn–Sham Density Functional Theory (KSDFT) is the most widely used electronic structure method in chemistry, physics, and materials science, with thousands of calculations cited annually. This ubiquity is rooted in the favorable accuracy vs cost balance of KSDFT. Nonetheless, the ambitions and expectations of researchers for use of KSDFT in predictive simulations of large, complicated molecular systems are confronted with an intrinsic computational cost-scaling challenge. Particularly evident in the context of first-principles molecular dynamics, the challenge is the high cost-scaling associated with the computation of the Kohn–Sham orbitals. Orbital-free DFT (OFDFT), as the name suggests, circumvents entirely the explicit use of those orbitals. Without them, the structural and algorithmic complexity of KSDFT simplifies dramatically and near-linear scaling with system size irrespective of system state is achievable. Thus, much larger system sizes and longer simulation time scales (compared to conventional KSDFT) become accessible; hence, new chemical phenomena and new materials can be explored. In this review, we introduce the historical contexts of OFDFT, its theoretical basis, and the challenge of realizing its promise via approximate kinetic energy density functionals (KEDFs). We review recent progress on that challenge for an array of KEDFs, such as one-point, two-point, and machine-learnt, as well as some less explored forms. We emphasize use of exact constraints and the inevitability of design choices. Then, we survey the associated numerical techniques and implemented algorithms specific to OFDFT. We conclude with an illustrative sample of applications to showcase the power of OFDFT in materials science, chemistry, and physics.



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1. INTRODUCTION

1.1. Motivation

The extraordinarily widespread use of density functional theory (DFT) in its conventional computational form via Kohn–Sham (KS) orbitals and their eigenvalues¹ masks a remarkable fact: those orbitals are not essential to Kohn–Sham DFT (KSDFT). This review focuses on exploitation of that fact in the electronic structure method known as orbital-free density functional theory (OFDFT). In a very real sense it is the more fundamental form of KSDFT. OFDFT literally puts the density back into centrality in DFT, both formally and in practice.

The immediate motivation is plain. Given the need to consider only the electron density as the main ingredient of the method, OFDFT can have a computational cost scaling linearly with system size *irrespective of system state*. Therefore, OFDFT is on the verge of becoming the alternative to conventional KSDFT in at least two limiting categories: (1) systems for which the size is too large or the number of simulations is too large to be tackled by conventional KSDFT and (2) systems with temperature and/or density too large to be tractable with conventional KSDFT. These two categories, especially the first, are becoming increasingly relevant as chemistry and materials physics push into systems of high molecular and structural complexity.

We say OFDFT “is on the verge” because it involves a problem analogous with one familiar in conventional KSDFT, namely construction of accurate approximate density functionals. The difference is that OFDFT needs kinetic energy density functionals (KEDFs) as well as the exchange–correlation (XC) functionals familiar in conventional KSDFT.²

The reader may have noticed the terminology “conventional KSDFT”. Perhaps this seems an unmotivated complication. But the reasons for making the distinction are deep and will become clear soon. The essential point is simple. OFDFT is a way of doing KSDFT. But it does not use the KS orbitals explicitly. Instead it uses KEDFs. KEDFs connect the theory to the earliest days of quantum mechanics. It is perhaps underappreciated that the earliest forms of OFDFT preceded

the Hohenberg–Kohn theorems³ of modern DFT. That earlier work provided the intellectual foundations and ample precedent for treating many-electron problems via a suitably constructed effective non-interacting system. We do not pretend to be historians of science. Nonetheless, the centrality of that non-interacting system concept motivates our historically oriented account of those pre-Hohenberg–Kohn roots. In it we try to do justice to the great effort of many researchers. That effort built the context that distinguishes conventional KSDFT in its ubiquitous practical form and KSDFT in its most basic form, namely OFDFT. We then lay out the relevant parts of formal DFT and use that to clarify the distinctive differences between conventional KSDFT and OFDFT. There follows a survey of modern KEDFs, with pointers to earlier ones via prior reviews. We devote a section to the software implementations of OFDFT in its various flavors and another to some of the most notable contemporary applications.

In all, we attempt to give a clear, candid account of the state of the art in OFDFT functional development and its algorithmic realizations in advanced electronic structure software, describing both the limitations and the opportunities for applications in chemistry and materials science.

1.2. Curse of Computational Scaling

In recent years, some chemists, materials scientists, and physicists have speculated upon “lab 2.0”.⁴ It is conceived as a scientific inquiry mode in which computer simulations replace experiments for tasks such as materials design and combinatorial search of chemical space. Even though computationally aided materials design and chemical discovery via tools such as high-throughput workflows now are daily occurrences,^{5,6} the computational cost scaling of adequately accurate quantum-mechanical methods hinders progress severely. “lab 2.0” thus still largely belongs to the future.

Disparity between desired computational approaches and available computational resources is not a new problem. At each stage of computational resource development, from pen and paper to abacus, slide rule, rotary calculator,^{7,8} desk calculator, analog machine, ENIAC,⁹ ..., it has been easy to define calculations intractable on currently available resources.

More profoundly, nature imposes some computational scaling lower bounds.^{10,11} The focus issue is electronic structure. It underlies the periodic table, which is normative for the chemistry of materials and their molecular constituents. In the simplest case, the nuclei are fixed and the electrons are treated by some model that makes them independent in a mean-field of the others, e.g. Hartree–Fock (HF)¹² or KSDFT.¹ In both, the straightforward implementation, in terms of a determinant (HF or conventional KSDFT) and standard XC approximants, causes the number of operations to grow as the cube of the number of electrons (i.e., $O(N_e^3)$ or simply $O(N^3)$ hereafter) because of the orthogonalization of the one-electron orbitals. In addition, the straightforward approach of expanding in a basis of local orbitals can lead to large additional costs caused by quadratic scaling for building the needed matrices. Scaling of this sort can be mitigated by employing techniques such as distance-based screening coupled with sparse linear algebra^{13–15} and/or fast Fourier transforms¹⁶ to recover almost linear scaling algorithms for matrix build and Hamiltonian-wave function operations, albeit with a large prefactor^{11,17} and restriction to systems wherein

Table 1. Chronology of Orbital-Free Density Functional Theory^a

Year	Milestone	Method
1927	Thomas and Fermi present their approximate method that satisfies the Pauli principle while avoiding the Schrödinger equation	TF ^{22,23}
1930	TF is applied to atoms such as Fe yielding qualitative agreement with experimental sequential ionization energies ²⁴	
1930	Dirac extends TF to include exchange effects	TFD ²⁵
1935	Slater extends TF to the condensed phase and applies it to metals, but no binding is found ²⁶	
1935	von Weizsäcker proposes the first gradient correction	TFvW, TFvWD ²⁷
1945	The ENIAC computer is built; patent filed in 1948 ⁹	
1949	Reitz implements TFD on ENIAC for computation of electronic properties of atoms ²⁸	
1949	Feynman, Metropolis, and Teller extend the finite-temperature Thomas–Fermi theory ²⁹	
1957–1982	Extensions of TFvWD and the gradient expansion approximation	GEA ^{30–34} and TF λ vW ^{35–38}
1962	Teller proves that TF cannot lead to molecular bonding ³⁹	
1964	Formal footing of DFT	Hohenberg–Kohn theorems ³
1965	Mapping to a non-interacting electron system and employment of the exact non-interacting KEDF	KSDFT ¹
1964–1965	Introduction of polarizability in the construction of the KEDF	Nonlocal KEDF ^{3,40}
1981–1983	Computational proof that a gradient correction to TF (e.g., TFvWD) leads to molecular binding ^{41,42}	
1985–ongoing	Nonlocal (two-point) KEDFs for condensed phases	CAT, ⁴³ WT, ⁴⁴ WGC, ⁴⁵ SM, ⁴⁶ Perrot, ⁴⁷ Chai–Weeks, ⁴⁸ XWM ⁴⁹ KEDFs, real space, ^{50–53} and reciprocal space ^{54,55} GGA XC ^{56,57}
1986	Generalized gradient approximation for the exchange and correlation energy functional	GGA KEDFs ^{58–63,63–76}
1991	Conjointness conjecture ⁵⁸	HC, ⁷⁷ WGCD, ⁷⁸ EvW-WGC, ⁷⁹ KGAP, ⁸⁰ MGP ⁸¹
2010–ongoing	Nonlocal (two-point) KEDF for semiconductors	
2012–ongoing	Machine-learnt KEDFs ^{82–84}	
2013–ongoing	General-purpose semilocal (one-point) KEDFs	VT84F, ⁸⁵ LKT, ⁸⁶ PGSL ⁸⁷
2019–ongoing	Nonlocal (two-point) KEDF for finite systems and subsystem DFT	LMGP, ^{88,89} LDAK ⁹⁰ and revHC ⁹¹

^aKEDF stands for kinetic energy density functional.

sparsity is physically realistic. More complicated explicit wave function methods simply worsen the scaling.^{18–21}

The curse of computational scaling in quantum mechanics has been known since its inception. Dirac's oft-quoted 1929 remark raises the problem at the end:

*"The underlying physical laws necessary for the mathematical theory of a large part of physics and the whole of chemistry are thus completely known, and the difficulty is only that the exact application of these laws leads to equations much too complicated to be soluble. It therefore becomes desirable that approximate practical methods of applying quantum mechanics should be developed, which can lead to an explanation of the main features of complex atomic systems without too much computation."*⁹²

Back then researchers addressed computational scaling barriers by making clever but drastic simplifications to the quantum mechanical approach. Only much later, in the mid-1940s, did the first Turing complete computer (i.e., a digital computer in the modern sense) surface, the ENIAC project.²⁸ In anticipation of the main focus of this review, we note that Dirac's ambition, to give "...an explanation of the main features...", was modest compared to today's ambition to achieve predictive calculations.

In any event, the late 1920s and the 1930s were replete with pen-and-paper calculations of atoms⁹³ and materials²⁶ which avoided the use of any computationally challenging manipulations such as diagonalization of large matrices. Heroic numerical solutions on manual calculators^{7,94–96} also were achieved.

It was in this environment that Thomas²³ and Fermi²² independently proposed a method that required no diagonalizations. Instead, their approach focuses on a single, central quantum-mechanical quantity, the electron number density, and results in a single differential equation. This is the justly famous Thomas–Fermi theory (TF hereafter). It is the primordial version of OFDFT. Table 1 provides an overview of OFDFT development milestones from then until today.

Because its clear analytical structure incorporates the basic features of many-Fermion systems generally, TF theory has been the subject of intense mathematical scrutiny; hence, much is known about its content and properties. See reviews by March,³⁰ Lieb and Simon,⁹⁷ Spruch,⁹⁸ March,⁹⁹ and Solovej¹⁰⁰ as well as the book by Englert,¹⁰¹ among many resources. Basic development of TF theory is treated in many books, e.g. by Parr and Yang,¹⁰² Slater,^{103,104} Giuliani and Vignale,¹⁰⁵ and Engel and Dreizler,¹⁰⁶ so we sketch only the idea and display relevant bits here.

The foundational insight in TF theory is that a set of electrons in a neutral system can be idealized as behaving much like a non-interacting, uniform gas. That is, one assumes that the real system of interest is nearly indistinguishable from a homogeneous electron gas (HEG). The HEG in turn is assumed to occupy a volume V uniformly with number density $n = \frac{N_e}{V}$. (Remark: We adopt the notation prevalent in the physics literature, n . In the chemistry literature, often ρ is used.) The hard-walled volume forces quantization into discrete energy levels. The electrons must satisfy the Pauli exclusion principle; thus, at most two of them can occupy a given level. This leads to a maximum occupied energy level

related to the charge density. In turn, there is a total kinetic energy (there is no interaction energy) given by

$$T = \frac{3}{10} (3\pi^2)^{2/3} n^{5/3} \equiv c_{TF} n^{5/3} \quad (1)$$

Note that here and henceforth, unless stated otherwise, we are using Hartree atomic units, $\hbar = m_e = q_e = 1$ with m_e and q_e the electron mass and charge magnitude, respectively.

The key conceptual leap then is to make the local density approximation, namely that locally a physical system behaves like an HEG of the density locally. The resulting generalization to inhomogeneous systems is

$$T_{TF}[n] = c_{TF} \int_{\mathbb{R}^3} d\mathbf{r} n^{5/3}(\mathbf{r}) \equiv \int_{\mathbb{R}^3} d\mathbf{r} \tau_{TF}(\mathbf{r}) \quad (2)$$

Along with this, one can associate an electrostatic potential that is as deep as the maximum energy (a simple shift of energy zero) and gives, via Poisson's equation, the proper charge density. This is the prescription for the TF atom, for example. For details not needed here, we refer the reader to Appendix 17 of ref 103.

It often has been remarked that TF theory is the ancestor of DFT (see Table 1). Early interest in TF theory arose, in fact, from its computational feasibility for treating many-electron atoms. When spherical symmetry is present, the TF equations are pseudo-one-dimensional and their solution can be approached with pen and paper. Among the early applications of TF theory is the work of Baker²⁴ aimed at predicting multiple atomic ionizations. This was an important topic that also was pursued by Hartree^{12,107} as an application of the HF method. It, unfortunately, is intractable for pen-and-paper calculation. Using TF theory, Baker²⁴ predicted semiquantitative agreement with experimental ionization potentials (IPs) for atoms such as O and Fe. See Figure 1.

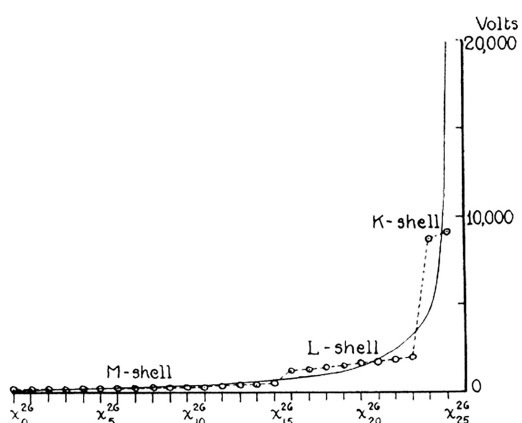


Figure 1. First computation of ionization potentials using Thomas–Fermi theory (full curve). Calculated values for Fe compared to experiment (dashed curve and data points). Reprinted with permission from ref 24. Copyright (1930) American Physical Society.

Dirac became interested in finding commonality between the HF ansatz and TF theory. In 1930 he showed²⁵ that application of the HF approach to an HEG produces a stabilizing exchange term proportional to the maximum momentum attainable by the HEG. It follows that the exchange energy density is proportional to $n^{4/3}$ and thus the associated exchange potential is proportional to $n^{1/3}$.

Dirac's analysis yielded two achievements. First, it justified TF theory (for which there was only a heuristic grounding until then) by constructing the quasi-classical counterpart of the HF problem for a nearly uniform density system. [Remark: Dirac²⁵ achieved the quasi-classical counterpart of the HF approach in a way that nowadays is standard practice.^{108,109} He took what we now call the Wigner transform (introduced in 1927 by Weyl¹¹⁰ and formally only two years later by Wigner¹¹¹) of a dynamical variable to transform the exchange and Coulomb operators into classical phase space functions (i.e., p and q). That led to an expression for the Hamiltonian also as a function of only p and q . Then he evaluated the Poisson bracket with the electron density matrix, which was also assumed to be Wigner-transformed to a function of p and q , and approximated by $\rho(p, q) = 2$ if $p < P$ and zero otherwise (P here is a maximum momentum attainable in the infinitesimal volume of phase space, $(2\pi\hbar)^3$, at point p, q). As the energy is conserved, the Poisson brackets need to be zero, $\{H, \rho\} = 0$. This procedure led him to find the Thomas–Fermi equation (i.e., the functional derivative of eq 2 with respect to the electron density) for atoms, augmented by an additional term stemming from the exchange operator and proportional to the maximum momentum P .] Second, Dirac showed that an exchange term should be added to the TF energy and potential.

However, already in the 1930s there were indications that the TF approximation did not support chemical bonds. Slater extended TF to treat metals by constructing the bulk as a sum of locally spherical atomic fragments.²⁶ Disappointingly, the result was repulsive energy curves. Thus it appeared that, at least for metals, TF theory could not predict binding. Years later Teller (who was a good friend of Fermi¹¹²) confirmed the difficulty rigorously³⁹ with the demonstration that molecular binding cannot happen in the TF model.

But in 1935 that problem was tackled by von Weizsäcker.²⁷ He contrived a correction to the TF model which involved the use of an ad-hoc orbital proportional to the square-root of the electron density. The kinetic energy from such an orbital certainly contains a density gradient term and, thus, introduces explicit treatment of inhomogeneity. The derivation itself is incorrect,¹¹³ but the idea is consequential.

Let us revisit the von Weizsäcker idea via an alternative development. Note that there are several other derivations, for example refs 114 and 115, that follow a density matrix-centric route. We anticipate OFDFT and suppose that one orbital is enough in some sense to capture the essential information in the wave function of the system. Such a “collective orbital” can take the form

$$\psi(\mathbf{r}) = (a + b(\mathbf{r})) e^{i\mathbf{p}\cdot\mathbf{r}} \quad (3)$$

with a a real constant such that $a \exp(i\mathbf{p}\cdot\mathbf{r})$ represents the part of the system most similar to the HEG, $b(\mathbf{r})$ is a real function that represents the deviation from uniformity, and \mathbf{p} is an average momentum associated with the collective orbital. The kinetic energy of the system can be evaluated as

$$\begin{aligned}
 T[\psi] &= \frac{1}{2} \langle \nabla \psi | \nabla \psi \rangle \\
 &= \frac{1}{2} \int d\mathbf{r} (-i\mathbf{p}a + \nabla b(\mathbf{r}) - ib(\mathbf{r})\mathbf{p}) \\
 &\quad (i\mathbf{p}a + \nabla b(\mathbf{r}) + ib(\mathbf{r})\mathbf{p}) \\
 &= \frac{1}{2} \int d\mathbf{r} (p^2 [a + b(\mathbf{r})]^2 + |\nabla b(\mathbf{r})|^2) \\
 &= \frac{1}{2} \int d\mathbf{r} (p^2 n(\mathbf{r}) + |\nabla \sqrt{n(\mathbf{r})}|^2)
 \end{aligned} \tag{4}$$

We recognize the first term, $\int p^2 n(\mathbf{r}) d\mathbf{r}$, as the TF functional by comparison with eq 1 and the realization that for the uniform system we can identify

$$p^2 = \frac{3}{5} (3\pi)^{2/3} n^{2/3} \tag{5}$$

Insertion of eq 5 in eq 4 leads to the Thomas–Fermi–von Weizsäcker (TFvW) functional,

$$\begin{aligned}
 T_{TFvW}[n] &= c_{TF} \int d\mathbf{r} n^{5/3}(\mathbf{r}) + \frac{1}{2} \int d\mathbf{r} \nabla \sqrt{n(\mathbf{r})} \nabla \sqrt{n(\mathbf{r})}
 \\
 &= T_{TF}[n] + T_{vW}[n]
 \end{aligned} \tag{6}$$

We remark that TF and TFvW can be combined with Dirac's exchange,²⁵ to lead to TFD and TFvWD, respectively. For later reference, we note

$$\begin{aligned}
 T_{vW}[n] &:= \frac{1}{2} \int d\mathbf{r} |\nabla n^{1/2}(\mathbf{r})|^2 \equiv \frac{1}{8} \int d\mathbf{r} \frac{|\nabla n(\mathbf{r})|^2}{n(\mathbf{r})}
 \\
 &\equiv \int d\mathbf{r} \tau_{vW}[n](\mathbf{r})
 \end{aligned} \tag{7}$$

At least so far as sophistication of formulation is concerned, TFvW clearly is superior to TF. But the differential equations involved are not amenable to analytical solution.³⁰ Widespread use began only decades after von Weizsäcker's original conception. Building from the work of Kirzhnitz,³¹ whose derivations led him to compute the first gradient expansion approximation (GEA) to the TF model, the first thorough probe of TFvW on realistic physical systems seems to be application to atoms by Yonei and Tomishima.^{38,116} They found that the $TF\lambda vW$ functional

$$T_{TF\lambda vW}[n] = T_{TF}[n] + \lambda T_{vW}[n] \tag{8}$$

with $\lambda = 0.2$ (rather than the GEA value of $\lambda = \frac{1}{9}$) yielded closest agreement with their quantum mechanical data, which were from the HF method with hydrogenic atomic orbitals (by the same authors).

Benguria et al.⁴¹ and later Berk⁴² did a mathematical analysis of the TFvW method and found that the von Weizsäcker correction must lead to binding between atoms. Thus, it is not surprising that investigations using TFvW theory appeared^{117,118} and have continued for decades.^{119–122}

Although more inclusive of quantum mechanical effects than TF, $TF\lambda vW$ has major drawbacks. As Figure 2 shows, no $TF\lambda vW$ variant reproduces atomic shell structure. As will become clearer below, this is a consequence of a more general limitation of $TF\lambda vW$, namely that it contains no information about the polarizability of the electronic system.⁴⁰ Similarly, Friedel oscillations¹²³ (density oscillations around an impurity in a metal) are among the so-called quantum oscillations that

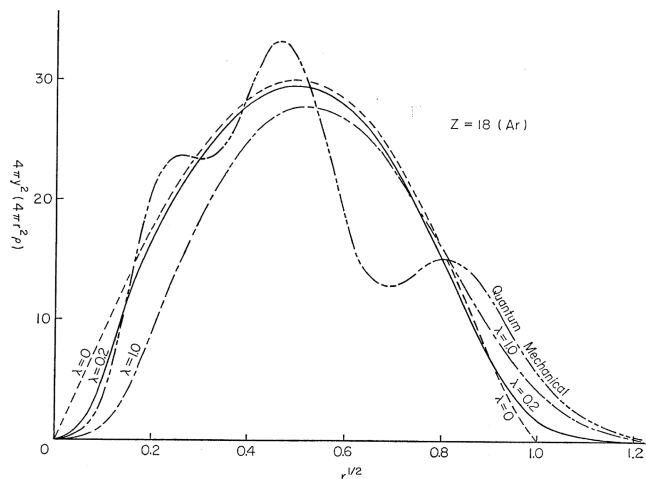


Figure 2. First probe of the $TF\lambda vW$ functional on realistic systems. Shown is the radial electron density for the Ar atom. “Quantum Mechanical” results are Hartree–Fock. Reprinted with permission from ref 38. Copyright (1965) The Physical Society of Japan.

TF and TFvW miss completely. The ability of some TFvW-type functionals to reproduce, at least semiquantitatively, the kinetic energy of atoms has been ascribed to major error cancellation.^{124,125} As we will explain in Section 2, $TF\lambda vW$ with $\lambda < 1$ are non- N -representable functionals and, therefore, are not expected to be generally accurate. Nevertheless, there have been interesting applications of $TF\lambda vW$ to metal clusters.^{119,126–128} In them, $TF\lambda vW$ was found to produce converged Carr–Parrinello or Born–Oppenheimer molecular dynamics and provide useful predictions related to melting and alloying; see Section 4.

1.3. Connecting the Early TF and TFvWD Functionals with Modern OFDFT

TF and TFvW, as already remarked, are density functionals. Given an electron density, each one gives back an energy. But the demonstration that the exact many-electron system ground state is expressible variationally as a density functional came only in 1964 with the famous Hohenberg–Kohn theorems,³ though there had been suggestive antecedents in the 1950s.^{129,130}

Only the barest essentials of DFT are needed at this point to connect TF and TFvW to modern advances. More details are in Section 2. The Hohenberg–Kohn existence theorems state that for a system of N_e electrons in an external potential $v_{ext}(\mathbf{r})$, there exists a functional of the density $\mathcal{E}[n]$ that gives the total ground-state energy E_0 variationally by

$$\begin{aligned}
 E_0[n_0] &= \min_n \{ \mathcal{E}[n] + E_{ext}[n] \} \\
 \mathcal{E}[n] &:= T[n] + U_{ee}[n]
 \end{aligned} \tag{9}$$

Here $T[n]$ is the total many-electron kinetic energy, U_{ee} is the total many-electron Coulomb interaction energy (Hartree, exchange, and correlation), and n_0 is the ground-state electron density. To make connection with the non-interacting electron concept of the TF and TFvW schemes, we insert the non-interacting kinetic energy (denoted, for reasons lost in history, as $T_s[n]$) and make a trivial addition and subtraction rearrangement

$$\mathcal{E}[n] = T_s[n] + \{ U_{ee}[n] + T[n] - T_s[n] \} \tag{10}$$

Structurally, this is the same as TFD or TFvWD: a non-interacting kinetic energy and some Coulombic (and other) corrections, all density functionals. Obviously the formal grounding and content are very different. The HK theorems prove that the TFvW, TFvWD concept is correct, even though the functionals themselves are not.

Because one-electron states (“Kohn–Sham orbitals”) commonly are introduced to make the DFT variational problem tractable by giving an explicit, calculable form for T_s , we emphasize that the form just given is “orbital-free”, OFDFT. (Remark: Shortly it will be apparent that “one orbital” would be more precise, in the same spirit as our summary of TFvW, eq 3 above. But by now “orbital-free” is standard.) The crucial distinction is that KSDFT¹ in the conventional orbital form has become the pre-eminent many-electron formulation for calculations on materials and their molecular constituents. Annual citation counts in the thousands in both quantum chemistry and materials physics¹³¹ attest to this point, as does the 1998 Nobel Prize in Chemistry. It was shared by Walter Kohn, for DFT itself,¹³² and John Pople for quantum chemical methods.¹³³ Conventional KSDFT with semilocal approximate XC functionals can produce remarkably accurate and useful results, despite known drawbacks.²

The dominant position of conventional KSDFT means that overcoming computational scaling barriers while simultaneously building more accurate XC density functional approximations (DFAs) are two critically important tasks. The KS orbitals put these two in tension. Interestingly, Pople was a late convert to DFT who focused on the explicit use of the KS orbitals in XC DFAs.¹³⁴ Conversion of the majority chemistry community soon occurred,¹³⁵ followed by a proliferation of orbital-dependent exchange-correlation DFAs. From the perspective of computational cost scaling, that proliferation is a retrograde step. Though introduction of the KS orbitals¹ was crucial because it meant that early band structure and molecular orbital codes could be adapted quite readily to DFT, the unintended consequence was the $O(N^3)$ cost scaling noted at the outset.

We fill out needed details of conventional and OFDFT forms of KSDFT in Section 2. Also there we catalog relevant exact properties of $T_s[n]$. For perspective, however, suppose again, for the moment, that one could construct all the terms in U_{ee} (the familiar challenge of XC DFAs). The remaining challenge would be to construct reliable, orbital-free approximations for $T_s[n]$ (KEDFs). That is the subject at hand. The remainder of this section is both a brief guide to the kinds of approximations that have been contrived as well as a guide to the subsequent sections of this review.

Both TF and TFW are what are called one-point functionals. Their origins in the HEG and inhomogeneity, respectively, suggest the generic form

$$T_{s,1}[n] = c_{TF} \int d\mathbf{r} n^{5/3}(\mathbf{r}) F_t(n(\mathbf{r}), \nabla n(\mathbf{r}), \nabla^2 n(\mathbf{r}), \dots) \quad (11)$$

TF ($F_t = 1$) is strictly local. Adding spatial derivative dependence is said to make a functional “semi-local”. The sense of that phrase is that while the mathematics is strictly local (one-point), the derivative dependence provides some indication of nearby density behavior. (Remark: We caution that some of the older DFT literature uses “non-local” rather than “semi-local” to describe gradient dependence. Mathematically it is strictly local.) KEDFs in which F_t depends only on n

and ∇n are called generalized gradient approximations (GGAs or also GEAs if the Kirzhnitz gradient expansion approximation³¹ is adopted; see eq 8) in correspondence with terminology already established for XC DFAs with the same dependencies. Those that depend upon $\nabla^2 n$ and higher order derivatives as well as n and ∇n are called “meta-GGAs”. As the names imply, development of these KEDFs is influenced strongly by the gradient expansion for the weakly inhomogeneous electron gas.^{117,125,136–139}

Nonlocal functionals go back to Hohenberg and Kohn,³ who proposed a solution to the inability of TF and TFvW to describe quantum oscillations by employing the Lindhard response function¹⁴⁰ for the HEG as an approximation to the electronic dielectric function. This function gives the response of the electrons at an arbitrary point to the introduction of a charge at some other point. Somewhat later, this line of thinking gave rise to investigation of nonlocal KEDFs. Most generally they would have the form

$$T_s^{\text{nl}}[n] = \int d\mathbf{r}_1 d\mathbf{r}_2 d\mathbf{r}_3 \dots d\mathbf{r}_m n^{5/3m}(\mathbf{r}_1) n^{5/3m}(\mathbf{r}_2) n^{5/3m}(\mathbf{r}_3) \times \dots n^{5/3m}(\mathbf{r}_m) \omega[n](\mathbf{r}_1, \mathbf{r}_2, \mathbf{r}_3, \dots, \mathbf{r}_m) \quad (12)$$

where $\omega(\mathbf{r}_1 \dots \mathbf{r}_m)$ is an m -body kernel to be determined. This form appears in the original Hohenberg–Kohn paper³ as a formal expansion of the exact density functional. After a brief discussion of the two-body term, they dismiss the rest: “The higher order terms will not be further discussed here.” In that same vein, so far as we are aware, the form has not been used for $m > 3$ and only once for $m = 3$.⁴⁴ In that case, the three-point structure was induced by a tour de force perturbation approach that strongly resembles the iterative form of the Dyson equation¹⁴¹ for the two-particle Green function.

Since the two-particle reduced density matrix (2-rdm) determines all the ground-state properties of a Coulombic fermion system,¹⁴² it similarly is appropriate to focus efforts at direct (nonperturbative) construction of KEDFs upon the two-point case. For reasons of physical clarity and constraint satisfaction (see Section 2), it is convenient to write two-point KEDFs in the generic form

$$T_{s,2}[n] = T_{vW}[n] + T_{TF}[n] + \int d\mathbf{r} d\mathbf{r}' \mathcal{K}[n(\mathbf{r}), n(\mathbf{r}'); \mathbf{r}, \mathbf{r}'] \quad (13)$$

[Remark: For simplicity, here we introduce nonlocal functionals through eq 13. We acknowledge that there are other formulations that preserve the structure of the T_{TF} functional (e.g., the Chacón–Alvarellos–Tarazona (CAT) family of functionals^{43,143}) and the structure of the T_{vW} functional (e.g., ref 144). These functionals are well-reviewed in refs 145 and 146. See further discussion in Section 2.] This form incorporates the exactness of T_{TF} in the uniform limit and of T_{vW} in the one-electron or two-electron singlet limit (see Section 2). The explicitly nonlocal contribution then is a sophisticated interpolation between those two extreme cases. Typically it has the structure of a density-weighted kernel

$$\mathcal{K}[n(\mathbf{r}), n(\mathbf{r}'), \mathbf{r}, \mathbf{r}'] = n^\alpha(\mathbf{r}) \omega[n](\mathbf{r}, \mathbf{r}') n^\beta(\mathbf{r}') \quad (14)$$

with α and β constants. Even though eq 13 is formally an approximation, it can be made exact (as can be seen by integration of eq 12 of all coordinates except two) using the general form of the kernel given on the rhs of eq 14. In practice, the kernel is determined, for example, by calibration

to linear response of the HEG (Lindhard response)¹⁰⁵ and the resulting KEDFs is approximate. We return to this in detail below.

Two-point KEDFs generally are expected to be more accurate than one-point KEDFs simply because the two-pointers probe and encode information about a larger spatial region than is possible with one-point functionals. Figure 3

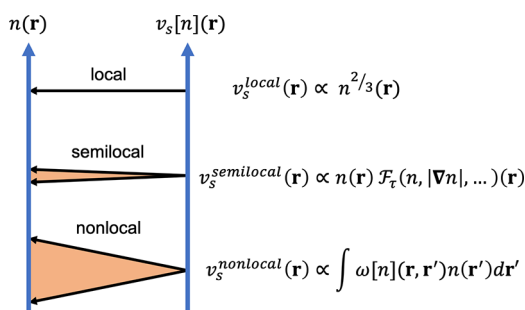


Figure 3. Conceptual depiction of the dependence on the electron density of local, semilocal, and two-point functionals. The corresponding potentials are depicted in the figure. Nonlocal or two-point functionals of $n(\mathbf{r})$ depend on an arbitrary neighborhood of \mathbf{r} whereas semilocal functionals only sample infinitesimally nearby and local functionals sample the single point, \mathbf{r} .

provides a simple pictorial comparison. That is, from local one-point, to semilocal one-point (GGA, metaGGA, ...), to two-point functionals, the kinetic energy potential, defined as $v_s(\mathbf{r}) = \frac{\delta T[n]}{\delta n(\mathbf{r})}$, is informed by a larger and larger neighborhood of the spatial point \mathbf{r} . TF and TFvW are extreme examples of awareness of only the point \mathbf{r} . Their forms are inadequate for diverse reasons which will be discussed in Section 2. Note, however, that the curse of computational scaling intrudes because two-point KEDFs depend on the density–density response function of the system; see Sections 2 and 3.

Machine-learnt (ML) KEDFs are the most recent type. They are discussed in Subsection 2.5. Because their explicit form typically is not known, they do not fit easily into the one-point, two-point categorization, though some are built specifically to have a certain maximum order of density gradients. Just as machine learning itself is relatively recent, so also the literature of ML KEDFs is comparatively recent and small.^{83,84,147–160}

The imperative for access to the full periodic table with high-quality KEDFs also has motivated efforts such as angular momentum-dependent¹⁶¹ and projector augmented wave (PAW) implementations¹²⁰ of OFDFT. We discuss those briefly at the end of Section 2, along with information-theoretical and other KE approaches. We conclude Section 2 with a brief discussion of some pitfalls of KEDF testing.

Implementation, algorithms, and associated issues are the subject of Section 3. For two-point functionals, implementation is simplified greatly by the employment of regular grids at any reasonable spacing, reciprocal space methods, and Fast Fourier Transforms (FFTs).^{53,55} FFT grids also can be used to encode one-point KEDFs, including free-energy versions.¹⁶² Irrespective of KEDF type, however, regular grids are unable to resolve the singular Coulomb potentials originating from point nuclear charges. Thus, several pseudopotentials have been developed.^{48,163–165} Those are, of necessity, local in the sense of pseudopotential terminology, that is, orbital-independent. This issue is crucial for expanded applicability of OFDFT, whatever the type of KEDF. We address it in Subsection 3.1.

In Section 4, we discuss a nonexhaustive selection of illustrative applications. Here we simply note that recent applications with one- and two-point KEDFs include warm dense matter^{166–168} and molten and zero-temperature metals and semiconductors (bulk,^{86,169–172} surfaces and other low-dimensional systems,¹⁷³ and clusters^{88,126,174}).

We conclude this section with a note on computational scaling of OFDFT versus conventional KSDFT. It is critical to remember that both use the KS decomposition, eq 10, to invoke a specific non-interacting reference system that generates the density. Conventional KSDFT uses the explicit kinetic energy expression for that system, $T_s[\{\varphi_i[n]\}]$, in terms of the N_e orbitals in the KS determinant, $\varphi_i[n](\mathbf{r})$. Obtaining those orthogonal orbitals when using semilocal XC functionals requires a formal $O(N_e^3)$ computational cost (from diagonalizing the KS Hamiltonian matrix) and a $O(N_e^2)$ memory requirement (from storing the wave functions and matrices involved). Therefore, several approaches attempt to evade the cubic scaling of standard implementations of conventional KSDFT by avoiding the diagonalization of the KS Hamiltonian and leveraging sparsity in a variety of different ways.

Other than OFDFT, several propositions have been put forth for achieving sub-cubic scaling (i.e., $O(N_e)$ or $O(N_e^2)$) of

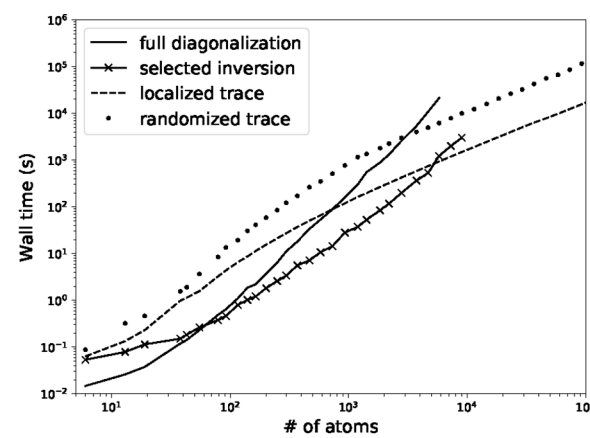
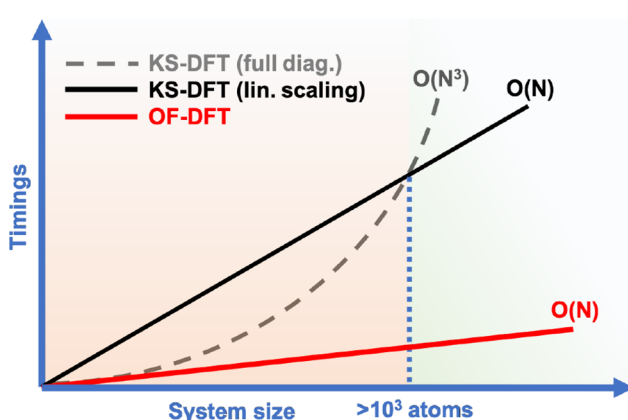


Figure 4. Estimation of the computational scaling of OFDFT (left panel) and conventional KSDFT (right panel) employing both cubic-scaling diagonalization and different sub-cubic-scaling approaches. Conventional KSDFT data from Moussa and Baczewski 2019.¹⁷

conventional KSDFT by combining algorithmic advances with efficient utilization of modern high-performance computer architectures.^{10,11,13,17,175–183} Underlying all such schemes, however, is exploitation of nearsightedness in the electronic structure^{184–186} to achieve sparse matrices (e.g., by screening certain one- and two-electron integrals or by effective cutoffs applied to the off-diagonal elements of the one-body reduced density matrix, 1-rdm), coupled with efficient solutions to the Poisson equation.

Sub-cubic-scaling approaches to conventional KSDFT are dependent on temperature and on the energy gap of the material. For example, for systems with gap, the 1-rdm decays exponentially, thus sub-cubic-scaling methods are very effective for these systems. Instead, for metals at zero and low temperatures, the 1-rdm decays slowly (algebraically, see eqs 117–119). Thus, sub-cubic-scaling methods are less effective in this regime. However, because at higher temperature the density matrix of metals decays exponentially,^{187,188} methods exploiting the sparsity properties of the 1-rdm can be useful and applicable also to metals.

We give an example of the performance of sub-cubic-scaling methods on the right-hand side of Figure 4, where we reproduce data from ref 17. On the left-hand side of the figure, we depict the expected timings for OFDFT compared with linear-scaling KSDFT and its straightforward implementations. The figure shows that both OFDFT and linear-scaling KSDFT are expected to yield linear timings with system size but with very different prefactors.

Given the advantages of OFDFT, the patient reader rightly may ask why it has not supplanted conventional KSDFT. The answer is that, until recently, adequately accurate KEDFs have not been constructed. As we show below, that situation is changing, we believe, rapidly. It is the purpose of this review to report that progress and outlook for achieving the promise of OFDFT in chemical and materials modeling. Additionally, recent years have witnessed the emergence of several schemes that combine conventional KSDFT and OFDFT. Subsystem DFT^{180,181,189–191} is one such scheme that is proving to be promising for several classes of systems.

For context, we note several prior OFDFT reviews (in addition to the Thomas–Fermi literature already cited): refs 115, 145, and 192–194 and a book.¹⁹⁵ A convenient listing of one-point KEDFs is in Chapter 16 of that book. Also see the University of Florida online listing.¹⁹⁶ To have some control on the scope of this article, we do not review free-energy (finite temperature) OFDFT except for a bit on successful computational use.¹⁶⁸ For that topic, see refs 194 and 197. Nor do we consider time-dependent OFDFT^{198–209} or OFDFT on a lattice.²¹⁰

2. KINETIC ENERGY FUNCTIONALS

This section sets forth DFT in pertinent form, with emphasis on known constraints, limits, and conditions relevant to constructing approximate KEDFs. Then we survey the specific approximations of various kinds (one-point, two-point, machine-learnt) and some of their strong and weak points. There follow some observations on testing, followed by a survey of various “road-less-taken” approaches to recognize angular momentum dependence, apply information theory, and convert to potential functional theory. We conclude the section with a look at OFDFT in the context of embedding schemes (subsystem DFT).

2.1. DFT Basics

For clarity about kinetic energy contributions and notation, we summarize the basic Hohenberg–Kohn (HK) DFT theorems³ and the KS construction¹ for spin-unpolarized systems. Details are in refs 102, 106, and 211. Nonrelativistic spin-polarization follows easily.²¹² For simplicity of presentation and notation therefore, the presentation is spin-unpolarized unless remarked to the contrary. Though DFT is general, chemical and materials interest is in the electronic structure. To that we specialize.

The Hamiltonian for N_e electrons in a fixed external field $v_{ext}(\mathbf{r})$ is

$$\hat{H} := \sum_i^{N_e} \left[-\frac{1}{2} \nabla_i^2 + v_{ext}(\mathbf{r}_i) \right] + \frac{1}{2} \sum_{i \neq j}^{N_e} \frac{1}{|\mathbf{r}_i - \mathbf{r}_j|} \quad (15)$$

in Hartree atomic units. For much of chemistry and materials physics, $v_{ext}(\mathbf{r})$ is the Coulomb potential of a fixed array of nuclei; hence

$$E_{ext}[n] = \int d\mathbf{r} n(\mathbf{r}) v_{ext}(\mathbf{r}) \equiv - \sum_{\alpha} Z_{\alpha} \int d\mathbf{r} \frac{n(\mathbf{r})}{|\mathbf{r} - \mathbf{R}_{\alpha}|} \quad (16)$$

with nuclear charges Z_{α} and positions \mathbf{R}_{α} . (Remark: Harmonic confinement potentials also have been useful for studying approximate functionals. See for example refs 213–221, among others.)

As already noted at eq 9, the HK theorems prove the existence and key properties of a functional of the electron number density $n(\mathbf{r})$,

$$\mathcal{E}[n] = T[n] + U_{ee}[n] \quad (17)$$

Recall, from Section 1, that $T[n]$ and $U_{ee}[n]$ are the *total* many-electron kinetic energy (KE) and Coulomb interaction energy, respectively. The number density integrates to the total electron number

$$\int d\mathbf{r} n(\mathbf{r}) = N_e \quad (18)$$

Because $\mathcal{E}[n]$ is independent of the external potential $v_{ext}(\mathbf{r})$, the functional is said to be “universal”. The Rayleigh–Ritz variational principle gives the ground-state energy E_0 and density n_0 as

$$\min_n \{ \mathcal{E}[n] + E_{ext}[n] \} = E_0[n_0] \quad (19)$$

(Remark: A mathematically careful description of the minimization involves differently defined functionals and their extrema. Formally there are three functionals, with different variational properties and different functional derivatives. For most chemistry and physics applications, the subtle mathematical differences among them do not matter enough to be distracted by them here. In this we follow the precedent of Engel and Dreizler,¹⁰⁶ p 36: “As a matter of principle, the subsequent development of the DFT formalism should therefore be based explicitly on the Lieb functional. We will nevertheless ignore the issue...and not distinguish between the various flavors...”.)

The HK theorems prove the existence of $\mathcal{E}[n]$ but do not supply an explicit constructive route to it. That fact motivates the connection to non-interacting systems already sketched in Section 1 at eq 10. The connection is made via the insight of Kohn and Sham,¹ namely that the HK theorems apply equally well to non-interacting systems.^{106,222} Thus, for most physical

systems, it is possible to invoke a special one-electron potential, the Kohn–Sham potential $v_{KS}(\mathbf{r})$, that causes a non-interacting electron system to have exactly the same electron density as the interacting physical system. Those systems for which $v_{KS}(\mathbf{r})$ exists are designated as non-interacting v -representable or simply v_{KS} -representable.¹⁰⁶

Because the KS system is a non-interacting many-electron system, its variational wave function is a single determinant with orbitals generated by¹

$$\hat{h}_s[n]\varphi_i(\mathbf{r}) = \varepsilon_i\varphi_i(\mathbf{r}) \quad (20)$$

$$\hat{h}_s[n](\mathbf{r}) := -\frac{1}{2}\nabla^2 + v_{KS}(\mathbf{r}) \rightarrow \hat{H}_s = \sum_i^{N_e} \hat{h}_s(\mathbf{r}_i) \quad (21)$$

$$n(\mathbf{r}) = \sum_i f_i |\varphi_i(\mathbf{r})|^2 \quad (22)$$

where \hat{H}_s is the many-Fermion KS Hamiltonian. At zero temperature the occupation numbers f_i ordinarily are integers (0, 1, or 2). An exception is the case in which the highest occupied level is degenerate.^{211,223} For fixed density, $n(\mathbf{r})$, the minimization of the KS energy (i.e., $E_{KS} = \langle \Phi | \hat{H}_s | \Phi \rangle$) is with respect to all single-determinant wave functions, $\Phi(\mathbf{r}_1 \dots \mathbf{r}_{N_e})$, that give $n(\mathbf{r}) = \langle \Phi | \hat{n}(\mathbf{r}) | \Phi \rangle$, with $\hat{n}(\mathbf{r}) = \sum_i^{N_e} \delta(\mathbf{r}_i - \mathbf{r})$. The minimizing one that yields the ground-state density $n_0(\mathbf{r})$ is the KS determinant $\Phi_{KS}(\mathbf{r}_1 \dots \mathbf{r}_{N_e})$. Note that the minimization is solely that of the KE expectation value, since the KE is the only many-body operator in the non-interacting Hamiltonian and the density is fixed.

The KS determinant therefore gives an explicit expression for the non-interacting kinetic energy $T_s[\{\varphi_i[n]\}]$:

$$\begin{aligned} T_s[\{\varphi_i[n]\}] &:= \frac{1}{2} \sum_i f_i \int d\mathbf{r} |\nabla \varphi_i[n](\mathbf{r})|^2 \equiv \int d\mathbf{r} \tau_s[n](\mathbf{r}) \\ &\equiv T_s[n] \end{aligned} \quad (23)$$

This object is the center of attention in OFDFT. (Remark: Notice that we use the positive definite form of the KS KE integrand, not the more commonly encountered $T_s[n] = -\frac{1}{2} \sum_i f_i \int d\mathbf{r} \varphi_i^*(\mathbf{r}) \nabla^2 \varphi_i(\mathbf{r})$. The two differ by a surface term that integrates to zero in virtually all cases.)

The KS determinant also defines the DFT exchange energy $E_x[n]$ (for spin unpolarized systems):

$$\begin{aligned} E_x[n] &:= -\frac{1}{4} \sum_{i,j} f_i f_j \\ &\times \int d\mathbf{r}_1 d\mathbf{r}_2 \frac{\varphi_i^*[n](\mathbf{r}_1) \varphi_j^*[n](\mathbf{r}_2) \varphi_i[n](\mathbf{r}_2) \varphi_j[n](\mathbf{r}_1)}{|\mathbf{r}_1 - \mathbf{r}_2|} \end{aligned} \quad (24)$$

It also is a density functional via the fact that the KS orbitals are density functionals.

The remaining step is to connect the physical and KS systems explicitly. In the expression for the total energy of the physical system, it is conventional to identify the classical Coulomb repulsion contribution (the Hartree energy) to the total Coulomb energy

$$E_H[n] := \frac{1}{2} \int d\mathbf{r}_1 d\mathbf{r}_2 \frac{n(\mathbf{r}_1)n(\mathbf{r}_2)}{|\mathbf{r}_1 - \mathbf{r}_2|} \quad (25)$$

The universal density functional eq 17 for the physical system then can be rearranged to expose the non-interacting and classical pieces as

$$\begin{aligned} \mathcal{E}[n] &= T_s[n] + E_H[n] + E_x[n] \\ &+ \{U_{ee}[n] - E_H[n] - E_x[n] + T[n] - T_s[n]\} \end{aligned} \quad (26)$$

$$:= T_s[n] + E_H[n] + E_x[n] + E_c[n] \quad (27)$$

This equation is the KS version of eq 10. The last term in eq 27 defines the KSDFT correlation energy as the sum of the final five terms of eq 26. In particular, the difference $T[n] - T_s[n]$ is lumped into E_c along with the Coulomb correlation. Though exchange and correlation are defined separately, they can be obtained (formally and operationally) together via the adiabatic connection.^{212,224} It also is common to develop an approximate exchange functional paired with an approximate correlation functional in order to achieve useful cancellation. See, for one much-used example, the PBE approximation.²²⁵ Sometimes exchange and correlation are approximated together in nonseparable form $E_{xc} = E_x + E_c$.²²⁶

Regardless of whether the functionals are expressed in terms of the KS orbitals or directly in terms of $n(\mathbf{r})$, the variational minimization of $\mathcal{E}[n]$ must be done over electron densities that integrate to N_e . That is equivalent to minimizing the Lagrangian

$$\mathcal{L}[n] = \mathcal{E}[n] + \int d\mathbf{r} n(\mathbf{r}) v_{ext}(\mathbf{r}) - \mu \left[\int d\mathbf{r} n(\mathbf{r}) - N_e \right] \quad (28)$$

The Euler–Lagrange equation follows after taking the functional derivative with respect $n(\mathbf{r})$ and requiring that it be null at all \mathbf{r} , namely,

$$\begin{aligned} \frac{\delta T_s[n]}{\delta n(\mathbf{r})} + \frac{\delta E_H[n]}{\delta n(\mathbf{r})} + \frac{\delta E_{xc}[n]}{\delta n(\mathbf{r})} + v_{ext}(\mathbf{r}) \\ \equiv \frac{\delta T_s[n]}{\delta n(\mathbf{r})} + v_{KS}(\mathbf{r}) = \mu \end{aligned} \quad (29)$$

Here μ is the electron chemical potential. The last equality comes from the fact that the physical and KS systems have the same chemical potential. That always holds because the potential v_{KS} is determined only up to a constant.

Thus the Kohn–Sham potential in eq 21 is

$$\begin{aligned} v_{KS}(\mathbf{r}) &= v_H(\mathbf{r}) + v_{xc}(\mathbf{r}) + v_{ext}(\mathbf{r}) \\ v_H(\mathbf{r}) &= \int d\mathbf{r}_2 \frac{n(\mathbf{r}_2)}{|\mathbf{r} - \mathbf{r}_2|}, \quad v_{xc}(\mathbf{r}) = \frac{\delta E_{xc}[n]}{\delta n(\mathbf{r})} \end{aligned} \quad (30)$$

For a fixed set of nuclei (ions, e.g. at each step in ab initio molecular dynamics, AIMD^{227–229}), v_{ext} is as in eq 16 and the electronic force on a given nucleus α is $-\nabla_{\mathbf{R}_\alpha}(\mathcal{E}[n_0] + E_{ext}[n_0] + E_{NN})$. Here $E_{NN}(\{\mathbf{R}\})$ is the internuclear (ion–ion) repulsion energy.

2.2. DFT Context Issues

Before proceeding, some context regarding present-day use of DFT, particularly in chemistry, is relevant. The vast majority of applications use conventional KSDFT, hence $T_s[n]$ in its explicitly orbital-dependent form, eq 23. The only approximate functionals of interest in that case are $E_{xc}[n] = E_x[n] + E_c[n]$. That is why much of the DFT literature uses the phrase

“density functional approximations” to mean XC approximations.

In contrast, OFDFT focuses on approximate expressions for $T_s[n]$ that do *not* rely on explicit evaluation of the orbital-dependent expression, $T_s[\{\varphi_i[n]\}]$, in eq 23. A consequence is clear. If OFDFT is to grow in utility, there must be sustained, concurrent development of improved orbital-free E_{xc} DFAs. Present-day materials physics practice and priorities illustrate the point. In that arena, the GGA rung of the Perdew–Schmidt²³⁰ ladder of XC DFA complexity dominates, with the PBE²³⁵ functional most widely used. A GGA XC functional has the generic, orbital-free form

$$E_{xc}^{GGA}[n] = \int d\mathbf{r} e_{xc}[n(\mathbf{r}), s] \quad (31)$$

where $e_{xc}[n(\mathbf{r}), s]$ is an XC energy density, the details of which are irrelevant here, except that it depends on both n and the dimensionless reduced density gradient

$$s(\mathbf{r}) := \frac{|\nabla n(\mathbf{r})|}{(3\pi^2)^{1/3} n^{4/3}(\mathbf{r})} \quad (32)$$

(Remark: Beware! In the chemistry literature for XC DFAs it is fairly common to define this variable *without* the constants. Occasionally that form shows up in the OFDFT literature as well.)

The contextual challenge that we do *not* address in this review is the explicit use of KS orbitals in $E_{xc}[n]$ approximations that are on higher rungs of the Perdew–Schmidt complexity ladder than GGAs. In the case of meta-GGA XC functionals, for example, explicit orbital dependence enters via the KS KE density $\tau_s[n(\mathbf{r})]$ as an ingredient. Recent popular examples include the SCAN $E_{xc}[n]$ DFA²³¹ and its successor, r^2 SCAN.²³² With certain exceptions, those two are broadly applicable to both molecules and extended systems. Some progress has been made in removing the $\tau_s[n(\mathbf{r})]$ dependence from both SCAN and r^2 SCAN.^{233–235} In stark contrast, the quantum chemistry community has sought accuracy improvements largely by moving to yet higher rungs on the Perdew–Schmidt ladder, specifically with hybrid E_{xc} DFAs, i.e. those that include a single-determinant exchange, eq 24, contribution.

Recall our earlier remark about Pople’s coming to DFT via interest in a hybrid XC functional. Inclusion of some single-determinant exchange introduces explicit orbital dependence in $E_{xc}[n]$. That orbital dependency seems to be completely at odds with the OFDFT objective of eliminating such dependence from $T_s[n]$. Conceptually the use of orbital-dependent E_{xc} DFAs poses no issues in conventional KSDFT, as the optimized effective potential (OEP) procedure^{236,237} can be used to retrieve the multiplicative KS potential. In practice, however, OEP requires considerable extra computation, so it is little used with orbital-dependent XC DFAs. The generalized KS Euler equation is used instead (i.e., in a short-hand notation, replacing $\frac{\delta E_{xc}}{\delta n} \frac{\delta n}{\delta \phi_i}$ by $\frac{\delta E_{xc}}{\delta \phi_i}$). Observe that the KS and generalized KS equations are not equivalent.^{238,239} Anyway, OEP is irrelevant here except to illustrate the potential payoff of OFDFT in an almost sardonic way. Because there is no orbital manifold in OFDFT, there is no generalized KS equation and, hence, no need for the elaborate machinery of OEP.

The irony of emphasis on orbital-dependent XC DFAs notwithstanding, from here forward we assume that the $E_{xc}[n]$

DFA is orbital-free. The OFDFT task then is to eliminate the explicit orbital dependence of $T_s[n]$ by constructing approximations to it that depend purely on the electron density, n .

Note also that the kinetic energy approximations discussed here are for $T_s[n]$, not for $T[n]$. Formally the focus upon $T_s[n]$ is to exploit the power of the KS logic of connection to non-interacting systems. Doing so avoids the formidable problem that an attempt to approximate $T[n]$ would amount to an attempt to approximate the entire $\mathcal{E}[n]$ by virtue of the Coulomb virial theorem. (This argument is not original; it was pointed out to SBT by So Hirata in a University of Florida seminar.) Practically, the reason for addressing $T_s[n]$ is that modern $E_{xc}[n]$ DFAs all have been developed in the KS context, i.e., the decomposition in eq 27.

With both $T_s[n]$ and $E_{xc}[n]$ approximations orbital-free, the constrained minimization can be carried out directly via the single variational quantity $n(\mathbf{r})$. See eq 29. This is the justification of the remark at the outset of Section 1 that OFDFT restores the centrality of the density in DFT.

2.3. Constraints

Because there are no mechanical procedures (e.g., order-by-order perturbation expansion) provided by the basic DFT theorems to develop approximate KEDFs, it is important to catalog provable properties of the exact functional $T_s[n]$. Such rigorous results are useful as constraints on KEDF development. An example is the concise list of constraints in ref 240.

The simplest relation is between the full KE $T[n]$ and the non-interacting $T_s[n]$. Because $T_s[n]$ is defined to be the minimum fermion expectation value of the many-fermion KE operator for specified n (recall above), it follows that

$$T_s[n] \leq T[n] \quad (33)$$

Constraints often are expressed in terms of $T_{TF}[n]$,^{22,23,241} eq 2, and $T_{vW}[n]$,²⁷ eq 6. Though $T_{TF}[n]$ does not support bound systems (recall Section 1 and ref 39), its simplicity and its exactness for the HEG have made it popular as a plausible initial approximation against which to make corrections. From the perspective of rigorous constraints, that is not, however, the optimal approach. The reason is clear from consideration of $T_{vW}[n]$. It is exact for a fermion system of one orbital (one electron or two electrons in a singlet state). Therefore, $T_{TF}[n]$ is simply wrong for the H atom and for the density associated with the tail region of any isolated molecule or atom.

$T_{vW}[n]$ also is exact for the ground state of any many-boson system, which leads to the observation that fermion antisymmetrization must raise the non-interacting KE over the corresponding bosonic case; hence

$$T_{vW}[n] \leq T_s[n] \quad (34)$$

This lower bound can be proven simply.^{242–245} We return to the point in the discussion of information-theoretic KEDFs below.

The T_{vW} lower-bound property motivates the so-called Pauli-term decomposition

$$T_s[n] = T_{vW}[n] + T_\theta[n], \quad T_\theta[n] \geq 0 \quad (35)$$

as a route to approximate KE functionals. The rigorous non-negativity^{246–249} of $T_\theta[n]$ is a powerful constraint on any approximate KEDF, either one- or two-point. Moreover, the functional derivative of $T_\theta[n]$ must be positive semidefinite²⁴⁹

$$\nu_\theta(\mathbf{r}) := \frac{\delta T_\theta[n]}{\delta n(\mathbf{r})} \geq 0, \quad \forall \mathbf{r} \quad (36)$$

Because this is a pointwise requirement, it is a particularly stringent constraint on approximate KEDF development.

The Pauli term decomposition enables the rewriting of the DFT Euler–Lagrange equation (eq 29) in a particularly useful way. Substitution of (35) in (29) and recognition that

$$\frac{\delta T_{vW}[n]}{\delta n(\mathbf{r})} = -\frac{1}{2}n^{-1/2}(\mathbf{r})\nabla^2 n^{1/2}(\mathbf{r}) \quad (37)$$

gives

$$\left\{ -\frac{1}{2}\nabla^2 + \nu_{KS}(\mathbf{r}) + \nu_\theta(\mathbf{r}) \right\} n^{1/2}(\mathbf{r}) = \mu n^{1/2}(\mathbf{r}) \quad (38)$$

This one-orbital expression is the OFDFT Kohn–Sham equation. It differs from the conventional KS equation (eq 22) by only a local potential, an advantage unattainable if one were to start from TF and add corrections.

A third, closely related constraint comes into play, namely

$$\nu_\theta(\mathbf{r}) \geq \frac{\tau_\theta(\mathbf{r})}{n(\mathbf{r})}, \quad \forall \mathbf{r} \quad (39)$$

$$\tau_\theta := \tau_s - \tau_{vW} \quad (40)$$

In this expression, the pointwise positive semi-definiteness of the Pauli term, $\tau_\theta \geq 0$ everywhere makes the constraint on ν_θ more stringent than the constraint of its basic semi-definiteness (eq 36).

A related constraint is on the limiting behavior of the Pauli energy density. In terms of the dimensionless reduced density gradient s , eq 32, the constraint is

$$\lim_{s \rightarrow \infty} \tau_\theta = 0 \quad (41)$$

This requirement arises from the fact that the density tail of an isolated many-electron atom is dominated by a single orbital that decays exponentially; hence, $s \rightarrow \infty$. For a single-orbital system, the von Weizsäcker functional is exact and eq 41 follows.

An upper-bound constraint to T_s that has been conjectured but apparently never proved^{250,251} also is useful. It is

$$T_s[n] \leq T_{TF}[n] + T_{vW}[n] \quad (42)$$

This inequality can be rationalized²⁵² by taking the $N_e \rightarrow \infty$ limit of the finite-system inequality due to Gázquez and Robles.³⁶ Note, however, that their inequality involves a local density approximation and hence is not an exact result. Violations of eq 42 have been discussed recently.²⁵³

Uniform density scaling²⁵⁴ is a property that any proper KEDF must have. For the scaled density $n_\lambda(\mathbf{r}) := \lambda^3 n(\lambda\mathbf{r})$ with λ a real, positive constant, the KS KE satisfies

$$T_s[n_\lambda] = \lambda^2 T_s[n] \quad (43)$$

The TF KE satisfies the scaling relation, which motivates the formulation of other KEDFs in terms of dimensionless factors multiplying τ_{TF} . A remarkably early example, prior to the well-known ref 254, is the explicit discussion of uniform density scaling in a KEDF by Alonso and Girifalco.²⁵⁵ Somewhat after ref 254, there is mention of this scaling in ref 44 in the context of the second kind of kernel they proposed (see their eq 3.39).

A different kind of constraint is associated with the large- Z limit of the neutral atom energy,²⁵⁶ (Z is the atomic number;

recall eq 16). Asymptotically in Z , the KS KE has the expansion

$$T_s = b_0 Z^{7/3} + b_1 Z^2 + b_2 Z^{5/3} + \dots \quad (44)$$

with $b_0 = 0.768745$, $b_1 = -1/2$, and $b_2 = 0.269900$. Reference 256 gives a procedure by which to extract the asymptotic behavior of a given candidate KEDF from atomic calculations. Comparison with the exact coefficients then provides a route to evaluate or refine the parameters in the candidate KEDF.

An important condition that often is ignored is the N -representability of a KEDF. Regarding N -representability, it is helpful to start with the density itself. The definition of what qualifies as a density²⁵⁷ is any smooth, differentiable function $f(\mathbf{r}) > 0 \forall \mathbf{r}$ (subject to boundary conditions) that integrates to N_e on the volume of interest and has finite von Weizsäcker KE on that volume. All such candidate densities are themselves N -representable; that is, for each such candidate there is an N_e fermion wave function that yields the specified density.^{244,258}

N -representability of functionals is a bit different. The concept is made precise in ref 259, theorem 4. It proves that if a KEDF delivers a KE below the conventional KSDFT value (obtained with the exact orbital-dependent T_s functional) for any system, that KEDF is not N -representable because there are no N -representable two-body density matrices (hence, no N_e fermion wave functions) that could yield that energy when traced with the Coulomb Hamiltonian plus ν_{ext} .

The risk in use of a non- N -representable KEDF is that it can yield energies below, even grossly so, the correct conventional KSDFT value. Ayers and Liu point out,²⁵⁹ for example, that T_{vW} is non- N -representable for electron densities integrating to 3 or more electrons because T_{vW} is a lower bound to T_s for those systems (recall above). Further, the TF plus scaled von Weizsäcker functional, $TF\lambda vW$, eq 8, is non- N -representable for $\lambda < 1$. They illustrate with the ground-state electron density of the xenon atom. For $\lambda = 1/9$ the result is "...far below the true answer: $T_{TF}(1/9) = 7083, \dots T_{accurate} = 7232$ hartree". Note also that, unhappily for developers of KEDFs, N -representability of a KEDF does not guarantee accuracy. An approximate KEDF could be N -representable simply by being a loose upper bound to T_s .

In contrast to that somewhat gloomy observation, we make two points. While it may not be straightforward to impose N -representability as a constraint in the course of constructing a KEDF, it is straightforward to test N -representability after the fact. Failures on simple systems should, at the least, be viewed as a warning. The other point is that constraints from N -representability have been developed for weighted-density approximation KEDFs based on model 1-rdms. See Chakraborty et al.^{260,261} and the discussion below.

Useful constraints on KEDFs also can come from known properties of the densities associated with certain classes of ν_{ext} . For ν_{ext} from a set of point nuclei, the density is known to satisfy the Kato cusp condition²⁶² at each nuclear site. For GGA and meta-GGA KEDFs, that has immediate, specific consequences for the form of the KS potential, $\nu_{KS}(\mathbf{r})$.^{249,263–267} Systems in which the physical (i.e., ionic) external potential has been replaced by a pseudopotential have, by construction, no cusp at each nuclear site but typically zero gradient instead. That has different consequences for KEDF construction.^{86,265,268} Observe that exploiting these distinctions introduces non-universality, in the sense already mentioned. In principle, the density functional is indifferent to the ν_{ext} and, hence, universal. In practice, an effective

strategy is to design a KEDF particularized to a specific class of ν_{ext} .

A less severe form of non-universality arises in another way. OFDFT calculations require the specification of both an approximation for T_s and an approximation for the XC functional E_{xc} . Another constraint ensues. Those two functionals are connected by the virial relationship²⁵⁴

$$T_s = \frac{1}{2} \int d\mathbf{r} n(\mathbf{r}) \mathbf{r} \cdot \nabla \nu_{KS}(\mathbf{r}) \quad (45)$$

Here ν_{KS} is the full KS potential, eq 30, which, of course, includes ν_{xc} . The implication is that if ν_{xc} is from an XC DFA, as virtually always is the case, then the KEDF to be used on the left-hand side should be constructed paired with that specific XC DFA. Otherwise, there is no guarantee that substitution of n_0 would satisfy that equation. One sees that pairing in the Kohn–Sham equation in fact. In it, up to a constant, the functional derivative of T_s evaluated at the equilibrium density n_0 is the negative of $\nu_{KS}[n_0]$. See also the related discussion in ref 269.

So far as we are aware, this pairing or matching requirement has been imposed strictly on KEDF development only once. The KEDF in ref 255 was developed from their XC approximation. A closely related, converse procedure, deorbitalization of a τ_s -dependent XC functional, was tried first by Perdew and Constantin²⁷⁰ and pursued more thoroughly by Mejía-Rodríguez and Trickey,^{233–235} but neither used the matching as a constraint.

There are several reasons for not using the virial form (but see below about potential functionals). The integrand on the right-hand side of eq 45 is not translationally invariant,²⁷¹ so one cannot identify the KE density τ_s with that integrand. For testing a candidate KEDF, the expression can be used directly with atoms (and probably should be). The invariance problem can be evaded via a translationally invariant KE density that integrates to the same value as the original translation-dependent form.²⁷² But a basic gauge problem remains. Anything that integrates to zero can be added to the integrand on the rhs of eq 45 without changing the functional relationship, so extraction of a KE density is ambiguous.^{271,272}

2.4. Orbital-Free T_s Approximations

In Section 1, we distinguished the three main KEDF types of contemporary interest, semilocal or one-point functionals, $T_{s,1}[n]$, two-point functionals, $T_{s,2}[n]$, and machine-learnt, $T_{s,ML}[n]$. As noted just after eq 13, one-point functionals generically are

$$T_{s,1}[n] = \int d\mathbf{r} \tau_s[n(\mathbf{r}), \nabla n(\mathbf{r}), \dots] \quad (46)$$

Recall from Section 1 that two-point functionals typically are written as interpolations between T_{TF} and T_{vW} in order to incorporate their exactness properties explicitly. Specifically,

$$T_{s,2}[n] = T_{vW}[n] + T_{TF}[n] + \int d\mathbf{r} d\mathbf{r}' \mathcal{K}[n(\mathbf{r}), n(\mathbf{r}'), \mathbf{r}, \mathbf{r}'] \quad (47)$$

with \mathcal{K} as in eq 14 of Section 1.

Machine-learnt functionals may combine both one-point and two-point contributions in ways that are not expressible in simple analytical form. As such, they may have characteristics that are not easy to analyze in the context of the conventional one-point and two-point formulation.

We consider each type in turn. At the end of the section we consider briefly some other variants.

Before proceeding, a remark on “empirical” functionals is in order. In the development of XC DFAs, it is customary to designate functionals that have multiple parameters fitted to substantial databases (experimental or calculated data or both) as “empirical”. In contrast, those DFAs that have parameters constrained by values calculated for small sets of atomic systems (“appropriate norms” per ref 231) are categorized as “nonempirical”. We follow that terminology here. Akin with DFA development, however, the construction of a KEDF involves a considerable element of design choice. There simply are not enough known constraints to determine a KEDF completely, so building in desirable physics or conceding some nonuniversality or choosing from a range of allowed parameter values are choices the constructor must address. One might argue whether this genuinely is “nonempirical”. We choose not to spend effort on that question and use the conceptual framework and vocabulary consistent with XC DFA development.

2.4.1. One-Point Functionals. Begin with an observation from experience. One-point $T_s[n]$ approximations in principle can depend upon $n(\mathbf{r})$ and all of its spatial derivatives, $\nabla n(\mathbf{r})$, $\nabla^2 n(\mathbf{r})$, ... In practice, the Euler equation that follows even from $\nabla^2 n(\mathbf{r})$ dependence is sufficiently complicated (fourth-order spatial derivatives) that there has been comparatively little consideration of a T_s approximation with higher than $\nabla^2 n$ dependence. Third-order spatial derivatives have been used in machine-learnt KEDFs,^{83,150,152,153} as we discuss in Section 2.5. While the sixth-order gradient expansion has a fourth-order spatial derivative contribution,¹³⁹ the expansion is divergent.^{136,139} Thus it is useless as a KEDF in itself. So far as we are aware, there have been no other candidate KEDFs with fourth-derivative dependence.

Recall from Section 1 that the implicit assumption in the one-point approach is that such functionals can probe nonlocal contributions adequately by selective exploitation of features of the gradient expansion. Because of the just-mentioned divergence, efforts began very early in DFT to formulate useful gradient contributions. Motivated by TF λ W eq 8 and a suggestion by Tomashima and Yonei,¹¹⁶ Plumer and Stott²⁷³ generalized the λ parameter to a density gradient dependence. A variation on the idea was put forth by Mazin in 1988²⁷⁴ but in an obscure journal and, hence, was not known in the OFDFT community until posted recently to arXiv. The idea reappeared in ref 69, where the specific form used is attributed to Pearson’s 1982 Ph.D. thesis. March and Santamaría²⁷⁵ presented yet a different type of gradient dependence, based on reasoning from the KS 1-rdm and recognition that exact exchange in KSDFT has the same form as HF.

Neither of those two schemes seems to have been pursued further. Instead, the strategy followed was adapted from density-gradient-dependent XC DFAs.^{57,276–278} Analogously with eq 31, generalized gradient approximation (GGA) KEDFs are constructed in the form

$$T_s^{GGA}[n] = c_{TF} \int d\mathbf{r} n^{5/3}(\mathbf{r}) F_t(s(\mathbf{r})) \quad (48)$$

$F_t(s)$, a function of the dimensionless reduced density gradient s introduced at eq 32, is the “enhancement factor” with respect to TF. It is dimensionless by construction so that the proper KE uniform scaling behavior (mentioned already²⁵⁴) provided by the Thomas–Fermi contribution is preserved. From eq 6, it

follows that the von Weizsäcker KEDF written as a GGA has the enhancement factor $(5/3)s^2$ and the Pauli term for a GGA KEDF T_θ^{GGA} has

$$F_\theta(s) = F_t(s) - \frac{5}{3}s^2 \quad (49)$$

as its enhancement factor. The GGA Pauli KE thus is

$$T_\theta[n] := \int d\mathbf{r} \tau_\theta[n](\mathbf{r}), \quad \tau_\theta = \tau_s - \frac{1}{8} \frac{|\nabla n|^2}{n} \quad (50)$$

The essence of GGA design strategy is to recover KE gradient expansion behavior for small s but modify it elsewhere so as to avoid the aforementioned^{31,136,139} divergent behavior. (Remark: The expressions for the fourth- and sixth-order gradient expansion terms are complicated and not of particular use here. See ref 195.) Thus, the small- s behavior of F_θ should have the form

$$F_\theta = 1 - c_2 s^2 + O(s^4) \quad (51)$$

In the gradient expansion itself, the quadratic coefficient is $c_2 = 40/27 \approx 1.482$.³¹

One way to develop GGA KEDFs is via rational construction, e.g. Padé approximants calibrated against such gradient expansion terms.⁶¹ Another is by extensive fitting⁷² to atomic data. Yet another is to exploit the “conjointness conjecture”⁵⁸ to rationalize use of the enhancement factor from a GGA exchange functional, E_x^{GGA} , in a GGA KEDF. That allows more than three decades of work on GGA XC DFAs to be exploited. While known not to be correct (see, for example, ref 268 or consider the fact that a GGA exchange DFA cannot satisfy all known constraints on it), the conjecture is instructive and, in fact, it fueled development of the first few GGA KEDFs.^{58,67,69,71}

However, none of those conjointness KEDFs would produce binding in test molecules even from post-SCF calculation with the correct conventional KSDFT densities⁶³ as input. (Here, “correct” is with respect, of course, to the XC DFA used.) The first GGA KEDFs that actually reproduced molecular binding from post-SCF calculation with KSDFT densities were the “modified conjoint” PBE n ($n = 2, 3, 4$) and exp4 functionals of Karasiev et al.^{63,73} The modification with respect to strict conjointness consisted of enforcement of the Pauli term positivity constraint, eq 35. Those modified conjoint KEDFs were not, however, purely nonempirical but were parametrized on very small training sets (a few molecules).

Recently therefore, three strategies have characterized efforts to construct GGA KEDFs: (1) satisfy as many known exact constraints as possible, the approach used to contrive the VT84F,⁸⁵ LKT,⁸⁶ and RATIONAL²⁴⁰ KEDFs; (2) devise an enhancement factor that yields as closely as possible the HEG linear response, the method used to obtain the PGint functional,²⁷⁹ (3) reproduce known semiclassical expansions for neutral atoms, leading to the APBEK and revAPBEK functionals.⁵⁹ All three involve, to varying extents, matching of the gradient expansion for small s . In the case of revAPBEK there also is an empirically adjusted parameter.

The APBEK, revAPBEK,⁵⁹ APBEKint, and revAPBEKint⁶⁰ KEDFs are worth examining a bit more. The PBE XC enhancement factor²²⁵ can be written compactly as

$$F^{pbe}(s) = 1 + \frac{\kappa \mu s^2}{\kappa + \mu s^2} \quad (52)$$

with κ and μ constants in the original XC GGA and in the simpler conjoint or conjoint-like KEDF cases. Thus, in APBEK and revAPBEK, $\mu = \mu^{\text{MGE2}} = 0.23899$ from the modified second-order gradient expansion, while $\kappa_{\text{APBEK}} = 0.804$ and $\kappa_{\text{revAPBEK}} = 1.245$. The modification in MGE2 is to match the asymptotic KE expansion (in Z) for neutral atoms. Though there is evidence of good performance on a small set of molecules, it is demonstrable that both APBEK and revAPBEK violate $v_\theta(\mathbf{r}) > 0$, eq 35, near the nucleus for Kato-cusped densities.²⁸⁰

A further refinement was to generalize the coefficient of the s^2 dependence (conventionally denoted μ) into an s dependence, leading to APBEKint and revAPBEKint.⁶⁰ The specific approach was to use an interpolation:

$$\mu^{\text{int}}(s) = \frac{3\mu^{\text{GE2}} + 5s^2\mu^{\text{MGE2}}}{3 + 5s^2} \quad (53)$$

with $\mu^{\text{GE2}} = 5/27$ from the KE second-order gradient expansion. It appears that both these KEDFs also violate v_θ positivity for Kato-cusped densities however.

Of these various GGAs, so far as we know, VT84F was the first nonempirical GGA KEDF to yield bound systems from solution of the OFDFT Euler equation.⁸⁵ Its Pauli enhancement factor reads

$$F_\theta^{\text{VT84F}}(s) = 1 - \frac{\mu s^2 e^{-\alpha s^2}}{1 + \mu s^2} + (1 - e^{-\alpha s^{m/2}})(s^{-n/2} - 1) \quad (54)$$

With $m = 8$ and $n = 4$, the functional has two parameters (μ and α) that are related by the gradient expansion at small s . For cusped densities, that relation can be satisfied while maintaining the $v_\theta > 0$ constraint, eq 36, only for a small range of parameter values. An Occam’s razor argument was used to choose among them.

The LKT⁸⁶ functional, in contrast, is specifically adapted to pseudodensities. It has the very simple Pauli enhancement factor

$$F_\theta^{\text{LKT}}(s) = 1/\cosh(as) \quad (55)$$

with the single parameter a set to 1.3 by enforcing $v_\theta > 0$ for low- Z atomic pseudodensities. Subsequently, the PG μ family⁸⁷

$$F_\theta^{\text{PG}\mu}(s) = \exp(-\mu s^2) \quad (56)$$

was parametrized to come close to reproducing F_θ^{LKT} by choosing $\mu = 0.75$.¹⁷² (Remark: μ here and in eq 57 is a parameter defining the specific Pauli enhancement factor and is not to be confused with the chemical potential.)

In Figure 5 we compare the Pauli enhancement factors of LKT, PG0.75, and VT84F. Because LKT and VT84F are adapted to specific near-nucleus behavior (zero density gradient or Kato cusp, respectively), they are nonuniversal. The effects of adaptation to nonuniversality are visible in the bump in the VT84F enhancement factor in the vicinity of $s = 1$. It provides v_θ positivity for Kato-cusped densities. Cuspless densities do not require the bump, as illustrated by LKT.

Numerical results illustrate the difference as well. Values from LKT and VT84F on simple metals and semiconductors are given in Table 2. Three metals, Li, Mg, and Al, each in simple cubic (sc), body-centered cubic (bcc), face-centered cubic (fcc), and hexagonal close-packed (hcp) structures, were treated. Nine zinc-blende structured III–V semiconductors also were treated, AlP, AlAs, AlSb, GaP, GaAs, GaSb, InP,

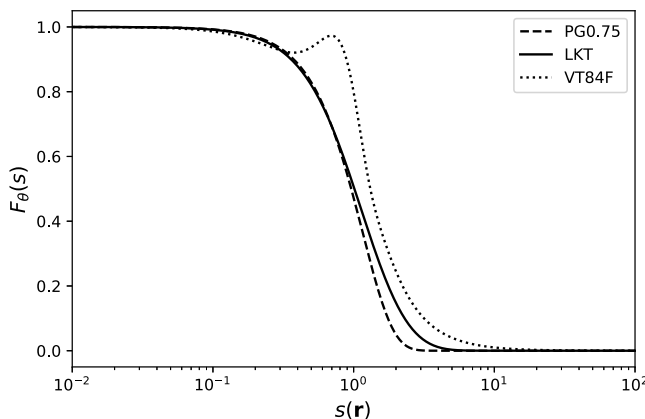


Figure 5. Comparison of the Pauli enhancement factors of LKT, VT84F, and PG0.75. By choice, PG0.75 is close to LKT up to $s \approx 1$.

Table 2. Effect of Nonuniversality Adaptation on KEDF Performance for Solid Metals and Semiconductors: MARE of Equilibrium Volumes, V_0 , Energies, E_0 , and Bulk Moduli, B_0 , as Percentages^a

KEDF	Metals			Semiconductors		
	V_0	E_0	B_0	V_0	E_0	B_0
LKT	4.0	0.2	7.7	2.1	2.8	4.3
VT84F	6.0	0.1	11.6	10.5	3.6	56.4

^aSee text for notation. Adapted from ref 86.

InAs, and InSb. Mean absolute relative errors (MAREs) referenced to the conventional KS values are reported in that table for equilibrium cell volume V_0 , total energy per cell E_0 , and bulk modulus B_0 . The Perdew–Zunger LDA XC DFA²⁸¹ was used. (For other technical details see ref 86 and its Supporting Information.)

All the OFDFT calculations were done with bulk-derived local pseudopotentials (see Subsection 3.1). LKT is adapted to such potentials. VT84F, in contrast, was built with constraint satisfaction for Kato-cusped densities. Table 2 shows substantial improvement when pseudodensities are adopted in all but one case, E_0 for the metals. Even that is an example of a big shift in a small number; the LKT MARE is 0.2%,

The PGint KEDF²⁷⁹ is interesting because of the unusual way in which it was parametrized. It takes the PG μ form, eq 56, and gives μ an s dependence, the same concept as in the “int” KEDFs just discussed. But instead of eq 53, PGint uses

$$\mu(s) = \mu_1 + (\mu_2 - \mu_1) \frac{\alpha s^2}{1 + \alpha s^2} \quad (57)$$

The parameters are $\mu_1 = 40/27$, $\mu_2 = 20/9$, and $\alpha = 10$. These values were chosen so that PGint recovers the second-order gradient expansion for $s \rightarrow 0$ and is close to PG μ with $\mu = 20/9$ for $s > 0.2$. The motivation for these design choices is to retain second-order gradient linear response in PGint. Though it gives some interesting results for atoms and jellium clusters, so far as we are aware, PGint has not been tested against thermochemical properties of molecules or solids nor does it seem to have been used to drive AIMD.

Two basic difficulties with any GGA KEDF motivate moving to inclusion of $\nabla^2 n$ dependence. First, for the case of a Kato-cusped density (the density from the physical ν_{ext} in the electronic structure problem), $\nu_\theta(\mathbf{r})$ is singular at the nuclear site, ²⁶⁴ $\lim_{r \rightarrow 0} \nu_\theta(\mathbf{r}) = a_\theta/r$ for any Pauli enhancement factor

that for small s goes as $F_\theta^{GGA} = 1 + a_\theta s^2$. Though the singularity is unphysical, the only GGA design choice is to make sure that the constant $a_\theta > 0$. That at least satisfies the $\nu_\theta > 0$ constraint. Many GGA KEDFs violate that constraint, for example APBEK and revAPBEK (recall above).

This dilemma motivated Karasiev et al.²⁶⁴ to consider reduced density derivative combinations of s and the dimensionless reduced density Laplacian defined as

$$q := \frac{\nabla^2 n}{4(3\pi^2)^{2/3} n^{5/3}} \quad (58)$$

(Remark: Beware of notational overload. The variable q should not be confused with the canonical position variable of classical phase space mentioned in Section 1 and with the Fourier space variable conjugate with $|\mathbf{r} - \mathbf{r}'|$ discussed later.) Then they rewrote the gradient expansion through fourth-order

$$T_\theta[n] = \int d\mathbf{r} [\tau_\theta^{(0)}[n](\mathbf{r}) + \tau_\theta^{(2)}[n](\mathbf{r}) + \tau_\theta^{(4)}[n](\mathbf{r}) + \dots] \quad (59)$$

with

$$\tau_\theta^{(2i)}[n](\mathbf{r}) = \tau_{TF}[n](\mathbf{r}) F_\theta^{(2i)}(s, q, \dots) \quad (60)$$

The corresponding enhancement factors are

$$\begin{aligned} F_\theta^{(0)} + F_\theta^{(2)} &= 1 + a_2 s^2 \\ F_\theta^{(4)} &= a_4 s^4 + b_2 q^2 + c_{21} s^2 q \end{aligned} \quad (61)$$

Instead of using the gradient expansion coefficients, however, they determined coefficients such that the ν_θ for a cusped density would be nonsingular at the origin. That led to, for example, the reduced density derivative combinations which are to be used as arguments of novel enhancement factors, such as $F_\theta(\kappa_4)$.

Reference 264 presented limited consideration of one KEDF constructed with the latter two variables but relied on determination of the parameters by fitting to conventional KSDFT results for two molecules and two atoms. The results were mixed, with some constraint violations but remarkably

$$\kappa_4 := s^4 + \frac{18}{13} q^2 - \frac{30}{13} s^2 q \quad (62)$$

$$\tilde{\kappa}_4 := \sqrt{s^4 + b_2 q^2}, \quad b_2 > 0 \quad (63)$$

$$\kappa_2 := s^2 + b_1 q \quad (64)$$

good performance in post-SCF calculations of T_s for 14 molecules, superior to that of the Perdew–Constantin meta-GGA KEDF²⁷⁰ that we discuss below. Post-SCF intramolecular forces (evaluated by finite differences) from the two were about equally good (or bad) in that some were rather accurate and some were quite off. Investigation of that ref 264 KEDF was not pursued. In retrospect, it may have been too complicated a form for initial exploration. Moreover, there was no compelling way to pick among possible reduced density derivatives nor to decide the form of the enhancement factor. What the exercise did show was the important role of $\nabla^2 n$ in getting rid of the unphysical ν_θ singularities of GGA KEDFs.

A few years later, Xia and Carter²⁸² pointed out the importance of $\nabla^2 n$ contributions in a different way. In our notation, they observe that the Pauli KE (be it approximate or exact) always can be written as

$$T_\theta[n] = c_{TF} \int d\mathbf{r} n^{5/3}(\mathbf{r}) G[n(\mathbf{r})] \quad (65)$$

(Remark: Beware that they use the form of the s variable *without* the constants, as we warned earlier.) For fcc Al, they calculated the KS KE density augmented with a bare Laplacian,

$$\tau(n; x) := \tau_s[n] + xq \quad (66)$$

with a constant mixing coefficient $-1 \leq x \leq 1$. (The calculations used PBE XC and a bulk-derived local pseudopotential, but neither choice affects the outcome.) Then they extracted $G[n; x]$. Plotted as a function of s for various values of x , complicated double-valuedness in G immediately is evident. Not only is $G(n; x) \neq G(s; x)$ but the double-valued behavior depends strongly on the q admixture coefficient x . The Xia–Carter results show that removal (or, at least, substantial suppression) of the double-valuedness would require the mixing to be a function of s , i.e. $x(s)$ at least. Since G is not solely a function of n and s , they concluded that the multivaluedness made the validity of GGA KEDFs questionable.

In a subsequent exchange of comments,^{252,283} Trickey, Karasiev, and Chakraborty argued that local imposition of the upper bound, eq 42, essentially cuts off the multivaluedness at least to the extent that a GGA KEDF is able to reproduce the most important features of $G[n]$ on the most important density range. In reply, Xia and Carter pointed out that local upper bound imposition does not remove what may be some critical regions of double-valuedness and that fundamentally such local imposition is not a rigorous constraint because it converts a sufficient condition into a necessary one. They did, however, agree that improved GGA KEDFs merited study.

The reduced density derivative and the $G[n]$ studies just discussed provide solid analytical and numerical motivation for considering Laplacian-dependent KEDFs. The investigation of such KEDFs has a long history because, among other things, of the gradient expansion. Recall from Section 1 that the simplest Laplacian-level functionals were explored in the mid-1980s^{124,284} to treat the known failure of GEA functionals to reproduce atomic shell structure in the KE density. A bare Laplacian, as in the Xia and Carter²⁸² study, has no effect on the kinetic energy potential, $v_s(\mathbf{r})$, delivered by a KEDF but changes the KE density (the gauge issue mentioned above). Empirically it was found that an appropriate admixture of a bare Laplacian term could describe atomic shell structure but at the cost of degrading the description of the near-nucleus kinetic energy density.^{124,284} Some recent studies also have included functionals with a bare multiplicative Laplacian term in an enhancement factor,^{270,285} as well as more complicated, constraint-based functions of s and q .

All of these can be subsumed in the Laplacian meta-GGA form^{268,270,285} already introduced, eq 60,

$$T_s^{mGGA-L}[n] = c_{TF} \int d\mathbf{r} n^{5/3}(\mathbf{r}) F_\tau^L(s^2(\mathbf{r}), q(\mathbf{r})) \quad (67)$$

The Perdew–Constantin (PC) meta-GGA has been among the more influential of these.²⁷⁰ It is designed to recover the fourth-order gradient expansion in the slowly varying case and T_{vW} in the rapidly varying one. Moreover, it is constrained to satisfy eq 34 locally. The form of the PC enhancement factor thus is another example of interpolation between limiting behaviors,

$$F_\tau^{PC} = \frac{5}{3}s^2 + \left(F_\tau^{mGE4} - \frac{5}{3}s^2 \right) f_{ab} \left(F_\tau^{mGE4} - \frac{5}{3}s^2 \right) \quad (68)$$

Here, F_τ^{mGE4} is a modification of the fourth-order gradient expansion

$$\begin{aligned} F_\tau^{mGE4} &:= \frac{F_\tau^{GE4}}{1 + \frac{\Delta_4}{\left(1 + \frac{5}{3}s^2\right)^{1/2}}} \\ F_\tau^{GE4} &= 1 + \frac{5}{27}s^2 + \frac{20}{9}q + \Delta_4 \\ \Delta_4 &:= \frac{8}{81}q^2 - \frac{1}{9}s^2q + \frac{8}{243}s^4 \end{aligned} \quad (69)$$

The interpolating function is

$$\begin{aligned} f_{ab}(z) &= 0, & z \leq 0 \\ f_{ab}(z) &= \left\{ \frac{1 + e^{a/(a-z)}}{e^{a/z} + e^{a/(a-z)}} \right\}^b, & 0 < z < a \\ f_{ab}(z) &= 1, & z \geq a \end{aligned} \quad (70)$$

with two parameters $0 < a \leq 1$ and $b > 0$. They were fitted to minimize the mean absolute relative error of the KS KE for a suite of neutral atoms, ions, neutral spherical jellium clusters, and liquid-drop model jellium spheres to yield $a = 0.5389$ and $b = 3$. Most of the testing was done on comparatively abstract systems, but the atomization kinetic energy, $T_s[\text{separated-atoms}] - T_s[\text{molecule}]$, was reported for post-SCF calculations on a set of 12 small molecules. Compared to the TF, GE2, and GE4, oddly the PC meta-GGA MAE was second best. TF was a bit better. However, TF got the sign of the difference wrong in four cases, whereas the PC meta-GGA had only two wrong. This illustrates a difficulty in testing. Seemingly innocent protocols can give misleading results.

A peculiarity in the PC meta-GGA was discovered and analyzed by Cancio and co-workers in the context of meta-GGA X functionals. In those functionals, it is typical to use an indicator function conventionally denoted as $\alpha := (\tau_s - \tau_{vW})/\tau_{TF} \equiv \tau_\theta/\tau_{TF}$. They found mishandling of the behavior for $\nabla^2 n < 0$ and associated difficulties in regions of strong electron localization. Negative $\nabla^2 n$ values are typical of covalent bond formation and for pseudodensities. To deal with these problems, they proposed a smoother enhancement factor^{268,286}

$$\begin{aligned} F_\tau^{CSK} &= \frac{5}{3}s^2 + 1 + z_{CSK} \{1 - \exp 1/z_{CSK} [1 - H(z_{CSK})]\} \\ z_{CSK} &:= \frac{20}{9}q - \frac{40}{27}s^2 \end{aligned} \quad (71)$$

Here $H(z)$ is the Heaviside unit step function and “CSK” denotes the authors (Cancio, Stewart, and Kuna). Results against the AE6 molecular data set²⁸⁷ (picked because of its diversity of bonding types) were illuminating. Post-SCF (density from PBE XC DFA in conventional KS) results are shown in Table 3. It compares atomization energy MAEs and KE MAREs. For scale, the MAE against experiment of the conventional KS calculation is 23.0 kcal/mol. Based on the atomization energies, VT84F is best. Thus, the table may seem to present a victory for GGA or a loss for meta-GGA KEDFs, depending on viewpoint. But the reality is more subtle. The

Table 3. Post-SCF Atomization Energy MAEs and KE MAREs against the AE6 Data Set Compared for Several mGGA KEDFs^a

KEDF	Atomization MAE (kcal/mol)	KE MARE
VT84F	182.1	0.2
mGGA-CSK	246.8	0.087
mGGA-PC	595.5	0.139
GE2	560.7	0.112
TF	671.1	0.162

^aAdapted from ref 286.

actual KE values are very different. For those, VT84F actually was the worst. The design priority for VT84F was to get good ionic forces in AIMD simulations while respecting as many constraints as attainable. The cost was a functional that generates KE values that are too large. Remember, however, that the table compares post-SCF values, not the results of solving eq 29 self-consistently with these KEDFs.

Other single-point functionals based on recovery of the fourth-order gradient expansion are found in refs 83 and 288–291. What is confirmed very convincingly by Golub and Manzhos⁸³ is that the fourth-order gradient expansion itself is not viable as a KEDF. Its KE density is notably worse than those from the second-order approximation in significant spatial regimes even for Si and Al. With coefficients adjusted for each system separately (hence, a nontransferable KEDF), agreement is much better for metals but not for covalently bonded systems. Those authors turned to machine-learning methods to seek further improvements; see Subsection 2.5.

Reference 291 also attempted to overcome limitations in the Perdew–Constantin KEDF.²⁷⁰ In our notation, those authors wrote

$$F_{\tau}^{\text{modGE4}} := 1 + \frac{5}{27}s^2 + \frac{20}{9}q - \gamma s^2q + \nu s^4 \quad (72)$$

The constants $\gamma = 0.2608$ and $\nu = 0.1826$ were determined by jellium surface energy and semiclassical neutral atom fitting. Unfortunately, it is difficult to set the recorded performance of this functional in the context of the rest of the literature. Though those authors did calculations on a test set of 21 molecules, they reported results on total KE and atomization KE differences only in terms of a “relative performance indicator” they devised. Ordinary MARE, MAE, etc. information was not provided nor were test results on periodic solids.

More recent investigation of Laplacian dependence has relied on devising an enhancement factor that reproduces as closely as possible the HEG linear response, for example to obtain the PGSL,⁸⁷ PGSLr,¹⁷² and GSE2 functionals.²⁷⁹ PGSL (“Pauli Gaussian supplemented with Laplacian”) has the Pauli term enhancement factor

$$F_{\theta}^{\text{PGSL}} := \exp\{-(40/27)s^2\} + \frac{1}{4}q^2 \quad (73)$$

The gradient expansion coefficient of q^2 would be 8/81. The value of 1/4 came from optimization against simple metals. PGSLr retains the good linear response of PGSL by modifying the Pauli term enhancement factor to be

$$F_{\theta}^{\text{PGSLr}} := F_{\theta}^{\text{PGSL}} - \lambda s^2q + \sigma s^4 \quad (74)$$

The empirical parameters, $\lambda = 0.4$ and $\sigma = 0.2$, were obtained from fitting to equilibrium volume, V_0 , total energy, E_0 , bulk

modulus, B_0 , and density errors for a set of 9 metals and 9 semiconductors. The authors noted that optimum values would be $\lambda^2 \approx \sigma$ but that numerical instability attendant with that choice prevented its adoption.

It is already clear that the best recent meta-GGA functionals provide some significant improvements relative to earlier GGA KEDFs but that the improvements are not systematic compared to the best current GGAs. Equally interestingly, however, the best of these GGAs and meta-GGAs are competitive with two-point functionals for calculations on bulk solids,^{86,172} on atomic systems,⁶⁰ and in subsystem DFT simulations.^{60,285,292–296} There is however a problem with Laplacians. High-order spatial derivatives in the Euler equation can cause numerical stability issues. We return to that in Section 3.

Table 4 gives a brief comparison (adapted from ref 172) of GGA and meta-GGA KEDF performance on a slightly larger

Table 4. KEDF Performance on Solid Metals and Semiconductors: MARE of Equilibrium Volumes, V_0 , Energies, E_0 , and Bulk Moduli, B_0 , as Percentages^a

KEDF	Metals			Semiconductors		
	V_0	E_0	B_0	V_0	E_0	B_0
SM (2-pt)	3.82	0.19	4.68	10.6	0.76	42.9
HC (2-pt)	5.84	0.43	11.7	1.98	0.51	11.67
LKT (GGA)	3.61	0.18	23.9	1.33	2.82	13.5
PG0.75 (GGA)	3.72	0.16	25.7	1.16	2.57	9.58
PGSLr (mGGA)	3.68	0.23	12.3	1.11	1.48	7.14

^aSee text for notation about KEDFs. Adapted from Ref 172.

set of simple metals and semiconductors than was used in the studies just discussed. Four metals, Li, Mg, Al, and Si, each in sc, bcc, fcc, and hcp structures were included. (Elemental Si is metallic in those structures.) Ten semiconductors, cubic diamond Si plus the nine III-Vs in the preceding data set, were used. For stringency, the table compares the performance of one- and two-point KEDFs. Details of the latter class are in the next subsection. Here note only that the two-pointers are the very early Smargiassi–Madden (SM)⁴⁶ and more recent Huang–Carter (HC).⁷⁷ Observe that HC is parametrized to semiconductors. Again, the OFDFT calculations were done with bulk-derived local pseudopotentials (see Subsection 3.1) but in this case with the PBE GGA XC DFA.²²⁵ Table 4 omits results from the WGC two-point functional²⁹⁷ because it was parametrized on metals and is known to perform poorly on semiconductors. Results from TF λ vW are omitted because it is non- N -representable for the values $\lambda < 1$ ordinarily used.

The table shows, somewhat surprisingly perhaps, that the best current GGA KEDFs, LKT and its closely related PG0.75, are rather good at yielding E_0 and V_0 in comparison with the more complicated two-point KEDFs. We do not have an explanation for the seemingly large increase in B_0 MARE for those two from the data set used to generate Table 2 to the data set used in Table 4. The independently calculated B_0 MAREs for LKT reported in refs 86 and 240 are much smaller and agree closely. The data set used in ref 172 differs from that used in the latter two references in having the Si systems. The calculations differ in the XC DFA. But all three studies find nearly the same results for LKT V_0 and E_0 MAREs, so it seems implausible that either of those differences (data set or XC DFA) would cause such a discrepancy in the B_0 MAREs. There

may be influences from different fitting procedures (energy vs volume), but we simply do not know.

This situation illustrates, however, a long-standing challenge for KEDFs. Bulk moduli often are a problem. One sees that here in the comparative performance of the PGSLr meta-GGA. Added Laplacian dependence (q dependence) and parametrization to a substantial data set do not make PGSLr any better than LKT for V_0 in either metals or semiconductors nor for metal E_0 , but it does improve for semiconductor E_0 . The big improvement achieved by PGSLr, at least on this small data set, is for B_0 . The step to a meta-GGA makes a major difference, roughly a factor of 2 MARE reduction.

Figure 6 gives a graphical view of the performance of several of the GGA and meta-GGA functionals treated in this section.

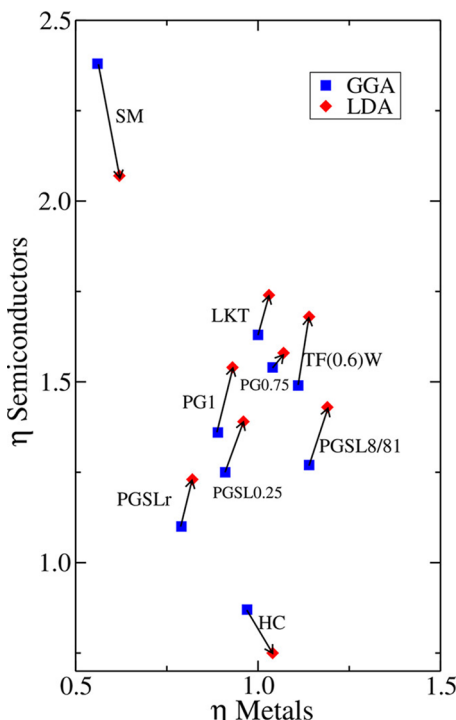


Figure 6. Global performance indicator, η , defined as a weighted average of deviations against benchmark values for metallic systems and semiconductor systems computed with LDA and PBE XC functionals and their associated local pseudopotentials (BLPS; see Section 3.1 for details on the pseudopotentials used in OFDFT simulations). Arrows connect results for LDA and PBE XC functionals for the same KEDF. Reprinted with permission from ref 172. Copyright (2011) American Chemical Society.

The so-called global performance indicator η was defined in ref 172 as a weighted average of MAREs for five properties, V_0 ; E_0 ; B_0 ; density error, $D_0 := \int d\mathbf{r} |n^{KS}(\mathbf{r}) - n^{OFDFT}(\mathbf{r})|$; and the error K_0 in the non-SCF non-interacting KE per cell. The weights were set on the basis of arguments for the relative importance of the five quantities. The indicator η was evaluated for both LDA and GGA (PBE) XC functionals and their associated pseudopotentials. A notable feature is that the one-point functionals do as well on metals as the HC two-point functional. Of course, HC was developed with semiconductors in mind, not metals, another example of the benefits and limitations of nonuniversality. An important observation also is that parametrized functionals, such as PGSLr, improve performance compared to the nonempirical LKT, hinting

that improvement of nonempirical one-point KEDFs can, and should be achieved.

We note that at least in the simple meta-GGA KEDFs contrived thus far, Laplacian dependence cannot generate density oscillations from the OFDFT Euler equation, even though such KEDFs give reasonable calculated energies and, often, quite good forces (energy gradients).

As an aside about GGA KEDFs, we remark that, although not shown here, among other somewhat recent ones, the LC94⁶⁷ and revAPBEK⁶⁰ functionals have been reasonably successful in the context of subsystem DFT simulations.^{180,181,190} Recall, however, the comments above about the lack of v_0 positivity in revAPBEK. Another recent development is the WPBEK KEDF,²⁹⁸ a different example of a modified conjoint GGA. It violates positivity constraints dramatically. That behavior led the authors to an interesting kind of dual representation, with a proper F_θ constructed so as to give the same non-interacting KE as the improper F_θ^{WPBEK} . We return to this idea in Subsection 2.4.3.

2.4.2. Two-Point Functionals. In Section 1 we remarked that interest in the nonlocality provided by two-point functionals originated in the first Hohenberg–Kohn DFT paper.³ There is a subtlety about that. The HK paper showed that effective approximate treatment of atomic density oscillations involves inclusion in the KE of excitations from occupied to virtual (above Fermi level) orbitals.⁴⁰ Because those orbitals are themselves density functionals (albeit implicit), the derived functional is still a density functional.

The HK approach did not catch on until 1985 with the first application of the scheme in materials physics⁴³ (jellium surfaces and atoms). That paved the way to several subsequent generations of two-point functionals, such as those of Wang–Teter,⁴⁴ Smargiassi–Madden⁴⁶ (already mentioned), Perrot,⁴⁷ Wang–Govind–Carter,⁴⁵ Huang–Carter⁷⁷ (already mentioned), and, recently, Mi–Genova–Pavanello⁸¹ and its density-dependent kernel version, LMGP,⁸⁸ as well as others.^{49,80,90}

Two-point KEDFs generally are expected to be more accurate than one-point KEDFs simply because the two-pointers probe and encode information about a larger spatial region than is possible with one-point functionals. Recall the discussion in Section 1 and Figure 3, but keep in mind the recent comparative success of one-pointers just surveyed.

Various of the earlier practical two-point KEDFs used somewhat near-sighted kernels⁴⁴ with Coulomb-like corrections for semiconductors.^{77,81,88} (Several^{45,77,80} also depended on parametrization beyond the level of isolated atoms and, hence, are semiempirical.) In that context it is informative to consider $T_s[n]$ in eq 23 from the perspective of probing an infinitesimal neighborhood of a sample point \mathbf{r} . $T_s[n]$ is derived explicitly as a one-electron operator acting on the KS orbitals. By virtue of orthogonality, those orbitals interfere, which gives rise to atomic shell structure. Thus, a proper KEDF must incorporate information about those shell–structure effects. That information is nonlocal because the nodal structure of low-lying orbitals forces the nodal structure of higher ones. This is one rationalization of the oft-stated claim that two-point functionals are essential for incorporation of shell structure.²⁹⁹ Note, however, that not all nonlocal KEDF formulations will yield good shell structure descriptions.³⁰⁰ (Remark: Even if one suspects that a one-point functional with sufficiently high-order spatial derivative contributions could give shell structure, the practical problem of numerical

instabilities so far seems formidable.) The Friedel oscillation argument (remarks just above and in Section 1) also is rooted in interference with states lying above the Fermi level.^{40,115}

This line of reasoning motivates two-point KEDFs developed from consideration of the perturbed electron gas.^{115,145,301} Suppose it to have nonzero average density n_{avg} . Then the weakly varying density is

$$\begin{aligned} n(\mathbf{r}) &= n_{avg} + \Delta(\mathbf{r}) \\ |\Delta(\mathbf{r})|/n_{avg} &\ll 1, \quad \forall \mathbf{r} \\ \int d\mathbf{r} \Delta(\mathbf{r}) &= 0 \end{aligned} \quad (75)$$

and to second order

$$\begin{aligned} T_s[n_{avg} + \Delta] &\approx T_{TF}[n_{avg}] + \frac{1}{2} \int d\mathbf{r}_1 d\mathbf{r}_2 \frac{\delta^2 T_s}{\delta n(\mathbf{r}_1) \delta n(\mathbf{r}_2)} \bigg|_{n_{avg}} \Delta(\mathbf{r}_1) \Delta(\mathbf{r}_2) \end{aligned} \quad (76)$$

The kernel of this integral, $\frac{\delta^2 T_s}{\delta n(\mathbf{r}_1) \delta n(\mathbf{r}_2)} \bigg|_{n_{avg}}$, has a simple, explicit dependence on the average density, n_{avg} and on spatial points \mathbf{r}_1 and \mathbf{r}_2 , thereby displaying the two-point structure. Recall the general expression for two-point functionals, eqs 13 and 14. Note, however, a significant difference. In general the kernel $\omega[n](\mathbf{r}, \mathbf{r}')$ has a more complicated and explicit density dependence than the simple n_{avg} in eq 76.

After eq 13 we noted that such nonlocal functionals are exact in principle. Here we provide an alternative route to realize this. It starts from recognition of the strict relationships among the derivatives of a functional and the functional itself. Included is the virial relation of eq 45.^{254,302} It is more instructive to consider functional integration relations.^{199,303,304} With them, the functional can be recovered from its first functional derivative, and the first functional derivative from the second. For a simple line integral with $n_t(\mathbf{r}) := f(t)n(\mathbf{r})$, T_s is

$$T_s[n] = \int d\mathbf{r} T_s[n(\mathbf{r})] = \int d\mathbf{r} \int_C dt \frac{\delta T_s[n_t]}{\delta n_t(\mathbf{r})} \frac{\delta n_t(\mathbf{r})}{\delta t} \quad (77)$$

where C is a smooth curve in \mathbb{R} parametrized by $f(t)$. For the simple case $n_t(\mathbf{r}) = tn(\mathbf{r})$ with $0 \leq t \leq 1$, we have^{81,304}

$$T_s[n] = \int d\mathbf{r} \int_0^1 dt \frac{\delta T_s[n_t]}{\delta n_t(\mathbf{r})} n(\mathbf{r}) \quad (78)$$

(This relationship has been exploited to compute KEDF energy corrections due to a nonlocal contribution to the potential.⁴⁸)

The procedure can be iterated to generate $\delta T_s/\delta n_t$ for insertion on the right-hand side of eq 78 to yield

$$T_s[n] = \int d\mathbf{r} \int d\mathbf{r}' \int_0^1 dt' \int_0^1 dt \frac{\delta^2 T_s}{\delta n_t(\mathbf{r}) \delta n_{t'}(\mathbf{r}')} n(\mathbf{r}) \quad (79)$$

and therefore the kernel in eq 14 can be expressed exactly as

$$\begin{aligned} \omega[n](\mathbf{r}, \mathbf{r}') &= n^{1-\alpha}(\mathbf{r}) n^{1-\beta}(\mathbf{r}') \int_0^1 dt' \times \\ &\quad \int_0^1 dt \frac{\delta^2(T_s - T_{TF} - T_{vW})}{\delta n_{t'}(\mathbf{r}') \delta n_t(\mathbf{r})} \end{aligned} \quad (80)$$

Practical utilization of this result thus involves finding a suitable approximation to the second functional derivative of T_s .

Hohenberg and Kohn³ established the connection between the KEDF kernel and the linear response function of the HEG. Defined as

$$\delta n(\mathbf{r}) = \int d\mathbf{r}' \chi_s(\mathbf{r}, \mathbf{r}') \delta v_{KS}(\mathbf{r}') \quad (81)$$

the connection of that response function with the KEDF usually is made by considering an additional functional derivative of the KS Euler equation (eq 29). Commonly it is written as¹¹⁵

$$\frac{\delta^2 T_s}{\delta n(\mathbf{r}) \delta n(\mathbf{r}')} = -\chi_s^{-1}(\mathbf{r}, \mathbf{r}') \quad (82)$$

The relationship was exploited and justified in several studies^{44,47,305} and also discussed in ref 115.

But care is needed because the KS Euler equation is not a general density functional. It is a ground-state expression. By virtue of the first Hohenberg–Kohn theorem, eq 29 establishes a one-to-one relationship between an external potential and its ground-state density; hence, $v_{ext}[n_0] \leftrightarrow n_0[v_{ext}]$. Thus, eq 29 should be written

$$\frac{\delta T_s[n]}{\delta n(\mathbf{r})} \bigg|_{n_0} + v_{KS}([n_0]; \mathbf{r}) = \mu[n_0] \quad (83)$$

Assuming ground-state v -representability and suitable restriction of differentiation to ground-state densities, the second functional derivative of T_s at n_0 is

$$\frac{\delta^2 T_s[n]}{\delta n(\mathbf{r}) \delta n(\mathbf{r}')} \bigg|_{n_0} = \frac{\delta \mu[n]}{\delta n(\mathbf{r}')} \bigg|_{n_0} - \frac{\delta v_{KS}(\mathbf{r})}{\delta n(\mathbf{r}')} \bigg|_{n_0} \quad (84)$$

Note the lack of explicit coordinate dependence in the $\mu[n]$. For each n_0 , the corresponding $\mu[n_0]$ is a spatial constant. This is crucial. The usual assumption (often unstated) is that μ is a constant from the original Lagrangian; therefore, its functional derivative is zero, and eq 82 follows. In general, however, different ground-state densities (and associated v_{ext}) at the same N_e have different μ . Consider, for example, the first ionization potential for a neutral atom and for the isoelectronic cation, e.g. Li and Be⁺. Thus, the usual claim about $\delta \mu/\delta n$ must be proved. So far as we are aware, the following argument is original.

The essential point is that because of Hohenberg–Kohn bijectivity, the first functional derivative of v_{KS} is symmetric in spatial arguments³⁰⁶

$$\frac{\delta v_{KS}(\mathbf{r})}{\delta n(\mathbf{r}')} = \frac{\delta v_{KS}(\mathbf{r}')}{\delta n(\mathbf{r})} \quad (85)$$

The same symmetry does not hold for the first functional derivative of $\mu[n_0]$. This follows from proof by contradiction. At fixed N_e , the presence of the proper $\mu[n_0]$ for each n_0 guarantees that

$$\int d\mathbf{r} \delta n(\mathbf{r}) = 0 \quad (86)$$

Now multiply [eq 84](#) by $\delta n(\mathbf{r})$ and integrate over the system volume:

$$\begin{aligned} & \int d\mathbf{r} \delta n(\mathbf{r}) \frac{\delta^2 T_s[n]}{\delta n(\mathbf{r}) \delta n(\mathbf{r}')} \Big|_{n_0} \\ &= \int d\mathbf{r} \delta n(\mathbf{r}) \frac{\delta \mu[n]}{\delta n(\mathbf{r}')} \Big|_{n_0} - \int d\mathbf{r} \delta n(\mathbf{r}) \frac{\delta v_{ks}(\mathbf{r})}{\delta n(\mathbf{r}')} \Big|_{n_0} \end{aligned} \quad (87)$$

Invoke the presumed symmetry of \mathbf{r} and \mathbf{r}' in μ and the actual symmetry in v_{ks} to get the second version of [eq 87](#):

$$\begin{aligned} & \int d\mathbf{r} \delta n(\mathbf{r}) \frac{\delta^2 T_s[n]}{\delta n(\mathbf{r}) \delta n(\mathbf{r}')} \Big|_{n_0} \\ &= \int d\mathbf{r} \delta n(\mathbf{r}) \frac{\delta \mu[n]}{\delta n(\mathbf{r})} \Big|_{n_0} - \int d\mathbf{r} \delta n(\mathbf{r}) \frac{\delta v_{ks}(\mathbf{r}')}{\delta n(\mathbf{r})} \Big|_{n_0} \end{aligned} \quad (88)$$

The difference is

$$\begin{aligned} 0 &= \int d\mathbf{r} \delta n(\mathbf{r}) \frac{\delta \mu[n]}{\delta n(\mathbf{r}') \Big|_{n_0}} - \int d\mathbf{r} \delta n(\mathbf{r}) \frac{\delta \mu[n]}{\delta n(\mathbf{r}) \Big|_{n_0}} \\ &= \frac{\delta \mu[n]}{\delta n(\mathbf{r}') \Big|_{n_0}} \int d\mathbf{r} \delta n(\mathbf{r}) - \int d\mathbf{r} \delta n(\mathbf{r}) \frac{\delta \mu[n]}{\delta n(\mathbf{r}) \Big|_{n_0}} \\ &= 0 - \int d\mathbf{r} \delta n(\mathbf{r}) \frac{\delta \mu[n]}{\delta n(\mathbf{r}) \Big|_{n_0}} \end{aligned} \quad (89)$$

Since this must hold at n_0 for arbitrary δn that integrates to zero, the conclusion is that $\frac{\delta \mu[n]}{\delta n(\mathbf{r}) \Big|_{n_0}} = 0$.

It follows therefore that

$$\frac{\delta^2 T_s[n]}{\delta n(\mathbf{r}) \delta n(\mathbf{r}') \Big|_{n_0}} = -\frac{\delta v_{ks}(\mathbf{r}')}{\delta n(\mathbf{r}') \Big|_{n_0}} \quad (90)$$

and [eq 82](#) follows.

Approximations must be considered for use of [eq 82](#) because knowledge of the KS response function itself would require solution of the KS equations ([eq 20](#)), exactly what OFDFT is meant to avoid. Computation of the response function via density functional perturbation theory³⁰⁷ or directly through the sum-over-states expressions also would be necessary. Since the formulation is a linear response, however, the obvious reference system is the HEG. Then the Fourier transform of the second-functional derivative is the inverse of the Lindhard linear response function.¹⁴⁰ The HEG is an advantageous reference because its response function depends only on $|\mathbf{r} - \mathbf{r}'|$ and, hence, its Fourier transform depends only upon a single reciprocal space variable, q . (Remark: Once again, consistency with the conventional literature notation forces us into symbol overload. But just to be sure, this q is not the reduced density Laplacian of the preceding subsection.) This makes inversion of the response function a simple “one-over” calculation rather than a general operator inverse as shown in [eq 82](#). That may not always be defined.²³⁷

Common practice therefore is to constrain the two-point kernel ω in [eq 13](#) to recover the HEG linear response in [eq 82](#)

evaluated at the equilibrium density. The consensus physical reasoning is that imposing that limit is beneficial, as the electronic structure of solids often resembles that of free electrons. That kind of behavior should be captured by the Lindhard function. A major challenge, however, is capturing the inverse response *beyond* Lindhard. To our knowledge there have been only two attempts at that. One, by Della Sala et al., aimed at encoding band gap behavior in a modified Lindhard function. The approach traces to Rey and Savin.³⁰⁸ The result was one GGA KEDF³⁰⁹ and one nonlocal KEDF with a density-independent kernel.⁸⁰ The other is by Takahashi.^{310–312} It is based on direct employment of the inverted KS response functions computed by a post conventional KS-DFT calculation of the KS response function. Initial results provide accurate but nontransferable functionals. Further work along either or both of these lines would improve the state of the art.

In any event, some implications of constraints immediately are apparent.

- The entire two-point term must vanish for uniform density.
- To obey the conjectured Lieb upper bound, [eq 42](#), requires

$$\omega[n](\mathbf{r}, \mathbf{r}') \leq 0 \quad (91)$$

- To obey Pauli term positivity, [eq 35](#), requires

$$T_{TF}[n] + \int d\mathbf{r} d\mathbf{r}' n^\alpha(\mathbf{r}) \omega[n](\mathbf{r}, \mathbf{r}') n^\beta(\mathbf{r}') \geq 0 \quad (92)$$

That suggests the kernel $\omega[n](\mathbf{r}, \mathbf{r}')$ should tend to a Dirac delta multiplied by a weight that goes like $n^{5/3-\alpha-\beta}$ for one-electron densities (i.e., when the kinetic energy density is that for von Weizsäcker, $\tau_s(\mathbf{r}) = \tau_{vW}(\mathbf{r})$), such as in the hydrogen atom or in the tail region of isolated atoms.

- Kernels must be dependent explicitly upon $n(\mathbf{r})$ and not solely on its average. Observe the notation, [eq 14](#), repeated here:

$$\mathcal{K}[n(\mathbf{r}), n(\mathbf{r}'), \mathbf{r}, \mathbf{r}'] = n^\alpha(\mathbf{r}) \omega[n](\mathbf{r}, \mathbf{r}') n^\beta(\mathbf{r}') \quad (93)$$

It is provable that density-independent kernels $\omega(\mathbf{r}, \mathbf{r}')$ yield KEDFs that are variationally unstable³¹³ whenever they are of the form given by Wang and Teter⁴⁴ (which is the most commonly used form). See discussion below.

- Uniform density scaling relations must be satisfied, so $\alpha + \beta = 8/3$ for functionals with density-dependent kernels, ω , which should be functions of dimensionless variables such as reduced density derivatives, $k_F(\mathbf{r})|\mathbf{r} - \mathbf{r}'|$, etc. so that they are homogeneous of degree zero under scaling. The local Fermi wave vector is

$$k_F(\mathbf{r}) := [3\pi^2 n(\mathbf{r})]^{1/3} \quad (94)$$

In principle, another constraint is that the entire two-point integrand should be symmetric with respect to interchange of \mathbf{r} and \mathbf{r}' ³¹⁴ but there are counterexample violations among practical approximations, e.g. ref 77.

Note also that two-point KEDFs with density-independent kernels depend on a reference uniform density, n_0 , which is an ill-defined quantity for any unenclosed system, e.g., an atom or molecule.^{297,299}

A significant deficiency of the Wang–Teter-like two-point KEDFs with density-independent kernels is the finding by

Blanc and Cancès³¹³ that such KEDFs are variationally unstable except with respect to near-uniform densities. There are many such functionals (Smargiassi–Madden,⁴⁶ Perrot,⁴⁷ Wang–Govind–Carter⁹⁸,⁴⁵ Mi–Genova–Pavanello,⁸¹ simplified versions of CAT-like functionals^{315,316}) which have been applied successfully to close-to-uniform systems (e.g., simple metals). The problematic variational instability does not seem to occur frequently in practical calculations of bulk systems. However, a recent study by Witt et al.³¹⁷ reports instabilities especially from the Perrot functional. The application of WT to warm dense matter also uncovered instabilities.³¹⁸

The issue was identified by Witt et al. as stemming from the violation of condition eq 92 by the Perrot functional as well as (even though infrequently) by the other two-point, density-independent kernel functionals. One can impose strict satisfaction of the positivity condition by applying an exponential stabilization scheme as follows. Rewrite the Pauli term in eq 92 as

$$T_{TF}[n] + \int d\mathbf{r} d\mathbf{r}' n^\alpha(\mathbf{r}) \omega[n](\mathbf{r}, \mathbf{r}') n^\beta(\mathbf{r}') = T_{TF}[n](1 + x) \quad (95)$$

Here $x = \frac{\int d\mathbf{r} d\mathbf{r}' n^\alpha(\mathbf{r}) \omega[n](\mathbf{r}, \mathbf{r}') n^\beta(\mathbf{r}')}{T_{TF}[n]}$. Whenever $x < -1$, Pauli-term positivity is violated. An elegant way to avoid this problem is to replace the $(1 + x)$ term by e^x . This substitution ensures the positivity condition always is satisfied and, in fact, no instabilities were observed once it was implemented.³¹⁷

The reader might be concerned that two-point functionals would be computationally expensive because a double spatial integral (six dimensions) must be evaluated. While true that direct evaluation would be unhelpfully costly, use of the HEG inverse response function in the formulation of ω in eq 80 reduces the evaluation cost to quasi-linear, $O(N \ln N)$. This matter is discussed in Subsection 3.4, about related computational aspects. The quasi-linear-scaling also is maintained for density-dependent kernels implemented by aid of spline techniques.^{77,88}

The first formulations of a nonlocal KEDF with density-dependent kernels resulted in the so-called average density approximation (ADA) and weighted density approximation (WDA),^{43,143,144,255,299,319–323} and later the WGC99 (WGC, hereafter) functional.²⁹⁷ (Remark: There is a section in ref 44 devoted to extending the Wang–Teter functional to a density-dependent kernel by discretizing the original one, $w_{WT}(\mathbf{r}_1, \mathbf{r}_2)$, over the distance $|\mathbf{r}_1 - \mathbf{r}_2|$ into N segments, evaluating the local Fermi wavevector on each segment, and then using that in a Gaussian basis expansion (Gaussians centered on each segment) of the original kernel. Structurally the scheme is reminiscent of the Trotter decomposition.³²⁴ To our knowledge, that WT scheme never has been used again.) In all of those kernels, except those of refs 43 and 255, explicit density dependence is introduced through an effective Fermi wavevector, ξ ,

$$\omega[n](\mathbf{r}_1, \mathbf{r}_2) = \omega[\xi(\mathbf{r}_1, \mathbf{r}_2)](\mathbf{r}_1, \mathbf{r}_2) \quad (96)$$

with

$$\xi(\mathbf{r}_1, \mathbf{r}_2) = \left(\frac{k_F^\gamma(\mathbf{r}_1) + k_F^\gamma(\mathbf{r}_2)}{2} \right)^{1/\gamma} \quad (97)$$

where $\gamma > 0$.

Beware that WDA/ADA KEDFs such as the symmetrized CAT functional²⁹⁹ are written in a somewhat different form than what we have used as canonical, to wit,

$$\begin{aligned} T_s^{CAT}[n] &= T_{TF}[n] + T_{NL}^{CAT}[n] \\ T_{NL}^{CAT}[n] &:= \frac{8}{5} c_{TF} \int d\mathbf{r} n(\mathbf{r}) \tilde{n}^{2/3}(\mathbf{r}) - \frac{3}{5} T_{TF}[n] \\ \tilde{n}(\mathbf{r}) &:= \int d\mathbf{r}_2 n(\mathbf{r}_2) [2\xi(\mathbf{r}_1, \mathbf{r}_2)]^3 \Omega(2\xi(\mathbf{r}_1, \mathbf{r}_2) |\mathbf{r}_1 - \mathbf{r}_2|) \end{aligned} \quad (98)$$

with $\gamma = -1/2$ in eq 97 chosen as a result of numerical experimentation. Ω has the role of a kernel. Note in particular that the “non-local” term in fact has a subtractive local (scaled TF) contribution.

The remarkable advance of the CAT functional was that it gave rather good atomic radial densities from SCF solutions of its Euler equation. From the perspective of 35 years later, CAT posed two difficulties. First, for two of the nine closed-shell atoms treated and one of the group V atoms, CAT gave lower total energies than the reference conventional KS (LDA XC) calculation, a sign of non- N -representability. The other difficulty was the explicit use of double spatial integrals. However, the original formulation of the CAT functional⁴³ did not feature a symmetrical integrand in the definition of $\tilde{n}(\mathbf{r})$ because it had a one-point effective Fermi wavevector. As we discuss in Subsection 3.4, use of one-point effective Fermi wavevectors enables techniques to avoid the double integration. There are simplified versions of CAT-like functionals^{315,316} in which a Fermi wavevector corresponding to an average electron density, k_F^0 , is used in place of ξ of eq 97. Simulations of liquid metals were successful with this strategy; see Section 4.3.

The WGC paper²⁹⁷ unified the formulation of two-point KEDFs in a single structure with the canonical form we have used here. For bulk Al lattice constants and energies per atom as test cases (sc, bcc, fcc, and hcp lattices), they explored the powers α and β from $(\frac{5}{6} \pm 0)$ to $(\frac{5}{6} \pm \sqrt{5}/6)$. The best results with respect to conventional KS calculations with LDA XC were for $(\alpha, \beta) = (\frac{5}{6} + \sqrt{5}/6, \frac{5}{6} - \sqrt{5}/6)$, hence, breaking spatial inversion symmetry ($\mathbf{r}_1 \leftrightarrow \mathbf{r}_2$). For eq 97 they found γ values from 1.9 to 2.7 were best. From that calibration, they calculated Al vacancy formation energies with success. However, Al surface energies were not as accurate, unless the reference average density used was the bulk value. Technically, the WGC formulation provided a clear step forward, namely reduction of the computational scaling by avoiding the double integral through the use of a Taylor expansion of the kernel around the reference constant density. As already mentioned, the reliance of WGC and its predecessors^{44,46,47} on such a reference density limits transferability and applicability of such KEDFs.

Several advances in the past decade have resolved the issue. The state of the art two-point functional (at least for bulk systems) is the Huang–Carter (HC) KEDF⁷⁷ mentioned already. Instead of the local Fermi-vector average $\xi(\mathbf{r}_1, \mathbf{r}_2)$ used in the WGC functional, the HC KEDF has a fully density-dependent kernel that has no HEG uniform density reference. Written in terms of eq 14, it is

$$\mathcal{K}^{HC}[n(\mathbf{r}_1), n(\mathbf{r}_2), \mathbf{r}_1, \mathbf{r}_2] = n^{8/3-\beta}(\mathbf{r}_1) \omega^{HC}[n](\mathbf{r}_1, \mathbf{r}_2) n^\beta(\mathbf{r}_2) \quad (99)$$

with

$$\omega^{HC}[n(\mathbf{r}_1)](\mathbf{r}_1, \mathbf{r}_2) = \omega(k_F(\mathbf{r}_1)[1 + \lambda s^2(\mathbf{r}_1)]|\mathbf{r}_1 - \mathbf{r}_2|) \quad (100)$$

Use of a one-point Fermi wavevector is explicit. In the foregoing equation, the s variable is the reduced density gradient defined in eq 32 (but *without* the constants) and β and λ are parameters determined empirically to be $\beta = 0.7143$ and $\lambda = 0.01177$ for III–V semiconductors. They are averages of optimal values over the set for the equilibrium volume, cohesive energy, and bulk modulus relative to conventional KS-DFT results from calculations with the same local pseudopotentials (BLPS; see Subsection 3.1 for details on the pseudopotentials used in OFDFT simulations). The simplified two-point dependence in the HC KEDF is important computationally, as we discuss in Subsection 3.4. Above we remarked (in the discussion of constraints) that this kernel also is not symmetric in \mathbf{r}_1 and \mathbf{r}_2 . Despite that, the calculational success of the HC KEDF on systems for which it was developed (discussed below) suggests, at least, that the symmetry violation does not have severe consequences for those systems.

The explicit nonlocality in ω^{HC} via the $\lambda s^2(\mathbf{r}_1)|\mathbf{r}_1 - \mathbf{r}_2|$ dependence is important. Nonzero even when the density is small (e.g., $s \rightarrow \infty$ in the tail of an isolated system), the dependence is included to approach the proper limit of the second functional derivative of the KEDF for $|\mathbf{r}_1 - \mathbf{r}_2| \rightarrow \infty$ (equivalently, $q \rightarrow 0$ in Fourier space; again beware of notational overload). This follows from exact properties of the response function^{325,326} given the relationship in eq 82.

In the HEG limit that is the Lindhard function

$$\chi_{\text{Lind}}(\eta) = -\frac{k_F}{\pi^2} \left[\frac{1}{2} + \frac{1 - \eta^2}{4\eta} \ln\left(\frac{1 + \eta}{1 - \eta}\right) \right] \quad (101)$$

where $\eta = \frac{q}{2k_F}$ for spin-compensated systems. (In the HEG, $k_F = (3\pi^2 n_0)^{1/3}$ is a constant of course.) (Remark: For sake of clarity, the variable q in the definition of η is to be considered $q = |\mathbf{G}|$, the absolute value of the reciprocal space variable, \mathbf{G} . For reference, the allowed \mathbf{G} vectors satisfy the relation $e^{i2\pi\mathbf{r}\cdot\mathbf{G}} = 1$.) Imposition of eq 82 leads to a first-order differential equation for the ω^{HC} that defines the kernel as a function of η . That ω^{HC} then is used in eq 100 by appropriate change of the kernel variable and consideration of the Fourier transform with respect to the $|\mathbf{r}_1 - \mathbf{r}_2|$ scalar variable.

The HC functional provided a major step forward for OFDFT applied to materials science. For example, for the first time an OFDFT functional could reproduce, at least nearly quantitatively, conventional KS-DFT results for bulk semiconductors. See Figure 7.

From the figure, it is clear that OFDFT can approach quantitative treatment of such semiconducting systems. Of course, optimism generated by that good HC KEDF performance is modulated by the reality that the tested semiconductor systems were part of the training set used to parametrize the functional. Subsequent application of the HC KEDF to systems outside the training set showed that HC is nevertheless a quality KEDF.³²⁷

However, that good performance is restricted to bulk systems. That limitation motivated a modification of HC to extend its applicability to highly inhomogeneous systems.⁹¹ The modification addressed a numerically difficult aspect of the functional, namely the $s^2(\mathbf{r})$ dependence in the kernel eq

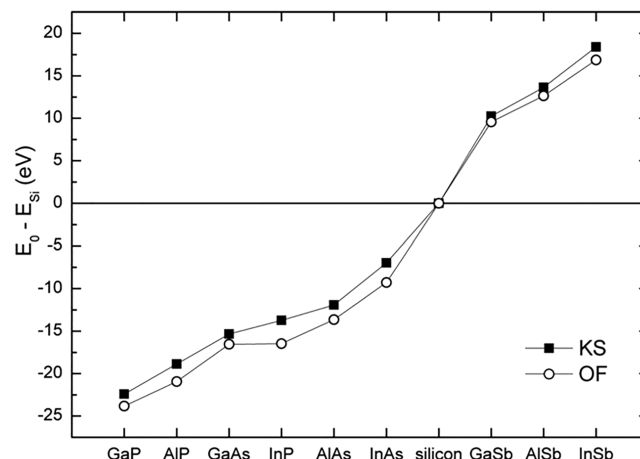


Figure 7. Energy differences (per primitive cell) between the diamond silicon and zinc blende phases of III–V semiconductors. OFDFT was carried out with the HC functional with parameters as in the text. Conventional KS-DFT (denoted as “KS”) calculations used the same local pseudopotentials for even-handed comparison. Reprinted with permission from ref 77. Copyright (2010) American Physical Society.

100. Simply replacing the $1 + \lambda s^2$ dependence with a PBE-like enhancement factor, $1 + \frac{as^2}{1+bs^2}$ with $a = 0.45$ and $b = 0.1$ (note that the dependence on λ disappears), delivered accurate surface energies and predicted surface morphologies for metals and selected semiconductors. The resulting KEDF is called revised HC (revHC). An important achievement was that revHC reproduced reasonably well the reconstructed Si(111) surface morphology predicted by conventional KS-DFT.

What about isolated systems, i.e., gas phase chemistry? The CAT family of functionals^{43,299,319–321} gave a clear hint that if nonlocality is taken into account, isolated systems also could be approached by OFDFT in close-to-quantitative accuracy similar to what HC and revHC deliver for bulk systems and surfaces.

To understand recent progress on that issue, it is useful to take a step back historically. Harking back to functional integration, e.g. eq 77, Chai and Weeks⁴⁸ went to the potential (in essence ν_θ) as the key. They developed a nonlocal functional that reduced properly to the TF and vW forms and had proper low- q or high- q linear response for weakly inhomogeneous systems. (Remark: We remark that q here is used in a way consistent with the literature. A more sensitive variable to use would be $\eta = \frac{q}{2k_F}$ because of its dependence on the local Fermi wavevector.) Both forms have the Pauli potential³²⁸

$$\nu_\theta^{CW2} = \frac{5}{3} c_{TF} n^{2/3}(\mathbf{r}) + \frac{10}{9\alpha} c_{TF} n^\beta(\mathbf{r}) \int d\mathbf{r}' f(|\mathbf{r} - \mathbf{r}'|; k_F(\mathbf{r}')) n^\alpha(\mathbf{r}') \quad (102)$$

with α and β constants related by $\beta = 2/3 - \alpha$. The value for the version that satisfies low- q constraints (“LQ”) is $\alpha = \frac{1}{2}$, while large- q constraint satisfaction (“HQ”) requires $\alpha = 2/3$. The Fourier transform of $f(|\mathbf{r} - \mathbf{r}'|; k_F(\mathbf{r}'))$ with respect to $|\mathbf{r} - \mathbf{r}'|$ for a fixed \mathbf{r}' is known explicitly, $f(\eta(\mathbf{r}')) = F_L(\eta(\mathbf{r}')) - 3\eta(\mathbf{r}')^2 - 1$ with $F_L(\eta)$ the inverse of the Lindhard response, eq 101.

Whether to use the LQ or HQ version would seem to be another example of design choice, but the advantage of hindsight, specifically the development of the N -representability criteria already discussed,²⁵⁹ enables a clear conclusion. For all but two of the 13 atoms they treated, self-consistent solution and use of the Herring path²⁴⁵ with the LQ version gave $E_{tot}^{LQ} < E_{tot}^{KS}$. In contrast $E_{tot}^{HQ} > E_{tot}^{KS}$ for all 13 atoms. At least on that set, the HQ functional imputed from the HQ kinetic potential is N -representable. The LQ one is not. Encouragingly, the MARE for the HQ functional on those 13 atomic total energies was only about twice that of the much more complicated CAT functional (discussed above).^{43,299,319–321}

The atomic densities at the nucleus $n(0)$ from HQ are much worse than from CAT however. Such differences led Chai and Weeks to discuss nonuniversality of KEDFs and KE potentials with respect to pseudopotentials (recall discussion of LKT above), apparently the first to do so.⁴⁸

For application to solids with near linear-scaling, $O(N \ln N)$ with N the grid count, the HQ weight function was expanded in a Taylor series around a reference density³²⁸ in a way similar to WGC and XWM two-point functionals.^{49,297} (We remark that such a Taylor expansion is not strictly needed because of the simple dependence of the Chai–Weeks kernel on the local Fermi wavevector. Thus, spline techniques, such as those discussed next for LMGP and HC, would be applicable to this functional. Those techniques were not in common use at the time the Chai–Weeks functionals were proposed.) For crystalline Al, both the lattice parameters and the energetic sequence of phases (fcc lowest, then bcc, sc, diamond) came out correctly with HQ. For Si, though the lattice parameters were reasonable, the order of crystalline phases was completely wrong, sc, fcc, bcc, dia (most to least bound), vs the proper order of dia, bcc, sc, fcc.

We return to the Chai–Weeks scheme below. Here, we focus on the next use of the functional integration approach, the recent development of the LMGP two-point functional with a density-dependent kernel.⁸⁸ Based on the Mi–Genova–Pavanello (MGP) KEDF,⁸¹ LMGP targets isolated systems by proposing a model second functional derivative of T_s . It is used in a manner similar to eq 79 (but limited to a single t integration) to yield a potential. The model second functional derivative is crafted so that it (1) reduces to the inverse Lindhard function for uniform densities; (2) is augmented by a term that imposes correct $1/q^2$ limiting behavior when $q \rightarrow 0$; (3) is symmetrized with respect to the interchange $n(\mathbf{r}) \leftrightarrow n(\mathbf{r}')$; and finally, (4) is fully density dependent, like HC and CAT. Full density dependence is computationally achieved in the LMGP family by a numerical scheme that is described in detail in Subsection 3.4.

The LMGP form for the second-functional derivative of the nonlocal part of the KEDF is

$$\frac{\delta^2 T_{NL}}{\delta n(\mathbf{r}) \delta n(\mathbf{r}')} = \frac{\pi^2}{(3\pi^2)^{1/3}} (n(\mathbf{r}))^{-1/6} G_{NL}[n(\mathbf{r}), n(\mathbf{r}')] (\mathbf{r} - \mathbf{r}') (n(\mathbf{r}'))^{-1/6} \quad (103)$$

Following eq 82, the Fourier transform of $G_{NL}[n(\mathbf{r}), n(\mathbf{r}')] (\mathbf{r} - \mathbf{r}')$ evaluated at the uniform density $n(\mathbf{r}) = n(\mathbf{r}') = n_0$ is related to the inverse Lindhard function:

$$G_{NL}[n_0](\eta) = -\frac{k_F}{\pi^2} \chi_{\text{Lind}}(\eta)^{-1} - 3\eta^2 - 1 \quad (104)$$

with η defined as previously with a constant $k_F = (3\pi^2 n_0)^{1/3}$. Fourier transformation of $G_{NL}[n_0](\eta)$ leads to $G_{NL}[n_0](\mathbf{r} - \mathbf{r}')$. This makes χ_{Lind} effectively a function of n_0 . The dependence of $G_{NL}[n_0](\mathbf{r} - \mathbf{r}')$ (shorthand notation G_{NL} below) on $n(\mathbf{r})$ and $n(\mathbf{r}')$ then is included approximately as follows

$$G_{LMGP}[n(\mathbf{r}), n(\mathbf{r}')] = G_{NL}[n_0 = n(\mathbf{r})] \quad (105)$$

The LMGP KEDFs include the correct $q \rightarrow 0$ limit by adding a term to eq 103 designed to mimic q^{-2} limiting behavior. Finally, an integration procedure brings the second derivative in eq 103 to the corresponding potential. The outcome is

$$\frac{\delta T_{NL}}{\delta n(\mathbf{r})} = v_{NL}(\mathbf{r}) = \int d\mathbf{r}' n(\mathbf{r}') \left[\int_0^1 dt \frac{\delta^2 T_{NL}}{\delta n(\mathbf{r}) \delta n_t(\mathbf{r}')} \right] \quad (106)$$

where $n_t(\mathbf{r}')$ is defined as eq 78. Equivalently, eq 79 gives the energy.

When computing the t -path integral, whether numerically or analytically, we note that because of eq 105, the LMGP kernel imposes $t = t'$ in the integral eq 106. To ameliorate this approximation, kernels of imposed symmetrization were considered. For example, arithmetic symmetrization leads to the LMGP functional

$$G_{LMGP}[n_0 t, n_0 t'] \rightarrow \frac{1}{2} [G_{NL}[n_0 t] + G_{NL}[n_0 t']] \quad (107)$$

That provides an improved t -path integration compared to LMGP because in obtaining the potential one needs to impose $t' = 1$. A geometric symmetrization also was considered: see the Supporting Information document of ref 81 for more details.

The kernel resulting from the integration then depends on n_0 and is used in a similar way to eq 100 to obtain the potential

$$v_{NL}(\mathbf{r}) = n^{-1/6}(\mathbf{r}) \mathcal{F}^{-1}[\Omega[n(\mathbf{r})](\eta(|\mathbf{G}|)) \mathcal{F}[n^{5/6}(\mathbf{r}')](\mathbf{G})](\mathbf{r}) \quad (108)$$

Here $\Omega[n(\mathbf{r})](\eta(|\mathbf{G}|)) = \int_0^1 dt \mathcal{F} \left[\frac{\delta^2 T_{NL}}{\delta n(\mathbf{r}) \delta n_t(\mathbf{r}')} \right]_{n(\mathbf{r})=n_0, t n(\mathbf{r}')=t n_0}$, \mathbf{G} is

the reciprocal space variable, as before, $\eta = \frac{|\mathbf{G}|}{2(3\pi^2 n_0)^{1/3}}$, and \mathcal{F} is the Fourier transform operator. It is important to remark that while the model second functional derivatives have been symmetrized with respect to the t -path integration variable, the underlying approximation $n(\mathbf{r}), n(\mathbf{r}') \rightarrow n_0$ remains. Thus, in eq 108 above, the potential and the density in the kernel are evaluated at point \mathbf{r} , effectively imposing $n_0 = n(\mathbf{r})$ in the definition of the kernel. Such an approximate algorithm is dictated by computational scaling, which is kept at $N \ln(N)$. It would be much steeper and computationally prohibitive for the kernel to be formulated as a function of both $n(\mathbf{r})$ and $n(\mathbf{r}')$.

The performance of LMGP is more than satisfactory for metal clusters and III–V semiconductor quantum dots; see the discussion in Section 4. As we discuss later in this section, LMGP also has been employed successfully in subsystem DFT simulations.⁸⁹

The development of HC and LMGP has motivated formulation of at least two further KEDFs, LDAK-X⁹⁰ and XWM.⁴⁹ “X” in LDAK-X gives the name of a parent functional from which the kernel is borrowed. For example, when X = MGP, the MGP kernel⁸¹ is used. For X = WT, the WT kernel is used. LDAK-X, in essence, exploits the same underlying

formalism as LMGP. However, it differs from LMGP in the definition of the energy. It avoids the potential-to-energy integration of eq 79 and instead defines an ad-hoc energy expression from which the corresponding potential is obtained by functional differentiation.⁹⁰ LDAK-MGP was shown to improve upon LMGP considerably for an array of clusters and other isolated systems.⁹⁰

The XWM KEDF⁴⁹ handles the Fermi wavevector in a way similar to that in symmetrized CAT and WGC (see text after eq 96). Also similarly to WGC, a computationally amenable algorithm arises from Taylor expansion of the kernel around a reference constant density, n_0 . As noted with other two-point functionals, that prescription makes XWM primarily suitable for bulk systems unless a recipe for n_0 is invoked (recall what is done with the WGC KEDF). A desirable alternative would be an algorithm or approximation that could evaluate WGC and XWM without explicit double integration or being tied to a reference density. However, as of now, such an algorithm has yet to be proposed.

A different approach has been pursued by the Della Sala group. They proposed two new types of nonlocal KEDFs with density-dependent kernels which are entirely defined in real space and do not rely solely on the Lindhard function: yGGA³²⁹ and uGE4m.²⁹¹ These KEDFs extend meta-GGA functionals with the inclusion of nonlocal ingredients. Their energy expressions are based on enhancement factors

$$T_s^{uGE4m} = c_{TF} \int d\mathbf{r} n^{5/3}(\mathbf{r}) F_s(s^2(\mathbf{r}), q(\mathbf{r}), y_\alpha(\mathbf{r})) \quad (109)$$

$$T_s^{yGGA} = c_{TF} \int d\mathbf{r} n^{5/3}(\mathbf{r}) F_s(s^2(\mathbf{r}), q(\mathbf{r}), u(\mathbf{r})) \quad (110)$$

Here q is the reduced density Laplacian introduced at eq 58 and

$$u(\mathbf{r}) \equiv v_H(\mathbf{r}) = \int d\mathbf{r}' \frac{n(\mathbf{r}')}{|\mathbf{r} - \mathbf{r}'|} \quad (111)$$

$$y_\alpha(\mathbf{r}) = \frac{3\pi^2\alpha^2}{4k_F(\mathbf{r})} \int d\mathbf{r}' \frac{n(\mathbf{r}')}{|\mathbf{r} - \mathbf{r}'|} e^{-\alpha k_F(\mathbf{r})|\mathbf{r} - \mathbf{r}'|} \quad (112)$$

The enhancement factors were developed to satisfy a number of conditions. For example, uGE4m recovers the semiclassical expansion of neutral atoms for $Z \rightarrow \infty$.²⁹¹ Instead, yGGA focuses on reproducing the Lindhard response for uniform densities in the low q region. To achieve this, several enhancement factors were proposed, such as (we use a short-hand notation for the functions involved with all of the following being functions of \mathbf{r} : s , q , u , y_α , η_w , ν_w , n , x_1 , and x_2),

$$F_s^{uGE4m} = +\frac{8}{81}q^2 - \left[\frac{1}{9} + \left(\nu - \frac{8}{243}f_1 \right) \right] s^2 q + f_2 \nu s^4 \quad (113)$$

$$F_s^{yGGA} = \frac{5}{3}s^3 + y_\alpha G(s^2, q) \quad (114)$$

For F_s^{uGE4m} the values are $\nu = 0.1826$, $f_1 = (1 + a \nu_u^{1/2})^{-1/2}$, $f_2 = (1 + \nu_u^{1/2} + b \nu_u)^{-1}$, and $\nu_u = \frac{1}{1 + \eta_u}$, where $\eta_u = \frac{u}{3(3/\pi)^{1/3} n^{1/3}}$.

The parameters defining f_1 and f_2 are $a = 450$ and $b = 240$. For F_s^{yGGA} , i.e., the *yuk4* functional from ref 329, the G function is

defined as $G(s^2, q) = T_{3,3}(x_1)T_2(x_2)$, where $x_1 = -\frac{40}{27}s^2$, $x_2 = \frac{40}{27}q$, and $T_{\lambda}(x) = \frac{4}{\lambda} \frac{e^{ix}}{e^{ix} + 1} + \frac{\lambda - 2}{\lambda}$.

Although the uGE4m KEDF is applicable solely to finite systems (the u function is undetermined for periodic systems due to its divergence, which is canceled by charge neutrality in those systems), in application to a variety of isolated systems (jellium clusters, atoms, and molecules) it delivered good results. The yGGA KEDFs were benchmarked against jellium clusters and their ground-state response to external perturbations as well as atoms (non-self-consistently) and showed promising results. An implementation of yGGA functionals for nonspherical systems is not yet available. The recent introduction of these functionals suggests the need and opportunity for further assessment of them.

Another interesting approach^{310–312} develops two strategies to obtain inverse KS response functions to be used to define KEDFs and their functional derivatives. The strategies differ in the way the response functions are represented. In the first case,^{310,311} the response functions are mapped on an ad-hoc 1D coordinate. The map applied to the electron density is given by $n^e(\varepsilon) = \int d\mathbf{r} n(\mathbf{r}) \delta(\varepsilon - v_{ext}(\mathbf{r}))$, which is valid only for isolated atoms and pure elemental crystalline solids. The KS response function therefore is transformed as $\chi_s(\mathbf{r}, \mathbf{r}') \rightarrow \chi_s(\varepsilon, \varepsilon')$, a form which can be handled more efficiently for inversion. In ref 312, the inversion is handled instead by a spectral representation of the response function. Once χ_s^{-1} is obtained, approximations for the non-interacting kinetic energy and its derivative are formulated as an expansion around a reference density, $n_0(\mathbf{r})$. Following eq 76

$$T_s[n] \simeq T_s[n_0] + \int d\mathbf{r} v_{T_s}(\mathbf{r}) \delta n(\mathbf{r}) + \frac{1}{2} \int d\mathbf{r} d\mathbf{r}' \delta n(\mathbf{r}) \chi_s^{-1}(\mathbf{r}, \mathbf{r}') \delta n(\mathbf{r}') \quad (115)$$

where $\delta n = n - n_0$. Even though the results in refs 310–312 seem encouraging, the approach needs further exploration and the underlying approximations need to be characterized and delineated more clearly before it can be considered viable for broad applicability.

Another interesting and potentially promising real-space route to construct KEDFs is to define a model 1-rdm that depends purely on the electron density. This was attempted by Chakraborty et al.^{260,261} Given a 1-rdm, $\gamma(\mathbf{r}, \mathbf{r}')$, the kinetic energy associated with it is

$$T_s[n] = -\frac{1}{2} \int d\mathbf{r} d\mathbf{r}' \delta(\mathbf{r} - \mathbf{r}') (\nabla_{\mathbf{r}}^2 \gamma[n; \mathbf{r}, \mathbf{r}']) \quad (116)$$

Chakraborty et al. proposed model 1-rdms of the form

$$\gamma[n; \mathbf{r}, \mathbf{r}'] = \sqrt{n(\mathbf{r})n(\mathbf{r}')} g(k_F|\mathbf{r} - \mathbf{r}'|) \quad (117)$$

This form is motivated by DFA development for the exchange energy, as well as having some resemblance to two-point functional ingredients already encountered. The effective wavevector, k_F , however, is a function to be defined. It can be either local or nonlocal. In principle eq 117 is exact, as it is derived directly from the formal definition of the exchange energy in terms of the exchange hole.¹⁰² Substitution of eq 117 in eq 116 gives

$$T_s[n] = \int d\mathbf{r} \frac{|\nabla n(\mathbf{r})|^2}{8n(\mathbf{r})} - \frac{3g''(0)}{2} \int d\mathbf{r} n(\mathbf{r}) k_F^2(\mathbf{r}) \quad (118)$$

where the double prime means second derivative.

Even though the function $g(x)$ should satisfy several known conditions (e.g., $g(0) = 1$, $g'(0) = 0$, $g''(0) < 0$, and $-1 < g(x) \leq 1$ for the case of $x \leq 0$), those are not sufficient to determine the function fully. In ref 260, the $g(x)$ for the HEG was chosen to ensure that the constructed KEDFs are exact for uniform densities, i.e.,

$$g_{HEG}(x) = 3 \left(\frac{\sin x - x \cos x}{x^3} \right) \quad (119)$$

The remaining important step to generate a KEDF in this approach is to construct the effective Fermi wavevector, k_F , that appears in the argument of g in eq 117. A straightforward option is LDA in eq 94. That choice simply yields TFvW. Another option is the symmetrized two-point form defined in eq 97. In principle there are others. What Chakraborty et al. did was to impose the diagonal idempotency condition

$$\int d\mathbf{r}' \gamma(\mathbf{r}, \mathbf{r}') \gamma(\mathbf{r}', \mathbf{r}) = n(\mathbf{r}) \quad (120)$$

That leads to nonlinear integral equations that correspond either to the weighted density approximation (WDA) in the one-point case (1WDA) or the two-point WDA (2WDA) for k_F .

Benchmarks for the kinetic energies of H-Ar atoms and some small molecules showed that those functionals do not deliver quantitative accuracy. However, both the 1WDA and 2WDA improve considerably on the LDA results (i.e., $k_F(\mathbf{r}) = (3\pi^2 n(\mathbf{r}))^{1/3}$). Table 1 of ref 260 shows that the 2WDA KEDF gives kinetic energies lower than the reference for atoms larger than Si, an indication of non- N -representability. Figure 8

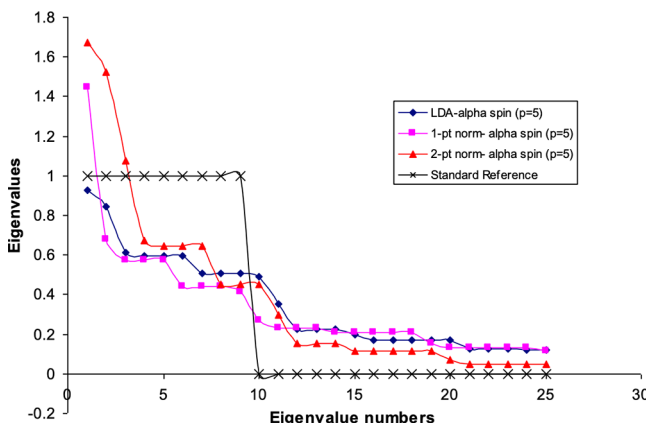


Figure 8. Natural orbital occupation numbers for the model 1-rdms for three choices of the effective Fermi wavevector to be used as the argument of the $g(x)$ function in eq 117: LDA, 1WDA (1 pt norm), and 2WDA (2 pt norm) for the argon atom. Calculations are spin polarized, and p corresponds to the value of the exponent γ (not to be confused with the 1-rdm which bears the same symbol) defined in eq 97, which is a parameter used in the symmetrization procedure of the model 1-rdms. Reprinted with permission from ref 260. Copyright (2017) Springer.

(taken from the paper) shows the eigenvalues of the model 1-rdms to have Pauli principle violations (i.e., eigenvalues larger than 1) for Ar, one of the atoms for which the KEDF gave a value below the exact T_s from conventional KSDFT. Interestingly, the figure also hints that the violations occur mostly for core orbitals. That suggests that non- N -represent-

ability might be more problematic for all-electron calculations than for pseudopotential-based OFDFT calculations.

Another interesting observation is the fact that, at least for Ar, the various approximations for the effective k_F function (LDA, 1WDA, 2WDA) lead to at most ensemble- N -representable densities with occupations smeared across several natural orbitals.

To recapitulate, though there is no two-point KEDF yet that works equally well for both isolated and bulk systems, substantial progress has been made. Density-dependent kernels are essential, both formally and practically. The double-integral computational burden can be avoided by a combination of splines and convolutions; see Subsection 3.4. At present, the HC and revHC functionals are perhaps still the best two-point functionals for bulk systems and surfaces. Encouragingly, as shown in Table 4 and Figure 6, there also are one-point KEDFs such as LKT⁸⁶ that exhibit comparable (or nearly so) accuracy. This suggests judicious use of both forms together as a possible way forward, as exemplified by the uGE4 and yGGA KEDFs we discussed. For isolated systems, the most advanced two-pointers are LDAK-MGP and LMGP. However, a better and more physical approach to the density dependence of the kernel is given by the CAT, WGC, and XWM functionals. The unresolved challenge to a full implementation of those functionals is to avoid double integrations or Taylor expansions.

2.4.3. Other Explicit KEDFs. Improvement of one- and two-point KEDFs of the types we have described thus far comprises the great majority of effort. We treat machine-learnt functionals separately below. But there are other lines of exploration of analytical forms. This section therefore provides a brief survey of those other lines of which we are aware, with a crucial caveat. Given the significance of DFT and of T_s in its utilization, it is essentially inevitable that we will have failed to notice some of those logic roads less-traveled. The reader thus cautioned, we press on.

Sporadically in DFT development since the publication of the HK theorems, the relationship of T_s to information theory concepts has drawn interest.³³⁰ Such investigations date to Sears, Parr, and Dinur²⁴³ in 1980. The part of their discussion that has remained relevant to OFDFT goes as follows.¹⁹² Factor the number density $n(\mathbf{r})$ out of the squared-modulus of some N_e -body wave function of interest to get a conditional $N_e - 1$ electron density:

$$N_e \frac{\Psi^*(\mathbf{r}_1 \dots \mathbf{r}_{N_e}) \Psi(\mathbf{r}_1 \dots \mathbf{r}_{N_e})}{n(\mathbf{r}_1)} := f(\mathbf{r}_2 \dots \mathbf{r}_{N_e} \parallel \mathbf{r}_1) \quad (121)$$

The expectation value of the N_e body KE operator then is

$$\begin{aligned} T[\Psi] &= \frac{1}{8} \int d\mathbf{r}_1 \dots d\mathbf{r}_{N_e} \frac{|\nabla_1 n f|^2}{n f} \\ &= T_{vW}[n] + \frac{1}{8} \int d\mathbf{r}_1 \int d\mathbf{r}_2 \dots d\mathbf{r}_{N_e-1} \frac{|\nabla_1 f(\mathbf{r}_1 \dots \mathbf{r}_{N_e})|^2}{f(\mathbf{r}_1 \dots \mathbf{r}_{N_e})} \end{aligned} \quad (122)$$

The von Weizsäcker term is identified with the Fisher information entropy, a measure of localization. If the state $\Psi(\mathbf{r}_1 \dots \mathbf{r}_{N_e})$ of interest is the full many-fermion ground state, the nonlocal term beyond T_{vW} has been called the correlation part^{65,66,331,332} and is discussed in terms of the Shannon

information entropy. (Note that this is *not* the correlation KE as defined in KS DFT; recall above.)

Of more direct relevance for OFDFT, if the state $\Psi(\mathbf{r}_1 \dots \mathbf{r}_{N_e})$ is chosen to be the KS determinant, $\Phi_{KS}(\mathbf{r}_1 \dots \mathbf{r}_{N_e})$, then eq 122 yields

$$T_s[\Phi_{KS}] = T_{vW}[n] + \frac{1}{8} \int d\mathbf{r}_1 \int d\mathbf{r}_2 \dots d\mathbf{r}_{N_e-1} \times \frac{|\nabla f_{KS}(\mathbf{r}_1 \dots \mathbf{r}_{N_e} \parallel \mathbf{r}_1)|^2}{f_{KS}(\mathbf{r}_1 \dots \mathbf{r}_{N_e} \parallel \mathbf{r}_1)} \quad (123)$$

Manifestly, the second term is $T_\theta[n]$, eq 35. Since $f_{KS}(\mathbf{r}_1 \dots \mathbf{r}_{N_e} \parallel \mathbf{r}_1) \geq 0$, this decomposition is a direct demonstration of Pauli term positivity and, hence, also of the lower-bound property of T_{vW} .

The same factorization route in fact was taken by Levy, Perdew, and Sahni³³³ to demonstrate the existence, independent of the HK formulation of DFT, of a density eigenvalue equation akin to eq 38,

$$\left\{ -\frac{1}{2} \nabla^2 + v_{\text{effective}}(\mathbf{r}) \right\} n^{1/2}(\mathbf{r}) = \mu n^{1/2}(\mathbf{r}) \quad (124)$$

but with detailed contributions directly from the N_e -electron wave function and $N_e - 1$ conditional factor. Equivalent routes have been explored^{334,335} in specific applications of the exact factorization approach.³³⁶

A different way to exploit the decomposition is to do a Levy–Lieb constrained search on the conditional factor $f(\mathbf{r}_2 \dots \mathbf{r}_{N_e} \parallel \mathbf{r}_1)$ via Monte Carlo calculations.^{65,337} The approach is to use a model form for $f(\mathbf{r}_2 \dots \mathbf{r}_{N_e})$ constructed by considering necessary conditions on the N_e -fermion wave function in conjunction with Monte Carlo sampling on the HEG over a finite range of comparatively high densities $0.55 \leq r_s \leq 1.81$. The result was a proposed KE functional which, in our notation, is

$$T_{GDS08}[n] = T_W[n] + \int d\mathbf{r} n(\mathbf{r}) [A_1 + B_1 \ln n(\mathbf{r})] \quad (125)$$

with $A_1 = 0.860 \pm 0.022$ and $B_1 = 0.224 \pm 0.012$. Later another version⁶⁶ led to what was denoted as GHDS10,

$$T_{GHDS10}[n] = T_W + T_{TF} + \int d\mathbf{r} n(\mathbf{r}) [A_2 + B_2 \ln n(\mathbf{r})] \quad (126)$$

with $A_2 = 1.02$ and $B_2 = 0.163$.

Both functionals have an evident information-theoretic structure and, as well, the same structure as the intrinsic free energy of the classical ideal gas; see ref 338, Section 3.1. That illustrates a problem discussed in ref 339. Both functionals violate the positivity constraint eq 35 except for certain density ranges. The analysis led to a proposed modified information-theoretical form that also is positive definite. How to implement that modified form is not obvious, since it relies on the use of a maximal bounding function for the density. The examples given in ref 339 suffice for the purposes of analysis but do not seem practical.

The GDS08 and CHDS10 KEDFs also violate uniform scaling requirements, a difficulty addressed by Delle Site³⁴⁰ with an exponential ansatz for the conditional factor. That paper also connected the conditional factor to a classical partition function. Additional analysis in the broader context of

information theory is in ref 341. More recently, Seshaditya et al.³⁴² have used the same approach to discuss changes in approximate functionals associated with changes in system dimensionality.

Two other connections deserve mention. There is a version of DFT formulated as a local thermodynamics which Nagy recently has shown to be connected with information theory. See ref 330 and citations therein. A version of the conditional probability factorization as an approach to the XC energy has appeared very recently.³⁴³ In it, a classical approximation for the effect of the selected electron is exploited. So far as we know, the approach has not been applied to the KEDF problem.

An alternative route to constructing an approximate KEDF is one we have mentioned already, namely to approximate $v_s := v_{vW} + v_\theta$ directly. Given that, one could solve eq 38 self-consistently for $n(\mathbf{r})$. Then the total energy could be evaluated by either the virial relation, eq 45, or the line integral, eq 77, already mentioned. The approach via the virial relation (or similar “bi-functional”; see below) is particularly well adapted to single-point approximations. Such schemes are examples of potential functional theory.^{344–347} They exemplify the observation that “An energy functional of the density is often useful even though the density is obtained by a separate method.”³⁴⁸

Regarding OFDFT specifically, the potential functional approach in the one-point context seems to have been taken up first in earnest by Chai and Weeks^{48,349} (their two-point scheme, discussed already, came later). In a way it was a confirmation of earlier study by Herring.²⁴⁵ He had concluded that it seemed formidably difficult to devise a v_s approximation that satisfied the counterpart to the nonlocal spatial symmetry differentiation requirement, eq 85. Chai and Weeks argued, on the other hand, that the nonlocality of τ_s is substantially more complicated than that of v_s because the integral of τ_s over all space is required (recall the two-point argument) whereas v_s is local. In ref 349 therefore, they contrived a one-point scheme for the kinetic potential that, in our notation, has the Pauli part

$$v_\theta^{MTF} = \frac{5}{3} c_{TF} n^{2/3} - \frac{|\nabla n|^2}{8n^2} + \frac{1-\alpha}{4} \frac{\nabla^2 n}{n} \quad (127)$$

(“MTF” means modified Thomas–Fermi, Chai and Weeks’ term.) The value $\alpha = 1$ gives the correct Kato cusp condition behavior of $n(r)$ at atomic nuclei but poor $n(r=0)$ values, a result that is unsurprising since the $\nabla^2 n$ contribution is removed from v_θ^{MTF} for that α . For the H atom for example, $v_{\theta}^{C-W} < 0$ for $r > 1.12$ au. To keep v_θ^{MTF} from going negative asymptotically for an exponential density requires $\alpha \leq \frac{1}{2}$. Chai and Weeks chose $\alpha = \frac{1}{2}$ for further investigation of MTF because that value gave the correct asymptotic behavior for n . That choice sacrifices the nuclear cusp condition, because v_θ^{MTF} has a negative singularity at the origin.

The bare density Laplacian in v_θ^{MTF} does not contribute to the linear response, so the MTF linear response is the same as from TFλvW with $\lambda = \frac{1}{2}$. Atomic total energies obtained from v_θ^{MTF} by the line integral method are below the KS values, so if v_θ^{MTF} corresponds to the functional derivative of some (unknown) T_θ^{MTF} , that putative KEDF is not N -representable.

As MTF is strictly local, Chai and Weeks addressed its deficiencies by constructing the nonlocal kinetic potential⁴⁸ that we discussed above; see eq 102 and text therewith.

Construction of explicit potential functionals for OFDFT via exploitation of virial-type relations seems to have lain fallow until taken up again by Finzel and co-workers.^{350–355} To summarize an extensive series of papers, introduction of explicit atomic shell structure is undertaken by exploitation of expressions akin to the virial relation, eq 45. Critically, however, they take it as a “bi-functional” of both $n(\mathbf{r})$ and some generating potential v_g

$$T_\theta = -\frac{1}{2} \int d\mathbf{r} n(\mathbf{r}) \mathbf{r} \cdot \nabla v_g(\mathbf{r}) \quad (128)$$

See Lemma 2, corollary in the Appendix of ref 35 for justification. If v_g were to be v_θ , the resulting T_θ value would be exact for the exact density. Instead what Finzel and co-workers consider is the construction of physically and chemically sensible approximations to v_g that may in fact not be the functional derivative of anything but nevertheless give T_θ estimates of useful accuracy.

Perhaps the starker example is to use a stepped potential.³⁵⁰ Here we give a slight generalization of that in the bifunctional setting to get to the proper result for a central-field atom of N_{shell} shells. Assume that the N_e -electron density is entirely inside a radius R_{shell} . Then

$$N_e = 4\pi \int_0^{R_{\text{shell}}} dr r^2 n(r) \quad (129)$$

Choose

$$v_g(r) = -\frac{1}{2\pi} \sum_{i=1}^{N_{\text{shell}}} k(r/R_{\text{shell}}) \ln(r/R_{\text{shell}}) \quad (130)$$

and positive constants k_i such that

$$k(r/R_{\text{shell}}) := k_i; \quad r_{i-1} \leq r < r_i; \quad i = 1, \dots, N_{\text{shells}} \quad (131)$$

with $k_{i+1} > k_i$ to mimic mean shell energies, $r_0 = 0$, and the other r_i spaced in some chemically sensible way. Observe that the only character this v_g shares self-evidently with a proper atomic v_θ is that it is everywhere positive and decreases radially. For candidate density, eq 128 then gives

$$\begin{aligned} T_\theta[n] &= -2\pi \int_0^{R_{\text{shell}}} dr r^2 n(r) r \frac{\partial v_g}{\partial r} \\ &= R_{\text{shell}} \sum_{i=1}^{N_{\text{shell}}} k_i \int_{r_{i-1}}^{r_i} dr r^2 n(r) \end{aligned} \quad (132)$$

Ergo, one can get a sensible numerical value for the Pauli KE with a step-structured potential v_g that has no obvious relationship (indeed, may have none other than generic character) to the functional derivative $\delta T_\theta / \delta n$.

Much more realistic generating potentials can be constructed from atomic constituents or fragments of molecules or prototype molecules (“pro-molecules”) assembled from atomic arrays. See refs 352–354. The challenge of this approach is, of course, that precisely because v_g does not have a known connection with some $\delta T_\theta / \delta n$, development of better v_g models requires very detailed and inventive analysis of the shell structure and bonding regions of highly diverse types of densities. The other issue is the candidate density itself. Again, pro-molecules are promising candidates.

Direct solution for the Pauli potential also has been considered via a differential equation. For that line, see refs

356 and 357. The essential idea is that the OFDFT Euler equation for $n^{1/2}(\mathbf{r})$, eq 38, and the differential equation for $v_\theta(\mathbf{r})$ can be solved self-consistently by cycling between solving for the density for a given $v_\theta(\mathbf{r})$ and then updating $v_\theta(\mathbf{r})$ with the new density. As actually implemented in the proof-of-concept calculation on the Be atom in ref 357, the differential equation for v_θ had contributions from the known solution to the ordinary KS problem. The argument in favor of pursuing the approach without such knowledge is that it may be easier to construct functions of \mathbf{r} that approximate those contributions than it is to construct KEDFs, which are functionals of n .

A different kind of decoupling of the Pauli KE and the Pauli potential appeared very recently. Recall mention, at the end of Section 2.4.1, of the WPBEK GGA KEDF.²⁹⁸ To cope with its positivity violations, those authors proposed a remedy that does not, in fact, depend on the details of F_θ^{WPBEK} but can be presented usefully for any GGA F_θ^{approx} .

As a first step, ref 298 introduces two reference enhancement factors [their eq 43],

$$F_{\theta,i}(s) = \frac{\exp \alpha s^2}{(1 + b_i s^4)^2} \quad i = 1, 2 \quad (133)$$

They set $\alpha = \mu_T - (5/3)$ with $\mu_T = 5/27$ (from the gradient expansion) or 0.23889 (from the semiclassical neutral atom). The parameters b_i were determined by calculating

$$I_i = \int d\mathbf{r} \tau_{\text{TF}}(\mathbf{r}) F_{\theta,i}(\mathbf{r}) \quad (134)$$

with the HF Li atom density for $i = 1$ and the HF Rn density for $i = 2$ and requiring that the results satisfy

$$\begin{aligned} T_{\theta,1} &< T_{\theta,\text{Li}} \\ T_{\theta,2} &> T_{\theta,\text{Rn}} \end{aligned} \quad (135)$$

The reference values on the RHS come directly from the atomic calculation by simple subtraction of the von Weizsäcker KE. The result is $b_1 = 50$ and $b_2 = 0.5$.

With the parameters set, the next step is to impose the condition

$$\begin{aligned} T_\theta^{\text{approx}} &:= \int d\mathbf{r} \tau_{\text{TF}}(n(\mathbf{r})) F_\theta^{\text{approx}} n(\mathbf{r}) = A_\theta I_1(n(\mathbf{r})) \\ &+ (1 - A_\theta) I_2(n(\mathbf{r})) \Rightarrow A_\theta = \frac{I_2 - T_\theta^{\text{approx}}}{I_1 - I_2} \end{aligned} \quad (136)$$

One therefore has a different Pauli enhancement factor that is properly positive but yields the same KE as the original F_θ^{approx} , namely

$$F_\theta^{\text{mix}} := A_\theta F_{\theta,1} + (1 - A_\theta) F_{\theta,2} \quad (137)$$

It is this F_θ^{mix} that would form the $v_\theta = \delta T_\theta / \delta n$ in the OFDFT Euler equation. Though ref 298 obtained numerical results solely post-SCF with HF atomic densities, a reasonably obvious next step would be to evaluate A_θ on a pro-molecule (to avoid having to calculate $\delta A_\theta / \delta n$ at each SCF cycle) and use the result as a GGA KEDF for the physical system for which the pro-molecule is the counterpart.

We conclude this section with pointers to three other elements of the literature. First, there is a relatively recent effort to formulate corrections to the TF density without the conventional gradient expansion.³⁵⁸ The essential idea³⁵⁹ is to

“trotterize” the density written in terms of the time evolution given by the KS Hamiltonian

$$n(\hat{H}_{\text{KS}}(\hat{\mathbf{q}}, \hat{\mathbf{p}})) := \int_C \frac{dt}{2\pi it} e^{-i\hat{H}_{\text{KS}}(\hat{\mathbf{q}}, \hat{\mathbf{p}})t} \quad (138)$$

The contour is the real axis from $-\infty$ to ∞ with the exclusion of a small circle in the lower half-plane centered on the origin. Expressed in terms of the quantum mechanical position and momentum operators, the KS Hamiltonian is

$$\hat{H}_{\text{KS}}(\hat{\mathbf{q}}, \hat{\mathbf{p}}) = \frac{\hat{\mathbf{p}}^2}{2} + \mu_{\text{KS}}(\hat{\mathbf{q}}|n) \equiv \frac{\hat{\mathbf{p}}^2}{2} + \mu - v_{\text{KS}}(\hat{\mathbf{q}}|n) \quad (139)$$

With these expressions, one can construct approximate factorizations of the exponential. For example, the first correction past TF given in ref 358 is

$$U_S(t) = e^{-iy_1 t(\hat{\mathbf{p}}^2/2)} e^{ix_1 t \mu_{\text{KS}}(\hat{\mathbf{q}}|n)} e^{-iy_0 t(\hat{\mathbf{p}}^2/2)} e^{ix_2 t \mu_{\text{KS}}(\hat{\mathbf{q}}|n)} e^{-iy_2 t(\hat{\mathbf{p}}^2/2m)} \quad (140)$$

with parameters x_1 , x_2 , y_0 , y_1 , and y_2 determined by analysis of the requirement $\nabla \nabla \mu_{\text{KS}} = 0$. Whether the various approximations that result from this procedure can be used to construct or constrain better KEDFs remains to be seen.

Second, semiclassical corrections to the TF model also have been attempted^{25,40} and still are an active avenue of research.³⁶⁰ For the sake of a contained presentation, we do not discuss these schemes further.

Third, though there has been considerable effort on local-scaling approaches to KEDFs, little seems to have occurred recently. Therefore, we refer the reader to refs 192 and 217 and references therein.

2.5. Machine-Learnt KEDFs

In recent years, machine learning (ML) methods have permeated most fields of science, including electronic structure theory.^{361–365} The main focus has been to accelerate the computationally intensive parts of an atomistic model, from electronic structure with correlated wave function methods,^{366–369} to the fitting of potential energy surfaces,^{370,371} to the prediction of quantities specific to KSDFT.^{82,83,147–154,372–381}

When fitting energy surfaces (either for small molecular systems or for condensed phases) the total electronic energy or potential energy surface is learned using an appropriate representation of the nuclear positions

$$\{Z_\alpha, R_\alpha\} \xrightarrow{\text{ML}} E(\{Z_\alpha, R_\alpha\}) \quad (141)$$

In the present context, this section focuses on ML techniques that can be used to predict electron densities, KEDFs, and other pure functionals of the density. The learning procedure therefore is centered on the founding theorems of DFT. Recall that they establish the bijective maps

$$n(\mathbf{r}) \leftrightarrow \mathcal{E}[n] \quad (142)$$

$$n(\mathbf{r}) \leftrightarrow v_{\text{ext}}(\mathbf{r}) \quad (143)$$

The map eq 142 has been exploited to learn XC DFAs in recent years with encouraging results for model systems³⁷⁶ and molecules.^{375,377,380} A general purpose double-hybrid XC DFA also has been optimized with supervised learning techniques.³⁸² Perhaps the most impressive achievement is the DeepMind21 (DM21) DFA.³⁸¹ It derives completely from a ML model trained on molecular data as well as fictitious

systems carrying a fractional number of electrons. DM21 is found to treat systems successfully that are problematic for physics-based XC functionals (i.e., over-delocalization of the spin density and the treatment of electronic strong correlation). In a follow-up analysis,³⁸³ DM21 was found to overestimate the hydrogen bond strength in water systematically, thereby pointing to the need for further improvement of the functional in order for predictive chemistry and materials science simulations to be undertaken with confidence.

Moving closer to the subject of this review, several works have focused on either the non-interacting kinetic energy or the general case,^{82,83,147–150,152–154,360,373,374,378} eq 17, of the universal functional. As of now, ML KEDFs have not supplanted those developed from physical and mathematical reasoning as discussed in the preceding subsections. However, the general trend is that ML functionals improve as ML techniques also improve.¹⁵⁴

An interesting and important hurdle that developers of ML-based functionals need to overcome is the inability, so far, of such functionals to provide accurate functional derivatives. Specifically, learning the energy functional does not guarantee the generation of an accurate functional derivative (the associated potential). Without that, the calculation of the density by direct minimization methods would be extremely challenging. Instead, the issue was tackled first by nonlinear denoising techniques.³⁷³ Meyer et al.¹⁵⁵ as well as Ryczko et al.¹⁶⁰ showed that a neural network trained concurrently on the functional and its derivative achieved excellent results for both one-dimensional and complex three-dimensional systems. Ryczko et al.¹⁶⁰ also showed that it is possible to increase the number of training data points by utilizing density-KEDF pairs for non-self-consistent v_{KS} potentials. They could train a neural network with data extracted from only two single-point conventional KSDFT simulations.

In a recent study, Imoto et al. reported¹⁵⁸ encouraging results for selected bulk materials, ranging from metals to semiconductors and rock salt. The electron densities obtained from a ML model (neural network) improve upon the densities resulting from modern GGA KEDFs (such as PGSL¹⁷² and LKT⁸⁶). However, bulk properties derived from the model still are not competitive with physics-based KEDFs.

A fresh approach is simply to learn both the density-to-energy map and the external potential-to-density map as in eq 143.¹⁴⁸ It is worthwhile to sketch how the map in eq 143 was learned in ref 148 by use of kernel ridge regression. The density, $n(\mathbf{r})$, is given as a functional of the external potential, $n[v_{\text{ext}}](\mathbf{r})$, using a regression with a Gaussian kernel,

$$n[v_{\text{ext}}](\mathbf{r}) = \sum_i^{N_s} \beta_i(\mathbf{r}) k[v_{\text{ext}}, v_i] \quad (144)$$

Observe that the kernel function depends on the external potential v_{ext} as well as the external potentials considered in the training set, $\{v_i\}$, whose number of samples is N_s . The chosen Gaussian kernel has the form

$$k[v_{\text{ext}}, v_i](\mathbf{r}) = \exp \left[-\frac{\int (v_{\text{ext}}(\mathbf{r}) - v_i(\mathbf{r}))^2 d\mathbf{r}}{2\sigma^2} \right] \quad (145)$$

where σ is a hyperparameter of the regression. (Remark: Hyperparameters are adjustable parameters to fine tune the regression.) The electron density and the unknown functions,

$\beta_i(\mathbf{r})$, were expanded in a plane wave basis. With that, eq 144 is cast as a linear system which is solved with order $O(N_s^3)$ operations. The scaling of the algorithm is deceptive because the linear system needs to be solved only once. Once the solution is found, the map in eq 143 has been learnt and can be evaluated with order $O(N_s)$ complexity through eq 144.

Once the map of eq 143 is known, it is natural to use the map in eq 142 to learn the electronic energy. This also was done in refs 84 and 148, which showed that the resulting method could be employed in an AIMD simulation of a medium-sized organic molecule recovering known dynamical processes. The regression in eq 144 naturally recovers the potential-to-density map for potentials close to the training set. Reference 84 further dwelled on this issue and showed that it is possible to learn a single map for a small set of chemically similar molecules (resorcinol, phenol, and benzene were tested).

The same research groups showed that the ML procedure can be improved further by relating high-level total energies (e.g., computed by coupled cluster methods) with the electron density from conventional KSDFT calculations with a GGA DFA (termed Δ learning). The outcome was results very close to those of the CCSD(T) reference. Those are much improved compared even to the conventional KSDFT values.⁸⁴

Besides the work in refs 84 and 148, other lines of research for generating ML KEDFs have been to construct density-based descriptors (such as $n(\mathbf{r})$, $|\nabla n(\mathbf{r})|$, $\nabla^2 n(\mathbf{r})$) to be used as a basis to train neural networks (NN). In principle, however, only the electron density should be needed as its derivatives are purely dependent on the density itself. Meyer et al.,¹⁵⁵ for example, employed only $n(\mathbf{r})$ as the main descriptor.

There are studies that seek to learn the kinetic energy density,^{83,150,153,154} an approach that implicates the gauge problem discussed previously. Interestingly, the ML focus in that context so far has been on the nonpositive definite KS KE density, i.e., $\tau_s(\mathbf{r}) = -\frac{1}{2} \sum_i^{\text{occ}} \varphi_i(\mathbf{r}) \nabla^2 \varphi_i(\mathbf{r})$. However, because it differs from the positive-definite form (see eq 23) only in a surface term that depends solely on the density, $\frac{1}{4} \nabla^2 n(\mathbf{r})$, the learning procedure can be carried out on either form of the KS KE density.

An alternative to that approach proposed recently¹⁵⁹ is to learn a quantity that carries information about the Pauli potential, defined in eq 36, rather than the KE density. Reference 159 proposes to use the source of the Pauli potential, defined as $\rho_c(\mathbf{r}) = 4\pi \nabla^2 v_\theta(\mathbf{r})$, rather than v_θ itself. Though the results seem encouraging for predicting the electronic structure of atomic systems, the concept is fairly unusual and needs additional exploration and validation.

In the studies mentioned above, once the descriptors and the KS KE density, $\tau_s(\mathbf{r})$, and/or potential, $v_s(\mathbf{r})$, are generated by a conventional KSDFT calculation of the electron density, the NN can be trained against $\tau_s(\mathbf{r})$. There are various options for approximating energy and potentials by regression. However, it appears that convolutional NNs provide superior representation of both potential and energy¹⁵⁵ in comparison to kernel ridge regression because of their ability to capture nonlocal correlations (i.e., the ones that are captured by two-point functionals).

Another important line of research related to ML and DFT has been the quest to predict molecular and periodic system electron densities directly. The challenge addressed is this: given atom-centered descriptors, is it possible to predict the

electron density of molecules and periodic systems? For example, in refs 384–387 the electron density is the target of various ML approaches. Virtually all methods surveyed utilize atom-centered basis functions to represent the electron density around each atom. The functions range from Gaussians multiplied by spherical harmonics to numerical functions. These produce electron densities with surprising transferability,³⁸⁵ even though the deviation of the predicted densities from the parent KSDFT method averages a few percent (or a few tens of meV/atom when energies are considered^{384,387}). While such accuracy can be regarded as already quite impressive, it still does not reach chemical accuracy.

It is clear that ML is making important contributions to KEDF development. They are almost certain to increase in number and impact, in parallel with ML development of XC functionals.^{381,388} It also is clear that the same exact conditions that constrain the development of physics-based KEDFs should play an increasingly important role in ML-based KEDFs to guide the training toward accurate and transferable KEDFs.^{149,372,389} See, for example, refs 381, 390, and 391.

2.6. Testing

From early on, KEDFs were tested for accuracy of their KE values calculated on HF densities. Though reasonably motivated in the sense of historical context, in our opinion there is little value to that procedure now (except perhaps for comparison with earlier literature). More than 15 years ago, ref 63 showed that a more meaningful post-SCF screening of KEDFs is to evaluate whether they give molecular binding when evaluated on the correct KS density (for a prescribed XC DFA of course). The argument is plain. A perfect KEDF would reproduce the binding energy and electron density (both as a function of geometric parameters) from the ordinary KS solution with a prescribed XC DFA. Evaluation of a candidate KEDF on the conventional KS density therefore tests whether that KEDF can deliver the right energy for the right density. If, at that stage, the deviations are larger than the target accuracy for the KEDF, then it is not worthwhile to test the v_θ for that KEDF in the OFDFT Euler equation. This is precisely the identification of functional-driven errors as distinct from density-driven errors as done for XC potentials in the conventional KS context, except that here the reference density is, of course, that from the conventional KS treatment.^{392,393}

Reference 63 also made an explicit choice of criteria for testing. The limitations of simple GGA KEDFs led them to a design choice: "... we are compelled to refocus on the original objective: an OF-KE functional parametrization solely for reproducing the KS forces, irrespective of the resulting total energy." Because the forces on the atoms depend solely on the electron density, the requirement to match the forces determined by OFDFT with an approximate KEDF with the conventional KSDFT forces is equivalent to requiring matching of the electron densities of the two methods.

It is not uncommon to study results from a KEDF for which the errors may be below the conventional (orbital) T_s . Such cases are clear violations of N -representability. Therefore, it would seem to be more appropriate to emphasize those energy deviations that are below the conventional (orbital) T_s compared to those that lie above. Doing so would bias construction of KEDFs to minimize the likelihood of non- N -representability.

As an example of retrospective testing, note that the non- N -representability of $\text{TF}\lambda v\text{W}$ with $\lambda < 1$ also means that both the HCD-1 and HCD-2¹²² KEDFs also are not N -representable, since $T_{\text{HCD}}[n] \leq T_{\text{TF}(1/S)VW}[n]$.

The gauge ambiguity of the KE density discussed above should be taken into consideration when devising measures for assessment of a KEDF. Garcia-Aldea and Alvarellos¹⁴⁶ introduced what they called a quality factor, defined as

$$\sigma_{\text{GAA}} := \frac{\int d\mathbf{r} |\tau_s(\mathbf{r}) - \tau_{\text{approx}}(\mathbf{r})|}{T_s} \quad (146)$$

This measure tends to favor TF compared to many other one-point functionals. Considering the Teller nonbinding theorem,³⁹ that is not good. (Note that testing based on the mean absolute relative error, or MARE, also can favor TF, for example, with the Perdew–Constantin KEDF; recall the discussion just after eq 70 above.) However, the Garcia-Aldea–Alvarellos measure does discriminate very clearly between Laplacian-dependent KEDFs and GGAs; see Table 1 in ref 233. This reinforces focus on the objective of the testing. The σ_{GAA} tests the approximate KE density, not the electron density or the possible non- N -representability of the functional.

Above we noted that other quality measures have been introduced and used sparingly, e.g. the “relative performance indicator”²⁹¹ and “global performance indicator”.¹⁷²

This summary illustrates that the OFDFT community is lacking an agreed-upon and comprehensive set of indicators of the quality of proposed KEDFs. Also missing are data sets of molecules, solids, and model systems on which to benchmark KEDFs. The contrast with quantum chemistry validation of XC DFAs via widely accepted data sets, for example refs 287, 394–398, is striking and needs to be addressed to expedite development and at the same time weed out unpromising avenues of research.

2.7. Going beyond Typical KEDF and Local Pseudopotentials

As already mentioned and as is discussed in detail in Section 3, current OFDFT simulation implementations for materials chemistry and physics applications rely on local pseudopotentials (LPPs). That effectively limits OFDFT simulations at present to that portion of the periodic table for which LPP-based KSDFT calculations reproduce the results of nonlocal pseudopotential conventional KSDFT calculations. We defer detailed discussion of LPPs to Section 3. Here we consider ways to avoid or reduce the challenge.

In addition to the LPP hurdle, as we have seen in the previous discussion, several of the currently best-available KEDFs are not broadly transferable. Recall, for example, the limitations from parametrization of the HC and WGC KEDFs.

Given these obstacles to wider use of OFDFT, it is unsurprising that approaches aimed at going beyond the basic OFDFT scheme have turned up. Some exploit a density decomposition approach in which $n(\mathbf{r})$ is split into regions that are treated individually with a suitably adapted KEDF. Others attempt to recognize orbital angular momentum locally in the KEDF and pseudopotentials. A third scheme is an approximate all-electron treatment via the projector augmented wave (PAW) scheme.³⁹⁹ We review these approaches in what follows.

2.7.1. Density Decomposition-Based OFDFT. The insight behind the density decomposition approach^{78,400} is

that it is possible to distinguish regions wherein the local density variations dominate contributions to τ_s from those regions wherein nonlocal (delocalized) variations dominate. (The conceptualization resembles the cellular decomposition into spherical and interstitial regions of the augmented plane wave method.^{401,402}) An early one-point version of the idea was put forth by Tal and Bader.²⁴⁶ They wrote the density as the sum of an atomic-core-like part $n_1(\mathbf{r})$ and a more diffuse remainder $n_2(\mathbf{r})$ and wrote the KE density as $\tau_{vW}[n_1] + \tau_{vW}[n_2] + \tau_{TF}[n_2]$. We do not know of any subsequent exploration of this scheme. The present-day versions of density decomposition take the perspective that a one-point KEDF, $T_s^{(1)}$, should be adequate in the regions of localized densities while a two-point KEDF, $T_s^{(2)}$, would be used in the delocalized (“interstitial”) type regions. As we have seen before, nonlocal KEDF kernels must have different limiting behaviors for $q \rightarrow 0$ for metallic or insulating systems. A density decomposition approach attempts to exploit this fact via the hope that the decomposed densities correspond closely enough to either metallic or insulating systems to be described by corresponding KEDFs.

A critical ingredient of any density decomposition scheme is a scale (or switching) function $F(\mathbf{r})$ which distinguishes the sought-after density regions, localized n_{loc} and delocalized n_{del} . Typically^{78,403} $F(\mathbf{r})$ is chosen as a simple function of $n(\mathbf{r})$. Thus, the delocalized density is

$$n_{\text{del}}(\mathbf{r}) = n(\mathbf{r})F(\mathbf{r}) \quad (147)$$

and the localized density is $n_{\text{loc}} = n - n_{\text{del}}$. The KS KE becomes

$$T_s[n] = (T_s^{(1)}[n] - T_s^{(1)}[n_{\text{del}}]) + T_s^{(2)}[n_{\text{del}}] \quad (148)$$

The accuracy of the approach depends on the choices for $T_s^{(1)}$ and $T_s^{(2)}$ (eq 148) and $F(\mathbf{r})$. A physically reasonable choice is to employ a semilocal functional for $T_s^{(1)}$ and either HC⁴⁰³ or WGC⁷⁸ for $T_s^{(2)}$. When the WGC KEDF is adopted, the functional is named WGCD.

Such a decomposition scheme can be cast in a subsystem DFT framework,⁷⁸

$$T_s[n] = T_s^{(1)}[n_{\text{loc}}] + T_s^{(2)}[n_{\text{del}}] + \frac{(T_s^{(1)}[n] - T_s^{(1)}[n_{\text{loc}}] - T_s^{(1)}[n_{\text{del}}])}{\text{Non-additive } T_s^{\text{nad}}[n_{\text{loc}}, n_{\text{del}}]} \quad (149)$$

However, as distinct from subsystem DFT¹⁹⁰ (see subsection below), the variational parameter is still the total electron density, n , not the decomposed densities. The variations in the localized and delocalized densities are propagated through eq 147, with the resulting contribution to the Euler equation

$$\frac{\delta T_s[n]}{\delta n} = \frac{\delta T^{(1)}[n]}{\delta n} - \left[\frac{\delta T^{(1)}[n_{\text{del}}]}{\delta n} - \frac{\delta T^{(2)}[n_{\text{del}}]}{\delta n} \right] \times \left(F(\mathbf{r}) + n(\mathbf{r}) \frac{\delta F}{\delta n} \right) \quad (150)$$

The switching function must be chosen judiciously to avoid overcomplicated structure that makes implementation itself overly complicated.¹⁶⁹ One issue of complication is whether to make $F(\mathbf{r})$ a function independent of the electron density. In ref 78 that was not the case. Instead, the density-dependence of $F[n(\mathbf{r})]$ was determined from an extra self-consistent loop on the density; see Figure 4 of that paper and the attendant

discussion. Such a loop amounts to approximating the $\frac{\delta F}{\delta n}$ contribution to eq 150. A simple numerical example⁴⁰⁴ shows that the functional derivative has a change in sign with respect to n/n_{del} , behavior not found in the double-loop implementation of ref 78. To our knowledge, the effects of that discrepancy have not been studied.

The Carter group applied the density decomposition scheme to transition metals,⁴⁰³ amorphous Li–Si alloys,¹⁶⁹ and covalently bonded molecules and materials.⁷⁸ The results, however, indicated that while the decomposition scheme can improve on the conventional OFDFT methodology (i.e., use of a single KEDF), it does not lead to a substantive, quantitative improvement. Renewed investigation may be an opportunity.

2.7.2. Angular-Momentum-Dependent OFDFT. Another spatial decomposition scheme to get around the limitations of LPPs for OFDFT applications was proposed by Ke et al.^{161,405} Angular-momentum-dependent OFDFT (AMD-OFDFT; almost a misnomer) is a local hybrid approach in which an orbital-angular-momentum-dependent KE for each atom is constructed by way of auxiliary atom-centered 1-rdms. They connect an interstitial density (with no orbital dependence) with ionic cores that are represented by both nonlocal (NLPPs) and local pseudopotentials. The cores are treated solely with the auxiliary 1-rdms. Thus, the KS KE is partitioned into core contributions and an interstitial contribution. The partitioning is achieved by the KE density ansatz

$$\tau_{\text{AMD}}(\mathbf{r}) = \sigma(\mathbf{r})\tau_s(\mathbf{r}) + (1 - \sigma(\mathbf{r}))\tau_{\text{KEDF}}(\mathbf{r}) \quad (151)$$

where $\sigma(\mathbf{r})$ is a switching function and τ_s and τ_{KEDF} are the KE densities from the auxiliary 1-rdms and the approximate KEDF chosen for the interstitial region, respectively. Clearly there are some similarities with the decomposition methods just discussed, but both the regional treatment and manner of switching are different.

The auxiliary 1-rdms are generated starting from a muffin tin (MT) geometry^{401,402} wherein atom-centered spheres are constructed, with the remaining volume being the interstitial region. Then the electron density is expanded in atomic-centered basis functions $\Psi_R(\mathbf{r})$ combined with the on-site density matrix elements, N_R , and the interstitial density, $n_I(\mathbf{r})$. Putting all this together

$$n(\mathbf{r}) = n_I(\mathbf{r}) + \sum_R N_R \Psi_R^*(\mathbf{r}) \Psi_R(\mathbf{r}) \quad (152)$$

In doing so, the angular-momentum-dependent parts of the NLPPs can be accounted for though the use of the nondiagonal parts of the auxiliary 1-rdms. Note that the switching encoded by $\sigma(\mathbf{r})$ takes place at a radius greater than or equal to the cutoff radius of the NLPP.

In this way, the energy functional can be expressed in the general form

$$E^{\text{OF}}[\{N_R\}, n_I] = T_s^{\text{KEDF}}[n] + E_{\text{XC}}[n] + E_H[n] + E_{i-e}^{\text{local}}[n] + E^{\text{NL}}[\{N_R\}, n_I] \quad (153)$$

(The subscript “ $i-e$ ” denotes ion–electron.)

The key part of this scheme is the nonlocal correction term which accounts for errors from the approximation of T_s^{KEDF} in the near-ion core region and includes the NLPP contributions. Ideally, E_{NL} would be precisely the difference between the conventional KSDFT and ordinary OFDFT total energies, namely

$$E^{\text{NL}} = E_{i-e}^{\text{NLPS}} + T_s - T_s^{\text{KEDF}} \quad (154)$$

AMD-OFDFT reproduces the bulk properties of titanium⁴⁰⁵ (a transition metal, thus outside of the applicability of OFDFT with LPPs, at least so far). However, since its introduction, to our knowledge there has been no follow-up work to demonstrate applicability to the broader periodic table.

2.7.3. OFDFT with Nonlocal Pseudopotentials. To employ a NLPP in OFDFT, one can use a model 1-rdm. Recalling the subject of the work by Chakraborty et al.,^{260,261} several approaches have been proposed for formulating a model 1-rdm. Xu et al.⁴⁰⁶ exploited a prescription in which a Gaussian decaying 1-rdm is linked to a local KEDF energy density as follows,

$$\gamma[n](\mathbf{r}, \mathbf{r}') = \rho_q(\mathbf{r}, \mathbf{r}') e^{-s^2(\mathbf{r})/2\beta(\mathbf{r}, \mathbf{r}')} \left[1 + A \left(\frac{s^2(\mathbf{r})}{2\beta(\mathbf{r}, \mathbf{r}')} \right)^2 \right] \quad (155)$$

Here $\beta(\mathbf{r}, \mathbf{r}') = \frac{\beta(\mathbf{r}) + \beta(\mathbf{r}')}{2}$ with $\beta(\mathbf{r}) = \frac{3}{2} \frac{n(\mathbf{r})}{\tau_s[n](\mathbf{r})}$ and $\rho_q(\mathbf{r}, \mathbf{r}') = \left[\frac{n^q(\mathbf{r}) + n^q(\mathbf{r}')}{2} \right]^{1/q}$.

This model 1-rdm, evaluated with approximate KEDF energy densities was employed to determine the nonlocal part of pseudopotentials by tracing the 1-rdm with the nonlocal pseudopotential operator,

$$E_{\text{NL}}[\gamma, v_{\text{NL}}] = \int d\mathbf{r} d\mathbf{r}' \gamma(\mathbf{r}, \mathbf{r}') v_{\text{NL}}(\mathbf{r}, \mathbf{r}') \quad (156)$$

Results reproducing the structure of molten simple metal alloys were presented. In addition, and most importantly, encouraging results for bulk Be and Cd were presented.⁴⁰⁶

2.7.4. OFDFT with the PAW Method. Another interesting development along somewhat similar lines as the AMD-OFDFT scheme is the work by Lehtomäki et al.¹²⁰ It provides an implementation of semilocal OFDFT within the PAW scheme. The PAW method³⁹⁹ is nowadays among the most widely used computational approaches to conventional KSDFT because of the clear advantage it provides over typical pseudopotentials. Specifically, PAW provides a way to reconstruct (approximately in practice) the total KS orbitals rather than just the pseudo-orbitals. This is an interesting advantage also in OFDFT because functionals can be evaluated on all-electron densities.

However, the PAW implementation of OFDFT is still in a relatively early stage. Thus far, it can support only semilocal KEDFs, thereby limiting its applicability and delivered accuracy. Comparison with two-point KEDFs (which only have implementations in more traditional pseudopotential schemes) also obviously is limited.^{120,407}

2.7.5. Subsystem DFT: A Distinct Form of OFDFT Challenge. Quantum embedding is a divide-and-conquer strategy to partition a system (the “full system”, hereafter) into interacting fragments with the aim of avoiding the calculation of full-system quantities^{408,409} while maintaining an accurate quantum-mechanical description at least in the neighborhood of a subsystem of interest and, perhaps, globally as well. Several methods of this type have been developed, e.g., subsystem DFT,^{180,181,189–191,410} density matrix embedding,^{411–414} and Greens function embedding.^{415–418} The quantum embedding methods differ mainly in two aspects. First is the way in which they define subsystems (using the electron density, the Greens

function, or other quantities). Second is the way in which they describe the interactions between subsystems. This review focuses on OFDFT; thus, we concentrate our discussion on subsystem DFT.

Subsystem DFT is an appealing approach for large-scale simulations because it defines subsystems based on a partition of the electron density. Specifically, it uses a sum of subsystem electron densities

$$n(\mathbf{r}) = \sum_I n_I(\mathbf{r}) \quad (157)$$

Those replace the density of the full system, n , as the variational functions. In the same spirit as ordinary OFDFT, subsystem DFT exploits this decomposition via a reformulation of the non-interacting kinetic energy functional. It does so with an ansatz similar to eq 149, to wit

$$T_s[n] = T_s \left[\sum_I n_I \right] = \sum_I T_s[n_I] + \underbrace{\left(T_s[n] - \sum_I T_s[n_I] \right)}_{\text{Non-additive } T_s^{\text{nad}}[\{n_I\}]} \quad (158)$$

This decomposed form provides an important opportunity. The subsystem-additive part of T_s can be evaluated with KS orbitals specific to each subsystem. Those subsystem KS orbitals are not the same as the full system KS orbitals. For one thing, they are not required to be orthogonal with orbitals of other subsystems. The subsystem nonadditive part of T_s can be evaluated in practice with pure density functionals. Therefore, the ansatz in eq 158 provides a route for combining conventional KSDFT with OFDFT in a way that benefits the speedy calculation of molecular condensed phases.

The full energy given as a functional of $n = \sum_I n_I$ is found by substituting eq 158 in eq 27 and including the interaction with the external potential,

$$E \left[\sum_I n_I \right] = \sum_I T_s[n_I] + T_s^{\text{nad}}[\{n_I\}] + E_H \left[\sum_I n_I \right] + E_x \left[\sum_I n_I \right] + E_c \left[\sum_I n_I \right] + \int d\mathbf{r} v_{\text{ext}}(\mathbf{r}) \sum_I n_I(\mathbf{r}) \quad (159)$$

Note the explicit replacement of the full system density in the arguments of the functionals by the sum of subsystem densities.

We acknowledge that the first use of eqs 157–159 (as well as the derived functional derivatives and KS-like equations which are presented below in eqs 160–161) is credited to the works of Senatore and Subbaswamy¹⁹¹ and later, provided a formal footing, by Cortona.⁴¹⁰

We remark that the energy functional in eq 159 is formally equivalent to $E[n]$. Therefore, it can be used in a variational minimization equivalent to eq 19 achieved by searching over particle-conserving variations of each of the subsystem densities, $\frac{\delta E}{\delta n_I} = 0$, $\forall I$. Doing so yields the so-called KS equations with constrained electron density,

$$\left[\frac{-\nabla^2}{2} + v_{\text{KS}}(\mathbf{r}) + v_{\text{emb}}^I(\mathbf{r}) \right] \phi_i^I(\mathbf{r}) = \epsilon_i^I(\mathbf{r}) \phi_i^I(\mathbf{r}) \quad (160)$$

In this expression, the KS-like potential, v_{KS} , is defined in eq 30 and is evaluated with the external potential $v_{\text{ext}}^I(\mathbf{r})$ associated with subsystem I , and $\phi_i^I(\mathbf{r})$ and $v_{\text{emb}}^I(\mathbf{r})$ are the KS orbitals and embedding potential of subsystem I , respectively. Equation 160 has the same structure as the KS equations for the isolated subsystem I . The crucial observation is that in eq 160 the effective external potential is not the external potential of the isolated subsystem but rather the sum of that external potential and the embedding potential, $v_{\text{ext}}^I(\mathbf{r}) + v_{\text{emb}}^I(\mathbf{r})$, which leads to a subsystem electron density, n_I , that generally differs from the electron density of the isolated subsystem. The embedding potential contains the functional derivative of the nonadditive KEDF given in eq 158. Specifically,

$$v_{\text{emb}}^I(\mathbf{r}) = \sum_{J \neq I}^{N_s} \left[\int d\mathbf{r}' \frac{n_J(\mathbf{r}')}{|\mathbf{r} - \mathbf{r}'|} + \sum_J v_{\text{ext}}^J(\mathbf{r}) \right] + \frac{\delta T_s^{\text{nad}}[\{n_I\}]}{\delta n_I(\mathbf{r})} + \frac{\delta E_{\text{xc}}^{\text{nad}}[\{n_I\}]}{\delta n_I(\mathbf{r})} \quad (161)$$

with $\{v_{\text{ext}}^J\}$ being the external potentials associated with each fragment relating to the total external potential as $v_{\text{ext}}(\mathbf{r}) = \sum_I v_{\text{ext}}^I(\mathbf{r})$. We also have introduced the nonadditive XC functional, $E_{\text{xc}}^{\text{nad}}[\{n_I\}] = E_{\text{xc}}[n] - \sum_I E_{\text{xc}}[n_I]$. We remark that the decomposition of the external potential into subsystem contributions is not a necessary step.

Clearly, the accuracy of practical implementations of subsystem DFT is determined by the quality of the nonadditive KEDF employed as well as by the accuracy of the XC DFAs used to solve the KSDFT problems for the fragments. To control computational costs, semilocal nonadditive KEDFs typically are employed,⁴¹⁹ at most at the Laplacian level of complexity.²⁸⁵ However, it is now common knowledge that semilocal nonadditive KEDFs do not provide a proper description of a regime in which the fragment electron densities overlap strongly.^{180,181,420} As originally indicated by Wesolowski and Weber,⁴²¹ subsystem DFT carried out with approximate nonadditive functionals should target weakly interacting subsystems.

Mi et al.⁸⁹ adopted new-generation nonlocal nonadditive KEDFs in subsystem DFT calculations and showed that they improve the computed interaction energies and electron densities considerably compared to commonly employed GGA nonadditive KEDFs. This is especially the case when interfragment electron density overlap is non-negligible. In Section 3 we discuss that application in more depth.

An interesting line of research exploits a “bottom-up” approach rather than a “top-down” approach. That is, what are called the nondecomposable, nonadditive potentials (most commonly for the kinetic energy) rely on an ad-hoc ansatz for the nonadditive potential in an effort to obtain accurate densities and to impose known conditions on the potential⁴²² comparatively easily. Lastra et al. proposed the first nondecomposable, nonadditive kinetic energy potential,⁴²³ followed by others.^{424,425} We refer to ref 422 for additional details and a review of the relevant literature. We also note that error cancellation benefiting the value of KEDF approximants does not necessarily translate to advantageous error cancellation for the derived nonadditive functionals and nonadditive potentials (and thus electron densities).^{189,426,427} Therefore, the bottom-up approach emerges as a viable alternative foregoing the definition of a parent KEDF and aiming to exploit beneficial

error cancellation tailored specifically for nonadditive potentials.

Though our survey is not a complete overview of subsystem DFT, we point out some theoretical and practical complications about this method of which the reader should be aware. Gritsenko⁴²⁸ showed that for subsystem DFT to have a properly defined minimum of the energy functional, eq 159, with respect to variations of each of the n_i , it is not enough to impose the requirement that n_i be ν_{KS} -representable¹⁰⁶ (see discussion above eq 22). This is because at most one can define an *infimum* of the functional. A simple mathematical counterpart problem is one in which a function of a single variable, $f(x)$, is expressed as a function of two variables, e.g., $f(x) \rightarrow f(y + z)$. While $f(x)$ may have a well-defined minimum at $x = x_{min}$ the same cannot be said about $f(y + z)$ because the equation $y + z = x_{min}$ has infinitely many solutions. Though this nonuniqueness problem is severe in principle, convergence problems related to it (or traced to it) never have been reported to our knowledge for practical calculations that used finite basis set expansions of the subsystem KS orbitals and approximate nonadditive KEDFs.^{180,181,429}

The formal framework of subsystem DFT requires that the subsystem densities be non-negative. That is, assuming a system is split into two subsystems with one having density $n_1(\mathbf{r})$ and the other $n_2(\mathbf{r})$, then the formal definition, from eq 157, requires $n_1(\mathbf{r}) = n(\mathbf{r}) - n_2(\mathbf{r})$. Therefore, if n_1 is chosen such that at some spatial points $n(\mathbf{r}) - n_2(\mathbf{r}) < 0$, the subsystem DFT procedure breaks down because of the non- N -representability of $n_1(\mathbf{r})$.⁴³⁰ Such violation of the non-negativity condition is a critical problem for frozen-density embedding simulations.⁴³¹ In those, the environment density, n_2 , typically is the one kept frozen. The violation is circumvented by subsystem DFT because the subsystem densities always are maintained as N -representable throughout the variational minimization by construction. We must remark that frozen-density embedding does not suffer from the nonuniqueness problem mentioned above,⁴²⁸ as the energy density functional admits a unique minimum for a given choice of the frozen-density environment. (We note that the unique solution does not guarantee that the total density, given by the sum of subsystem and frozen densities, yields the true ground-state density; see ref 432.)

Beyond non-negativity, it is useful to consider cases in which a subsystem density vanishes. While that is unexpected in practical calculations, the implications nonetheless must be considered.⁴³³ A clear limitation of the definition of the nonadditive functionals in eq 159 is that the functionals should share the same domain of functional differentiation. However, that can be the case only if all subsystem densities are nonzero everywhere in space. If one subsystem density goes precisely to zero inside a Lebesgue measurable volume element, then such a density is not ν_{KS} -representable in a space that includes that volume. Therefore, the existence of the functional derivatives (and therefore the existence of the embedding potential in eq 161) on the entire space is subject to a positive definite condition for all subsystem densities.

Returning to the formal nonuniqueness of the partitioning in eq 158, it motivated the development of a number of embedding methods that provide a unique embedding potential. For example, partition DFT (PDFT)⁴³⁴ and density functional embedding theory^{435,436} partition the electron density in a way that reproduces the total density of a reference KSDFT calculation (even though the conventional

KSDFT calculation is avoided⁴³⁷). There are approximate formulations of PDFT that exploit nonadditive KEDFs in a fashion akin to subsystem DFT.⁴³⁷ When applied to PDFT, semilocal nonadditive KEDFs seem to yield comparatively better results than in subsystem DFT.²⁹⁴ For example, preliminary results show that covalent bonds could be tackled by this method when a parametrized GGA nonadditive KEDF is employed.²⁹³

3. OFDFT IN PRACTICE

Along with construction of accurate KEDFs, it is critical to have appropriate OFDFT algorithms implemented in reliable, highly efficient codes. Unsurprisingly, for achievement of quasi-linear scaling, $O(N \ln N)$, the devil is in the details, with each of the total energy terms needing attention to obtain a fast codebase. Particular issues include construction of appropriate pseudopotentials for OFDFT (which, as already pointed out, differ from those for conventional KSDFT), approaches for optimization of the electron density, choices of basis sets for representing the density and potentials, and efficient algorithms for system sizes up to and beyond millions of atoms.

3.1. The Need for Local Pseudopotentials

In conventional KS calculations, treatment of the sharp cusps in the KS orbitals induced by nuclear attraction often is avoided by use of pseudopotentials (PPs) or effective core potentials. The reasons are twofold: (1) in periodic systems, PPs generate pseudo-orbitals that are smooth and, thus, can be expanded in a modest number of plane waves, and (2) elimination of explicit core electrons reduces the number of KS states to be computed, be they in a molecule or periodic system.

In OFDFT, PPs are needed for related reasons. First, use of a rather smooth pseudodensity allows evaluation of convolutions [an important class of integrals as will appear later, e.g., in eq 171] with numerical, discrete Fourier transforms. Second, as described in the preceding section, many modern KEDFs are calibrated in some way to the HEG and, thus, can be expected to yield best results for systems with comparatively smooth electron densities. In parallel, therefore, with many implementations of conventional KSDFT,^{178,438–441} accurate and transferable electron–ion PPs are crucial ingredients for OFDFT simulations.^{121,163–165,442}

In the context of conventional KSDFT, PPs have a local part and a nonlocal part and, hence, are nonlocal PPs, or NLPPs. To reproduce the effect of orthogonality between the valence and core orbitals, they use spherical harmonic projectors centered on the atoms. Those give rise to the so-called angular momentum channels of an NLPP. Observe that the meaning of “nonlocal” in connection with NLPPs is quite distinct from the meaning for nonlocal KEDFs. For NLPPs, the term signifies the explicit orbital dependence of the potential, not the spatial separation dependence of two-point functionals.

The typical approach is motivated by the plain fact that since the KS orbitals do not appear explicitly in OFDFT, any PP employed for OFDFT must not include angular momentum dependence and, hence, must be “local”. These are denoted (recall above) as LPPs. However, it is possible, in principle, to use orbital-free prescriptions for the 1-rdm, for example eqs 118–120 as developed in ref 260 as well as prescriptions developed recently by Xu et al.⁴⁰⁶ (see Subsection 2.7.3). Given the novelty of those approaches, they have not seen

wide usage yet. Therefore, this section focuses on LPP development.

3.1.1. Early OFDFT Pseudopotential Prescriptions.

Various LPPs proposed early on for OFDFT can be put into three broad categories.

Empirical or model LPPs^{443–450} typically are comparatively simple analytic functions with parameters adjusted so that several physical properties (e.g., electron density, equilibrium volume, bulk modulus) calculated with the LPP fit experimental results or values from other (presumably reliable) electronic structure calculation such as conventional KSDFT. Empirical LPPs played an important role in early KEDF development, though they were derived mostly for metallic systems.^{46,305,451–453} Empirical LPPs can be adapted successfully to complex simulations whenever the fitting procedure is successful.^{450,454} However, fitting limits their transferability. That, no doubt, is a major reason why such PPs no longer see wide use.

An LPP can be contrived by selecting one of the l angular momentum channels of an extant NLPP. Such a simplified local pseudopotential cannot describe correctly the energetic properties of solid phases, even for simple alkali metal solids (bcc Cs and Rb).⁴⁵⁵ Therefore, these no longer are used.

Atomic LPPs (ALPs) are derived by inverting the Kohn–Sham equations to reproduce the pseudodensity from a NLPP KSDFT calculation.^{163,456} For group-IV elements, conventional KSDFT simulations with such potentials yield good results for bond lengths and lattice constants, but not for total energies and KS orbital energies.⁴⁵⁶ Additionally, ALPs have been found to lack sufficient transferability and accuracy to be applicable to simple semiconductors such as Si.¹⁶³

3.1.2. LPPs Currently Available for OFDFT.

To obtain transferable LPPs, approaches have been devised to account for the specific chemical environment of an ion. Among them the bulk-derived LPP [BLP] type is perhaps the most widely used.^{163,164,457} (Remark: In this section and beyond we use BLP, and the more frequently used acronym in the literature, BLPS or BLPSs, to indicate bulk-derived LPPs.) Compared with ALPs, BLPSs are obtained by inversion of the KSDFT electron pseudodensity from one or more bulk crystal systems instead of from atom densities. The first BLPSs, proposed by Watson et al.,⁴⁵⁷ used OFDFT with a selected KEDF to reproduce, exactly, the reference density from a conventional KSDFT calculation with NLPP. The resulting LPP therefore was *KEDF-dependent*. Those LPPs gave good results for metallic systems, e.g., Li, Na, and Al, but had limited transferability beyond that range. To improve transferability, Carter's group proposed *KEDF-independent* BLPSs based upon inversion of the conventional KSDFT equations rather than the OFDFT equation.^{163,164,458} The main algorithmic steps involved are

1. Obtain the target bulk crystal electron pseudodensity from conventional KSDFT calculations with a norm-conserving NLPP.
2. Obtain the effective conventional KSDFT potential, $v_{eff}(\mathbf{r})$, for the entire crystal from that target pseudodensity via the inverse approach⁴⁵⁹
3. Obtain the corresponding LPP for the bulk crystal, $v_{loc}(\mathbf{r})$, from that $v_{eff}(\mathbf{r})$, by removing the Hartree and XC potentials,

$$v_{loc}(\mathbf{r}) = v_{eff}(\mathbf{r}) - v_H(\mathbf{r}) - v_{xc}(\mathbf{r}) \quad (162)$$

4. Obtain the atomic pseudopotential, $v^{atom}(g)$ (where $g = |\mathbf{G}|$ and \mathbf{G} is a reciprocal space vector), by dividing the Fourier transform of $v_{loc}(\mathbf{r})$, $v_{loc}(\mathbf{G}) = \mathcal{F}[v_{loc}(\mathbf{r})]$, by the structure factor $S(\mathbf{G})$ (which is the Fourier transform of the density of nuclear point charges (see Section 3.5.1):

$$v^{atom}(g) = \frac{1}{N_g} \sum_{|\mathbf{G}|=g} \frac{v_{loc}(\mathbf{G})}{S(\mathbf{G})} \quad (163)$$

Here N_g is the number of \mathbf{G} vectors satisfying $|\mathbf{G}| = g$.

To gain transferability of the BLPSs, multiple bulk structures are considered in the process so as to include a large, varied sample of \mathbf{G} points in eq 163. (The number of allowed \mathbf{G} points is dictated by the specific crystal lattice.) Thus, the utility of $v^{atom}(g)$ depends strongly on the sampling of $v^{atom}(\mathbf{G})$ for the low- \mathbf{G} vectors which, unfortunately, are not plentiful. Moreover, the inversion scheme to obtain $v_{loc}(\mathbf{r})$ typically does not yield the correct Coulombic tail decay. Therefore, additional fitting schemes are needed to obtain reliable BLPSs. Here, we do not review those technical details. The main principles are to try to obtain as closely as possible the correct Coulombic tail of the PP, while reproducing the bulk properties as well as possible. See refs 163, 164, and 442, 456.

BLPSs have been applied successfully to various bulk materials,^{163,164} including the elemental Li, Mg, Al, Si, Ag, Ga, In, P, As, and Sb crystals and various III–V semiconductors;^{77–79} liquid Li,⁴⁶⁰ and to the melting temperature of Li.⁴⁶¹ In sum, KEDF-independent BLPSs seem to be the most accurate, transferable LPPs available presently for bulk systems.⁴⁶²

There have been attempts to improve BLPSs for liquid phases, leading to a strategy to construct “globally optimized” LPPs (goLPSSs).⁴⁴² In that approach, BLPSs are modified by adding Gaussians with parameters fitted to minimize the difference between OFDFT and conventional KSDFT for calculated bulk properties and for forces on individual atoms in liquid phases. The goLPSSs have been constructed for elements such as, Li, Ge, and Ga but are not available yet for other atom types. We remark that the procedure to obtain goLPSSs does not rest on a formally justifiable footing and should, in our view, be considered to be a semiempirical parametrization of the OFDFT procedure.

Another constructive approach is the neutral pseudoatom (NPA) method.^{463–465} It constructs LPPs referenced to an atom immersed in an approximate environment. Specifically, the atom is centered in a spherical cavity, subject to a uniform positive background of density equal to the mean valence electron density of the system under consideration. Often that reference is a liquid metal.^{316,466,467} The cavity radius is determined by equalizing the positive charge removed from the hole with the total number of atomic valence electrons, hence the NPA name.⁴⁶⁸ Anta and Madden⁴⁶⁹ used such a reference system to develop KEDF-dependent ab initio LPPs, which were applied to liquid metal studies.⁴⁷⁰ See also ref 471. Essentially the same approach has been used for many years in high-temperature systems, both as NPA and in “average atoms”. The topic has become a specialty in its own right that no longer is strongly connected to OFDFT, so we leave it to refs 472–475 for leads to that literature.

Returning to explicit LPPs, the fitted normalized linear combination (FNLC) potentials are quite different. Karasiev and Trickey¹²¹ proposed that alternative scheme. It uses information from each of the angular momentum channels,

$v_{NL}^l(\mathbf{r})$, of a reference NLPP in the form of a normalized average, to wit,

$$v_{FNLC}(\mathbf{r}) = \frac{1}{\sum_{l=0}^{l_{\max}} c_l} \sum_{l=0}^{l_{\max}} c_l v_{NL}^l(\mathbf{r}) \quad (164)$$

The weight of the contribution of each angular momentum channel, c_l , is adjusted so that conventional KSDFT calculation with the FNLC yields a good fit to selected equilibrium bulk material properties calculated with the original NLPP. Benchmark studies of Li and Al indicated that this approach can yield very good LPPs for Li and Al. However, those authors recognized the issue in their approach: “A remaining challenge for the OFDFT agenda is to construct a good LPP from an existing non-LPP without appeal to any bulk or aggregate system KS calculations”.¹²¹

Mi et al.¹⁶⁵ took up that challenge with their proposal of optimal effective LPPs (OEPP). Inspired by the importance of the norm-conserving conditions satisfied by NLPPs for conventional KSDFT, they uncovered a fundamental relation between a norm-conserving PP and what could be construed as the ideal norm-conserving LPP, $v_{NCLPP}(\mathbf{r})$,

$$\int_{\Omega} d\mathbf{r} (v_{NCLPP}(\mathbf{r}) - v_{NL}^l(\mathbf{r})) n_l(\mathbf{r}) = 0 \quad , \forall l \quad (165)$$

Here $v_{NL}^l(\mathbf{r})$, $n_l(\mathbf{r})$, and Ω are, respectively, the l -th angular momentum channel of a norm-conserving NLPP, the pseudodensity of the l -th atomic orbital, and the region inside the core radius. By combining this relation with the optimized effective potential (OEP) concept already mentioned,^{236,237,476–479} they showed that an optimized LPP should be constructed via a Slater-type pseudodensity-weighted average of the orbital-dependent norm-conserving PPs subject to preservation of the norm-conserving condition,

$$v_{OEPP}(\mathbf{r}) = \frac{1}{n(\mathbf{r})} \sum_{l=0}^{l_{\max}} v_{NL}^l(\mathbf{r}) f_l n_l(\mathbf{r}) \quad (166)$$

In the foregoing expression, f_l is the occupation number and $n(\mathbf{r}) = \sum_{l=0}^{l_{\max}} f_l n_l(\mathbf{r})$.

The corresponding algorithmic steps are simple. For a given electronic configuration and a given set of core radii r_{ell}^c in the selected NLPP,

1. Obtain the LPP by calculating the density-weighted average as in eq 166;
2. For a single atom, compute the norm-conserving conditions and the deviations of the eigenvalues and pseudo-orbitals from a conventional KSDFT calculation with the OEPP versus those from the original NLPP;
3. Repeat the process by adjusting the r_{ell}^c and electron configuration to optimize the OEPP.

We should clarify that there is a certain freedom in choosing r_{ell}^c when constructing NLPPs as long as their transferability and accuracy are maintained. However, OEPP depends on the parent NLPP through eq 166. Thus, r_{ell}^c can be used within that range to fine-tune the accuracy of the OEPP.

Benchmark tests demonstrated that high-quality OEPPs indeed can be constructed from a given norm-conserving PP without empirical fits for several atom types (e.g., most *s/p*-block elements except for the second row). Unfortunately, for second-row and 3–5d metals, a valid LPP with high

transferability cannot be constructed because the NLPP potentials for different l 's differ greatly and the norm-conserving condition of eq 165 for the LPP in eq 166 cannot be satisfied. Figure 9 provides a pictorial view of the

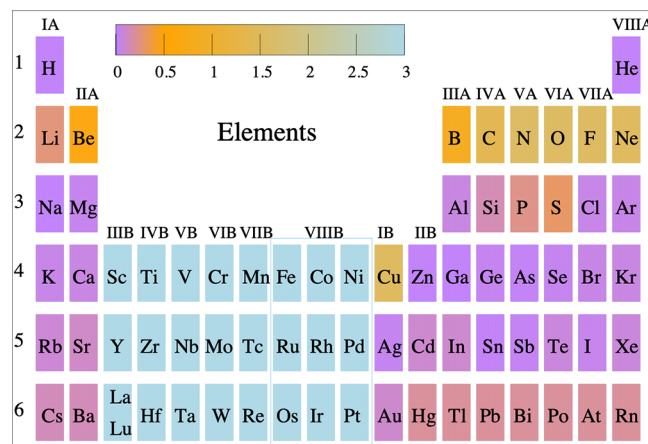


Figure 9. Color map representing the accuracy of OEPP pseudopotentials. The accuracy is assessed by the ability of the OEPP to reproduce the atomic pseudodensity of the parent NLPP as quantified by $\delta_n := \int dr 4\pi r^2 |n_{OEPP}^{ps}(r) - n_{NLPP}^{ps}(r)|$. Here $n_{OEPP}^{ps}(r)$ and $n_{NLPP}^{ps}(r)$ are the corresponding OEPP and NLPP radial pseudodensities, both computed with conventional KSDFT. The LDA XC functional was adopted. Reprinted with permission from ref 165. Copyright (2016) American Institute of Physics.

effectiveness of the OEPP method for each element of the periodic table. The metric used is the preservation of pseudodensity from NLPP to OEPP. Observe that the foregoing remark about the second row and the challenge of the 3d, 4d, and 5d metals is quite apparent.

In summary, there are two important comments to be made regarding pseudopotentials use in OFDFT. First, even though reasonably accurate and transferable LPPs are available for most light metallic elements and some main group elements, for most transition metals and second-row elements, the construction of LPPs with high transferability has proven difficult and may be essentially intractable. Second, our discussion makes clear that developing LPPs for OFDFT is a daunting exercise. Several guiding principles and procedures are available for their construction, but a clear winning strategy has yet to emerge. Thus, the requirement for LPPs is still very much an open problem for OFDFT and perhaps may be its Achilles' heel. New approaches, such as the one based on model 1-rdms summarized in Subsection 2.7, and very recent efforts involving machine-learning pseudopotentials⁴⁸⁰ might prove to be breakthroughs for extending the applicability of OFDFT to a wide selection of elements in the periodic table.

3.2. Variational Optimization

Determination of the OFDFT variational minimum obviously is an optimization problem. The function to minimize is the number-conserving total energy Lagrangian, $\mathcal{L}[n]$, eq 28,

$$n_0 = \operatorname{argmin}_n \left\{ \mathcal{L}[n]; n \geq 0, \int |\nabla n^{1/2}|^2 d\mathbf{r} < +\infty \right\} \quad (167)$$

$$E_0[n_0] = \mathcal{E}[n_0]; \quad (168)$$

The second line reproduces eq 19. Recall that the Lagrangian includes the chemical potential μ to enforce electron number

conservation. The system volume for the integrations, Ω , can be the simulation cell for the case of periodic boundary conditions or the entirety of \mathbb{R}^3 for isolated systems.

As with conventional KSDFT, at least superficially there would seem to be two main approaches for carrying out the optimization. An “indirect” approach converts the minimization into a nonlinear self-consistent field (SCF) problem, while a “direct” approach proceeds as a constrained minimization problem.^{481,482} We summarize key aspects of both.

3.2.1. Solution of the Euler Equation: The SCF and OE-SCF Methods. The indirect approach begins with the one-orbital, OFDFT Euler equation, eq 38, for the square root of the density,

$$\left\{ -\frac{1}{2}V^2 + v_\theta[n](\mathbf{r}) + v_{KS}[n](\mathbf{r}) \right\} \psi(\mathbf{r}) = \mu\psi(\mathbf{r}) \quad (169)$$

where $n(\mathbf{r}) = |\psi(\mathbf{r})|^2$.

Because of the kinship of this equation to the conventional KS equation, one might expect that efficient basis-set methods developed for solution of that equation could be applicable straightforwardly to the OFDFT problem. There are statements to that effect in the literature.³³³ The idea is that a conventional KSDFT SCF code could be converted to SCF solution of eq 169 simply by adding v_θ to v_{KS} , changing the normalization (from 1 to $N_e^{1/2}$), and selecting the lowest eigenvalue. What Chan et al.¹¹⁹ found is that the prescription does not work in practice and, hence, that more robust minimization techniques were needed.¹¹⁹ Subsequently, Karasiev and Trickey¹²¹ confirmed that finding for both atoms and diatomic molecules and for both a Gaussian-type and numerical orbital basis. Note that both papers reported all-electron calculations. Quoting from ref 121, “... use of standard KS codes to solve the OFDFT Euler equation as a modified KS eigenvalue problem is problematic at best. At least for the all-electron case, it seems implausible as a productive route to routine OFDFT calculations.”

Shao et al.⁴⁶² explored this issue with the pseudopotential code DFTpy and showed that the main reason for the failure of customary SCF techniques in the OFDFT context is HOMO–LUMO orbital swapping that arises quite commonly in nearly degenerate systems. As system sizes increase, the HOMO–LUMO gap computed from the OFDFT Hamiltonian with eq 169 decreases; see Figure 10.

To resolve that near-degeneracy problem in conventional KSDFT calculations, it suffices to level shift or to open the minimization to the set of *ensemble* N -representable densities computed from several states above the HOMO level and then to smear the occupations⁴⁸³ (typically with the Fermi–Dirac distribution). Doing so for the OFDFT one-orbital equation would defeat the purpose, since it would involve solving the eigenvalue problem for several levels, hence a regression to a $O(N^3)$ scaling method.

A modified SCF procedure, termed the one-orbital ensemble SCF (OE-SCF), solves that problem.^{462,484} To consider ensemble N -representable densities in OFDFT, OE-SCF begins by direct minimization of $T_{vw}[n] + E_H[n] + E_{xc}[n]$. The Pauli potential is fixed for that minimization and then updated and a new optimization done. These steps are cycled until self-consistency (hence the name OE-SCF, even though no diagonalization ever occurs). Tests show that OE-SCF converges as well as the commonly adopted direct energy minimization of the full energy functional (see next section)

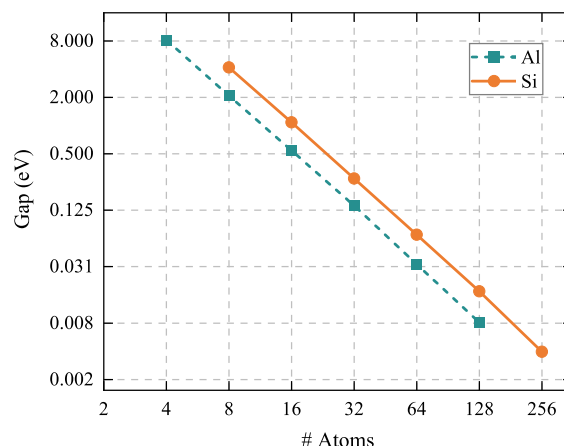


Figure 10. Energy gap between the lowest two energy levels associated with the eigenvalues of the OFDFT Hamiltonian in eq 169 for supercells of Al and Si bulk systems. The plot, in log scale, displays a monotonically decreasing gap with system size. The LMGP KEDF and LDA XC functionals were used. Reprinted with permission from ref 462. Copyright (2021) American Chemical Society.

with one important additional benefit. The Pauli potential needs to be evaluated only a handful of times during the minimization (i.e., once per SCF cycle). For complicated two-point KEDFs, that can yield a large reduction in computational cost, as much as by orders of magnitude.⁴⁶² In Section 4, we report how OE-SCF enables large-scale simulations of complex systems to be done with otherwise computationally expensive nonlocal functionals with a density-dependent kernel, such as LMGP.

OE-SCF currently is implemented in the DFTpy⁵⁴ and ATLAS⁵³ codes. As OE-SCF is rather new, most OFDFT codes still exploit direct energy minimization, to which we turn next.

3.2.2. Optimization by Direct Minimization. There is an immense literature on generic direct optimization, so we note only the aspects directly pertinent to OFDFT implementation here.

Algorithms for direct address of eq 167 must explore variations of a single function of three-dimensional space. To do so typically requires parametrization of the argument function, n , either by expansion in a basis set or by direct discretization of space (which logically is a basis set of delta functions). Thus, the problem becomes the minimization of a function of many variables (proportional to the number of discrete points or basis functions). That can be a simple task for atomic systems rendered in an atom-centered basis. But even in modeling of clusters, the number of variational parameters becomes large enough to require the development of ad-hoc optimizers.^{119,485} When bulk materials are considered, discretization of space with regular grids is the usual choice. That allows use of discrete Fourier transforms to represent the electron density and the potentials and for solving the Poisson equation associated with the classical Coulomb energy (see next section).

Energy minimization algorithms typically include the following steps to obtain the quasi-orbital, ψ , see eq 169:

1. Obtain the optimization direction vector in the space of the variational parameters, \mathbf{p}_k .
2. Find a reasonable step size λ_k along the vector \mathbf{p}_k .

3. Generate an improved quasi-orbital, $\psi_{k+1}(\mathbf{r})$, updating the one from the previous optimization step, $\psi_k(\mathbf{r})$, with \mathbf{p}_k and λ_k .
4. Iterate until convergence criteria are satisfied.

Many algorithms from the optimization literature can be employed for step (1). Some of those provided in the available OFDFT software^{53,54,486} include steepest-descent (SD), nonlinear conjugate gradient (CG),^{487–493} limited memory Broyden–Fletcher–Goldfarb–Shanno (L-BFGS),⁴⁹⁴ and truncated Newton (TN) methods.⁴⁹⁵ Because of efficiency limitations on SD and storage limitations for L-BFGS, CG is the most widely used of these. It is possible to improve its computational efficiency by adopting a precondition procedure, an option that has been explored for OFDFT by Hung et al.⁴⁹⁶

We note that L-BFGS stands out for its stability, with several available options for updating the search direction, \mathbf{p}_k . Because, however, it requires the storage of prior $\psi_k(\mathbf{r})$ and gradients, its memory cost is larger than that of any of the other methods mentioned.

For step (2), the step size, λ_k could be determined by direct approximation⁴⁸⁵ (as in the SD method) or by line search.^{53,54,486} Determination by direct approximation, however, causes the stability of the optimization to be quite system dependent. In contrast, line search is numerically stable, so it is used rather generally in OFDFT codes. Line search stability arises from imposition of the so-called Wolfe conditions; see ref 497.

The most direct way to accomplish the third step, update of ψ_{k+1} , would be $\psi_{k+1} = \psi_k + \lambda_k \mathbf{p}_k$, followed by normalization of the total density to N_e . That procedure is not always stable. The problem has an elegant solution in the form of a scheme that makes the \mathbf{p}_k contribution orthogonal to ψ_k . Thus, the update becomes⁴⁸⁵

$$\psi_{k+1} = \psi_k \cos(\lambda_k) + \mathbf{p}_k \sin(\lambda_k) \quad (170)$$

which maintains the normalization of electron density at every step.

The Trust-Region Image Method (TRIM)⁴⁹⁸ is another elegant solution to the problem of energy minimization subject to the constraint $\int d\mathbf{r} n(\mathbf{r}) = N_e$. In short, TRIM exploits a grand canonical formulation of OFDFT, finding that the minimum problem with respect to $n(\mathbf{r})$ is actually a saddle point minimax optimization with respect to n and the chemical potential, μ . The energy functional is found to be convex with respect to n and concave with respect to μ . Ryley et al.⁴⁹⁸ derive expressions for the gradient and the Hessian with respect to variation of the chemical potential and the electron density (which they parametrize in terms of Gaussian-type orbitals). That provides a density optimization method that is robust and requires fewer iteration to convergence than the method of Chan et al.¹¹⁹ Unfortunately, no comparisons to the method of Jiang and Yang⁴⁸⁵ have been reported. Therefore, it is difficult to assess the relative robustness of TRIM compared to the density optimization methods exploiting eq 170 as commonly implemented in pseudopotential codes such as PROFESS and DFTpy.^{54,486}

3.3. Basis Sets

As just surveyed, whether the energy minimization is direct or indirect, it is always necessary to express functions in terms of a basis. Though any complete set would do in principle, in practice the choice of basis is critical to determining the range

of applicability of the software. Basis sets employed currently include plane waves (PWs),^{481,482,499} atomic orbitals (AOs) (e.g., Gaussian-type-orbitals (GTOs)⁵⁰⁰), finite differences (FDs),^{501,502} finite elements (FEs),⁵⁰¹ and wavelets.^{503–505} The majority of OFDFT software to date is targeted at materials problems (hence, periodic boundary conditions); thus, a discretization of space well-adapted to that periodicity is used. Therefore, in this section we discuss the PW, FE, and FD approaches and refer the reader to the references above regarding the other ones.

3.3.1. Plane Wave Basis. Plane waves (PWs) are the most widely used basis functions in electronic structure calculations for materials science.^{481,482,499,506} Generally, PWs confer the following advantages:

1. Independence from nuclear (ionic) positions avoids Pulay forces⁵⁰⁷ and provides a uniform resolution description in all space;
2. Calculation of derivatives of $n(\mathbf{r})$ requires only algebraic operations in reciprocal space. [Remark: The derivative of any real-space function, $f(\mathbf{r})$, is straightforward in reciprocal space, $\nabla f(\mathbf{r}) = \frac{1}{(2\pi)^3} \int d\mathbf{G} i\mathbf{G} \tilde{f}(\mathbf{G}) e^{i\mathbf{G}\cdot\mathbf{r}}$, where $\tilde{f}(\mathbf{G})$ is the Fourier transform of $f(\mathbf{r})$.] Depending on the pseudopotential used, derivatives of the total energy with respect to geometrical parameters (e.g., force and stress) are simple;
3. Imposition of periodic boundary conditions is straightforward;
4. Systematic improvement of the basis set is provided simply by setting the wave vector cutoff;

5. Quantities expressed in real space can be transformed efficiently to reciprocal space by use of Fast Fourier Transforms (FFTs), with a computational effort that is almost linear scaling, $O(N \ln N)$.

Even though recent progress on one-point KEDFs^{86,87} has resulted in functionals capable of describing condensed phase systems (recall Section 2), two-point functionals continue to dominate in materials science applications.¹⁴⁵ For them, the basis set employed must permit efficient KEDF evaluation to retain the quasi-linear scalability of OFDFT. PWs are well suited for that because the double-integration formula in eq 13 can be simplified greatly when a homogeneous system is considered. In that case, the kernel $\omega(\mathbf{r}, \mathbf{r}') \equiv \omega(|\mathbf{r} - \mathbf{r}'|)$ and the convolution theorem can be exploited to turn eq 13 into

$$T_s^{\text{nl}}[n] = T_{\text{TF}}[n] + T_{\nu W}[n] + \int d\mathbf{G} \tilde{n}^\alpha(\mathbf{G}) \tilde{\omega}[n](|\mathbf{G}|) \tilde{n}^\beta(-\mathbf{G}) \quad (171)$$

Because $n(\mathbf{r})$ is a real function, the integral can be simplified further by exploiting $\tilde{n}(-\mathbf{G}) = \tilde{n}(\mathbf{G})$. Overall, the $O(N \ln N)$ scaling is retained.

The quasi-linear scaling is maintained roughly even when more complicated nonlocal functionals are considered, such as those with density-dependent kernels, $\omega(n(\mathbf{r}), |\mathbf{r} - \mathbf{r}'|)$ as in HC,⁷⁷ LMGP⁸⁸ and LDAK,⁹⁰ or $\omega(\xi(\mathbf{r}, \mathbf{r}')|\mathbf{r} - \mathbf{r}'|)$, as in WGC²⁹⁷ and XWM.⁴⁹ Recall the discussion in Subsection 2.4.2. At first glance, the computational cost for such functionals would seem to scale quadratically with the number of grid points. In Subsection 3.4, we discuss the details of the evaluation in a PW basis set for functionals of this type such that linear scaling with system size is maintained with a moderate prefactor.

PW basis sets are not without drawbacks. Probably the most significant is the one we have discussed already, the necessity

of pseudopotentials.^{481,482,499,506,508} Recall Subsection 3.1. That removes all-electron calculations from the picture. (There is KSDFT software for the all-electron linearized augmented plane wave basis (LAPW).^{402,509–512} However, that basis is atom-centered and muffin-tin-based and, hence, quite ill-suited to AIMD.) An excellent compromise for the PW basis is provided by the PAW method.^{120,399} In conventional KSDFT calculations, PAWs seem largely to have supplanted NLPPs. As we noted in Subsection 2.7.4, the PAW ansatz needs substantial further exploration in connection with OFDFT.

Another drawback of the PW basis is the inherently three-dimensional periodicity in the computation of the classical electrostatic potential. This is introduced by use of the periodic Coulomb kernel, $f_h(G) = \frac{4\pi}{G^2}$ for reciprocal space vector \mathbf{G} , yielding a Hartree potential in reciprocal space

$$\nu_H(\mathbf{G}) = \frac{4\pi}{G^2} n(\mathbf{G}) \quad (172)$$

Unfortunately, for intrinsic system periodicity lower than three dimensions, the common algorithm is not ideal.⁵¹³ Corrections, however, exist, both at the level of the energy functional⁵¹⁴ and for the electrostatic potential.^{515–517} Those typically only add a small prefactor to the simulations. Alternatively, real-space Poisson solvers can be employed.^{501,502,517} Note that this problem occurs in both OFDFT and conventional KSDFT.

A potential drawback for massively parallel implementations is the need for “all-to-all” communication among the processors needed by the FFT.^{502,518} Though this problem cannot be circumvented entirely, currently available parallelization schemes are robust and have been shown to maintain linear scalability of the algorithm.^{519,520} We discuss this aspect further in Subsection 3.5.2. Again, we point out that parallel limitations for FFTs are not specific to OFDFT. Conventional PW-basis KSDFT codes have the same problem.

3.3.2. Real-Space Representations. Among real-space basis sets, perhaps the most direct and easy approach is FD.⁵²¹ In it, the values of both $n(\mathbf{r})$ and the potentials are given on a discrete grid of real-space points.⁵²² Accordingly, the Hamiltonian also must be discretized. Thus, the key aspects of the FD method are the discretization of the Laplacian operator for the evaluation of the von Weizsäcker kinetic potential and solution of the Poisson equation for the electrostatic potential. Denoting the grid point positions as (x_i, y_j, z_k) , the expansion of the Laplacian operator acting on ψ for coordinate x is

$$\frac{\partial^2 \psi(\mathbf{r})}{\partial x^2} = \sum_{n=-N_f}^{N_f} C_n \psi(x_i + nh, y_j, z_k) + O(h^{2N+2}) \quad (173)$$

where h and N_f are the grid spacing and order of the FD expansion, respectively, and C_n are Taylor expansion coefficients. Algorithmic details and results are found in refs 501 and 502.

The truncation error $O(h^{2N+2})$ introduced by the Taylor expansion can be controlled by tuning the expansion order and grid spacing. For low-order FD expansions, sparse linear algebra leads to further computational efficiency. Note that when $\psi(\mathbf{r})$ is used to evaluate physical quantities such as the Hartree energy, the error originating from the series truncation can be of either sign and cannot be controlled systematically.

Thus, the FD approximation may lead to nonpositive definite Hamiltonian approximations that could result in variational breakdown.⁵⁰¹

For a non-Cartesian grid, an additional important detail is that a direct discretization of the Laplacian operator contains $O(N_f^2)$ terms in the summation because of the need to include off-diagonal neighboring points in a square of size $(2N_f + 1)^2$ around any given point. That would impose a high computational scaling prefactor. To solve the problem, Tiago et al.⁵²³ presented a generalized high-order FD method that bypasses the evaluation of mixed-derivative terms by employing a weighted combination of derivatives along the three original directions and three additional nearest-neighboring directions. That reduces computational cost significantly. The approach has been adopted in the ATLAS code.⁵³ To improve FD computational efficiency further, it is possible to define grids of different resolution depending on the spatial region.^{502,524} Though double-grid and multigrid methods^{501,525} have been deployed in ordinary KSDFT solvers, their usefulness in OFDFT implementations has yet to be assessed.

FD provides some advantages over PW for the evaluation of semilocal functionals. From Figure 11 it is clear that for large

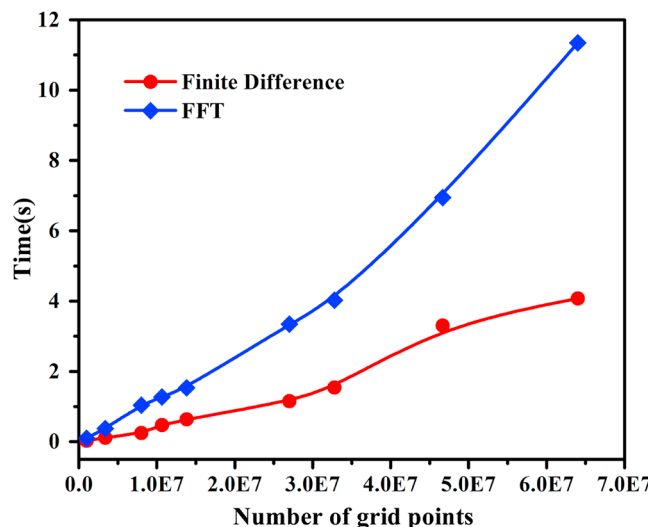


Figure 11. Timings for the computation of the von Weizsäcker potential in different numbers of grid points. The FD basis set implemented with efficient computation of the Laplacian outperforms the PW basis set formulation implemented with FFTs. Reprinted with permission from ref 53. Copyright (2016) Elsevier.

systems FD outperforms PW for the von Weizsäcker functional. FD, however, is not well-suited to evaluate nonlocal functionals. Thus, typically PW and FFTs are favored for them. However, real-space approaches to evaluate nonlocal functionals exist (*vide infra*).

Lastly, because FD methods represent the electron density in real space, they are well-suited to aid AIMD accelerators. This was described in detail in ref 526, where a simple FD method was employed to track the electron density and its associated potential with free-space (open) boundary conditions in adaptively restrained particle simulations (ARPSs). An ARPS decomposes the system into slow- and fast-moving particles. An ARPS⁵²⁷ accelerates AIMD by freezing some of the slow-moving particles along a molecular dynamics

trajectory. Coupling ARPSs with an FD solver for OF-DFT provides the possibility to compute good initial guesses for the electron densities and their associated Hartree potential during the AIMD easily. That is, because of the real-space representation of the density, the frozen ions remain fixed also during the OF-DFT density optimization, thereby achieving a computational saving. Even though such schemes could be implemented in a PW basis (as PW basis methods have access to a real-space representation of the electron densities on the FFT grid), to our knowledge such PW counterparts have not appeared so far in the literature.

Moving on to the finite element method (FE), generically it uses sets of strictly local piecewise polynomials as basis functions. Those functions only overlap with immediate neighbors.^{528–530} Various choices for those basis functions involve their dimensionality, polynomial order, and continuities (preservation of value and derivatives across interelement boundaries).^{528,529,531} The two main steps needed to construct a FE basis set are

1. Partitioning of the domain Ω into subdomains (elements).
2. Definition of the local basis function within each element and piecing them together to obtain the final piecewise polynomial basis $\{\phi_j\}$. Generally, any function ψ defined in the domain Ω can be expanded with

$$\psi(\mathbf{r}) = \sum_j u_j \phi_j(\mathbf{r}) \quad (174)$$

In step 1, the elements are chosen to optimize computational efficiency. For example, triangles and quadrilaterals commonly are used as two-dimensional elements, while tetrahedra, parallelopipeds, and wedges are common in three-dimensional systems.^{528,529,531}

In step 2, interpolation nodes are set in each element for the selected piecewise polynomial functions. The basis functions typically are chosen such that ϕ_j is 1 on the j -th FE interpolation node while it is 0 on all the other nodes. Typically only function continuity is imposed. First- and higher-order derivative continuity typically is not. Thus, differential equations for solution of the electronic structure problem need to be reformulated into integral equations. Once this is accomplished, such problems as the real-space Laplacian, Poisson equation, Poisson–Boltzmann equation, and eigenvalue equation can be transformed to a corresponding sparse matrix problem, which can be solved in various efficient and scalable approaches. Details of these techniques can be found in refs 530 and 532.

A general drawback of FE methods is that typically they require more memory than FD-based approaches and are more difficult to implement as well. Until now, they have proven advantageous primarily for very large system sizes.

A specific issue arises in the context of density-independent two-point KEDFs. Equation 171 shows how the PW basis and the convolution theorem combine to form an elegant reduction of the 6-dimensional integrals. In real-space basis set schemes an alternative is needed. It has two requirements, handling of the long-range Coulomb interaction and of nonlocal functionals (two-point KEDF and, sometimes, XC). Choley and Kaxiras⁵⁰ proposed a method with scaling close to $O(N)$. They noted that the kernel of a density-independent two-point KEDF can be fitted with reasonable accuracy by

rational functions. Further, its Fourier transform can be approximated as a sum of subkernels

$$\omega(g) = \sum_{j=1}^m \frac{P_j g^2}{g^2 + Q_j} \quad (175)$$

where P_j and Q_j are complex numbers determined by fit to the original Fourier transformed kernel. With this expression, the convolution of a function $f(\mathbf{r})$ with a kernel $\omega(|\mathbf{r} - \mathbf{r}'|)$

$$v(\mathbf{r}) = \int d\mathbf{r}' f(\mathbf{r}') \omega(|\mathbf{r} - \mathbf{r}'|) \quad (176)$$

becomes

$$v(\mathbf{G}) = \sum_{j=1}^m v_j(\mathbf{G}) \quad (177)$$

where

$$v_j(\mathbf{G}) = \frac{P_j g^2}{g^2 + Q_j} f(\mathbf{G}) \quad (178)$$

Then, the $v_i(\mathbf{r})$ can be obtained efficiently by solving the corresponding Helmholtz equation.

$$\left[-\frac{1}{(2k_F)^2} \nabla^2 + Q_j \right] v_j(\mathbf{r}) = -\frac{P_j}{(2k_F)^2} \nabla^2 f(\mathbf{r}) \quad (179)$$

Benchmarks for the WGC²⁹⁷ functional (which, by Taylor expansion of the kernel around a reference constant density, requires the evaluation of integrals involving only density-independent kernels; see below) indicated that the error introduced by the approximation is negligible. The scheme also was adopted to simulate tens of thousands of atoms with full atomic resolution and millions of atoms using coarse-graining techniques.⁵³³

3.4. Computationally Efficient Evaluation of Nonlocal Functionals

For nonlocal KEDFs with density-dependent kernels, the convolution integral no longer is directly applicable because such functionals involve integrals (in both the energy and the potential) of the type

$$P(\mathbf{r}) = \int d\mathbf{r}' f(\mathbf{r}') \mathcal{K}(k_F(\mathbf{r}), \mathbf{r} - \mathbf{r}') \quad (180)$$

$$Q(\mathbf{r}) = \int d\mathbf{r}' f(\mathbf{r}') \mathcal{K}(k_F(\mathbf{r}'), \mathbf{r} - \mathbf{r}') \quad (181)$$

As just noted, the WGC and XWM functionals^{49,297} bypass that complication by use of a Taylor expansion of the kernel (i.e., the \mathcal{K} function in eqs 180 and 181) around a constant density, n_0 (or equivalently around a constant Fermi wavevector, $k_F^0 = (3\pi^2 n_0)^{1/3}$). The remaining integrals are evaluable by simple convolutions. While the technique reduces computational complexity, it jeopardizes transferability. Specifically, the WGC and XWM KEDFs are applicable only to bulk systems for which a reference density is well-defined and in which $n(\mathbf{r})$ does not exhibit large variations. Without a system-dependent prescription for the reference density, WGC and XWM are inapplicable to finite systems. In some instances they also are affected by convergence problems.^{49,79}

The LMGP KEDF provides a way around this difficulty by defining the potential with an integral equivalent to eq 180. It can be evaluated by a spline technique based upon use of a

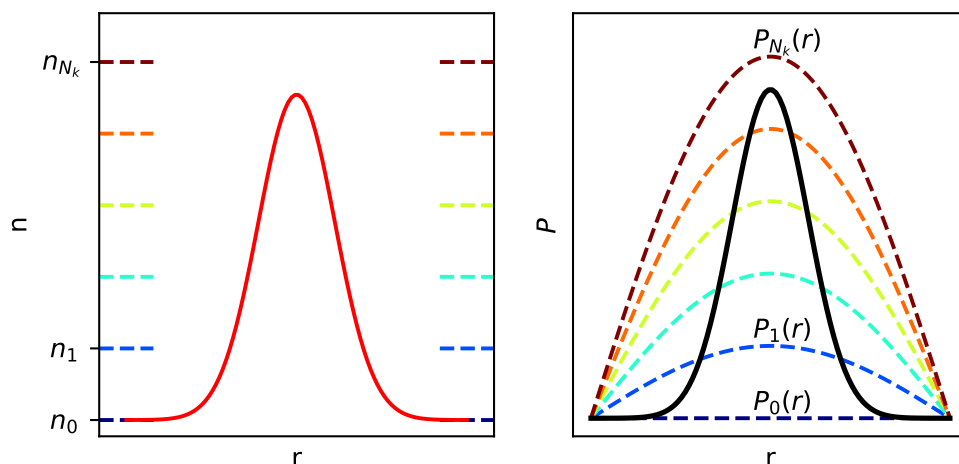


Figure 12. Depiction of the procedure to compute the integral, $P(r)$, in eq 180. On the left, the electron density is sampled by N_k values, $\{n_k\}$. On the right, the function $P(r)$ is approximated at every point by splining the values for each of the $P_k(r)$ functions.

discrete set of reference densities. For N_k selected density values $\{n_k\}$, there is a corresponding set of Fermi wavevectors, $\{k_F^k\}$. It follows⁵³⁴ that

1. For each k_F^k value, the integral eq 180 can be evaluated by convolutions to generate N_k functions, $P_k(r) = P(k_F^k, r) = \int d\mathbf{r}' f(\mathbf{r}') \mathcal{K}(k_F^k, \mathbf{r} - \mathbf{r}')$.

2. The final objective function, $P(r)$ then can be recovered by spline representation of the $P_k(r)$ functions for each \mathbf{r} point. That is, for each \mathbf{r} , the spline identifies k_F^k points near $k_F(r)$, $\{..., P_{k-1}(\mathbf{r}), P_k(\mathbf{r}), P_{k+1}(\mathbf{r}), ...\}$ and splines them to find the best estimate for $P(r) = P(k_F(r), \mathbf{r})$.

This spline procedure is depicted in Figure 12. For it, a linear quadrature was chosen for $\{n_k\}$ (i.e., $n_k = k \times \Delta$, with Δ a positive real number). In practice, it often is advantageous to employ a logarithmic quadrature (i.e., $n_k = \alpha n_{k-1}$ with α a positive real number). The motive is to improve accuracy and aid convergence for nonperiodic systems (e.g., clusters and surfaces). In them, the electron density has large variations, from a maximum to arbitrarily close to zero, in contrast to bulk systems. For nonperiodic systems, large N_k values are reported⁴⁶² to be necessary, so judicious choice of grid becomes critical to computational performance.

The HC and LDA-K KEDFs define an energy functional of the kind shown in eq 47. The functional derivative (potential) involves integrals of both types, eq 180 and eq 181. For HC and LDA-K, evaluation of the integral eq 181 therefore requires additional steps that typically are done in reciprocal space. For a periodic system with cell volume V_{cell} the Fourier transform of $Q(\mathbf{r})$ is⁵³⁵

$$\tilde{Q}(\mathbf{G}) = \frac{1}{V_{cell}} \int d\mathbf{r}' f(\mathbf{r}') e^{-i\mathbf{G} \cdot \mathbf{r}'} \tilde{\mathcal{K}}(k_F(\mathbf{r}'), \mathbf{G}) \quad (182)$$

thus

$$\tilde{\mathcal{K}}(k_F(\mathbf{r}'), \mathbf{G}) = \int d\mathbf{r} \mathcal{K}(k_F(\mathbf{r}'), \mathbf{r} - \mathbf{r}') e^{-i\mathbf{G} \cdot (\mathbf{r} - \mathbf{r}')} \quad (183)$$

The resulting set of functions, $\{\tilde{\mathcal{K}}(k_F^k, \mathbf{G})\}$, then is used to approximate the integral in eq 182 by Riemann summation. Inverse Fourier transform to real space then recovers $Q(\mathbf{r})$ of eq 181.

It is worth mentioning that it is more computationally demanding to evaluate eq 181 than eq 180 because all N_k

points need to be considered for all \mathbf{r} values in eq 181 compared to the spline for eq 180. That typically requires only the k_F^k points in the neighborhood of the $k_F(\mathbf{r})$ of interest. In conclusion, the computational evaluation of both integrals scales quasi-linearly because the FFTs involved scale like $O(N \log(N))$ and at most the algorithm requires N_k such FFT evaluations, independent of system size.

3.5. Algorithms for Million-Atom Simulations

Application of the increasingly accurate one- and two-point KEDFs we have described to simulation of extremely large systems (10^6 atoms or more) exposes some further algorithmic challenges. That also is true for exploitation of modern high-performance computing architectures. Issues of interest relate to the cost of evaluating the ion–ion and electron–ion interaction contributions and, hence, the structure factor. Related to that is solution of the Poisson problem (needed for the electrostatic potential). In what follows we summarize advances regarding both as embodied in contemporary OFDFT software. We also describe common strategies for distributing data and work in the most computationally intensive steps of the OFDFT simulation.

3.5.1. Structure Factors. Use of FFTs (and splining) gives $O(N \ln N)$ computational cost scaling for evaluating two-point KEDFs. But the electrostatic interaction terms (Hartree, ion–ion, and electron–ion) involve a computational cost that at first sight is quadratic scaling. Though the separate electrostatic sums are divergent in periodic systems,^{536–540} the divergences cancel for neutral cells.⁵⁴¹ The Hartree term can be evaluated by using FFT to achieve $O(N \ln N)$ scaling; see eq 172. But if the ion–ion interaction is evaluated by standard Ewald summation,⁵⁴¹ the quadratic computational cost with the number of atoms in the simulation cell M , $O(M^2)$, becomes the bottleneck at system sizes of a few thousand atoms. The problematic term is the ionic structure factor. To simplify, here we consider only the ion–electron interaction E_{ext} but a similar treatment applies to the ion–ion interaction. Also for simplicity we give the argument for the bare Coulomb interaction. Use of pseudopotentials changes only the details, not the essence.

Equation 16 can be re-expressed as

$$E_{ext}[n] = - \int d\mathbf{r} d\mathbf{r}' n(\mathbf{r}) \frac{n_a(\mathbf{r}')}{|\mathbf{r} - \mathbf{r}'|} \quad (184)$$

where

$$n_a(\mathbf{r}) = \sum_{\alpha}^{N_a} Z_{\alpha} \sum_{k}^{n_{\alpha}} \delta(\mathbf{r} - \mathbf{R}_k) := \sum_{\alpha}^{N_a} Z_{\alpha} S^{\alpha}(\mathbf{r}) \quad (185)$$

Here N_a is the number of atomic species (or atom types), \mathbf{R}_k is the position of the k -th atom of type α , n_{α} is the number of atoms of type α , and Z_{α} is their ionic charge. $S^{\alpha}(\mathbf{r})$ is the structure factor. Fourier transformation and application of the convolution theorem then gives

$$E_{ext}[n] = \sum_{\alpha}^{N_a} Z_{\alpha} \int d\mathbf{G} \tilde{n}(\mathbf{G}) \tilde{S}^{\alpha}(\mathbf{G}) \frac{4\pi}{|\mathbf{G}|^2} \quad (186)$$

$$\tilde{S}^{\alpha}(\mathbf{G}) = \sum_{k}^{n_{\alpha}} \exp(i\mathbf{G} \cdot \mathbf{R}_k) \quad (187)$$

$\tilde{S}^{\alpha}(\mathbf{G})$ is simply the Fourier transform of $S^{\alpha}(\mathbf{r})$ in eq 185.

As there are N_g grid points for \mathbf{G} , the direct evaluation of eq 187 scales as $O(n_{\alpha} \cdot N_g)$.^{53,486,542} This is effectively quadratic scaling with system size. For system size $n_{\alpha} \approx 1,000$, the structure factor calculation begins to dominate the computational time relative to other terms in the OFDFT energy functional. That degrades the total computational cost to quadratic scaling.^{53,486,542,543} This crossover seldom is encountered in conventional KSDFT simulations, as their inherent $O(N^3)$ scaling dominates, with the result that system sizes greater than a few thousand atoms rarely are treated by conventional KSDFT.

To address the problem, Darden et al. proposed the particle-mesh Ewald (PME) method.⁵⁴⁴ In it, Lagrangian interpolation is used to evaluate the reciprocal space Ewald sums and the resulting convolution is evaluated via FFT. The approach reduces the cost to $O(N \ln N)$. The Lagrangian interpolation is effectively an approximation of the real-space structure factor in which the Dirac delta functions in eq 185 are approximated by spline basis functions with compact support (i.e., they decay exactly to zero within a finite number of grid points). Cardinal B-splines typically are chosen for this.⁵⁴⁵ The compact support is important, as it assures the locality of the functions around the ions that is the cause of the linear scalability of the method. Typically only a handful of grid points in each direction from the ion is needed. (Note that the ion position may not coincide with a grid point.)

Choley et al. apparently were the first to use the B-spline plus FFT convolution PME scheme for evaluating the structure factor in electronic structure calculations. The first implementation in OFDFT software was by Hung et al.⁵⁴²

It also is possible to avoid the computation of the structure factor altogether by representing the atomic local pseudopotential (LPP) through its source pseudocharge density. The pseudocharge is localized and can be represented fully on a small subset of grid points (a crucial property also shared by the cardinal B-spline function). After summing all the pseudocharges, followed by multiplication by the Coulomb kernel ($\frac{4\pi}{q^2}$), a single FFT operation recovers the full LPP. The scheme has been implemented in the ATLAS software package and benchmarked on large-scale simulations.⁵⁴³

3.5.2. Parallelizing FFT for Large Systems: Small Box FFT and 2D-FFT. It should be abundantly clear by now that an important component of the computational cost in OFDFT

with two-point KEDFs is the ubiquitous use of FFTs. Though they provide favorable $O(N \ln N)$ scaling (N total number of grid points), exploitation of that fact is hampered by the need for all-to-all communications among the CPUs involved. Jiang et al.⁵⁴⁶ proposed an efficient “small-box” FFT (SBFFT) algorithm to address this problem for the Hartree term and, in fact, for any convolution integral with a radially decaying kernel. The kernel is presumed to be decomposable into long- and short-ranged parts. As depicted in Figure 13 (in cubic

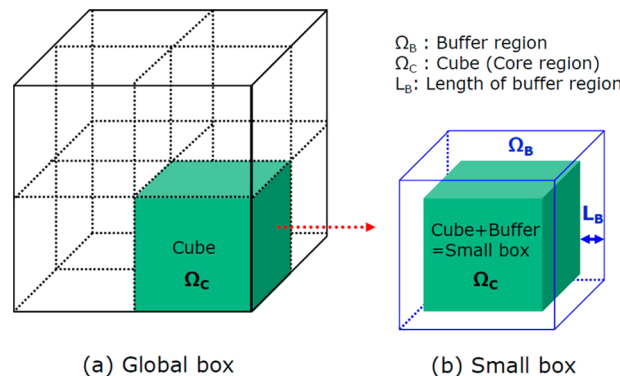


Figure 13. Small-box FFT algorithm distributes work and data for the computation of Coulomb and KEDF terms in the OFDFT energy functional. The large simulation cell is divided into small boxes. Each is composed of a physical small box and a buffer region which is introduced for evaluating the derivative of the density function needed for the evaluation of KEDFs. Reprinted with permission from ref 519. Copyright (2016) American Chemical Society.

symmetry for simplicity), SBFFT is a divide-and-conquer strategy. The simulation cell volume is subdivided into small boxes. Each small box has an internal structure consisting of a cube decorated by a buffer region. In the small box, the Poisson equation for the short-range kernel is solved on a dense FFT grid. Then a long-range (but smooth) kernel is used in a Poisson equation solver over the entire simulation cell on a coarse FFT grid. The Hartree potential thus is the sum of localized potentials over the small boxes and a long-range potential represented on a coarse grid which provides efficient evaluation. SBFFT yielded a speed-up of 8.6 for a 10-fold increase in the number of CPUs, which is quite good for a parallel algorithm for massive systems. In addition, the relative error was estimated to be 4×10^{-5} (or 0.1 meV maximum error in the potential) for a massive simulation using 24,000 processors with a large FFT grid of $2,400^3$ points.⁵⁴⁶

Following the same strategy, SBFFT can be used to evaluate two-point KEDFs by introducing a real-space truncation of the nonlocal KEDF kernel. Chen et al.⁵¹⁹ performed an optimization of 1,024,000 Li atoms by distributing the work on 65,536 processes. That is a striking improvement over a “vanilla” FFT parallelization.

Two additional methods for implementing parallel FFTs, slab and pencil decompositions,^{547,548} have been introduced recently to OFDFT by Shao et al.⁵²⁰ in the ATLAS package.

In the slab decomposition, slices of the simulation cell along one axis are distributed evenly among all processes, while in the pencil decomposition the spatial partitions are done along two axes. As a result, the theoretical limit on the number of processors is larger in the pencil decomposition. The limits can be estimated from the mesh grid. With an $N = N_x \times N_y \times N_z$ point grid, the maximum number of processes is $\max(N_x, N_y)$

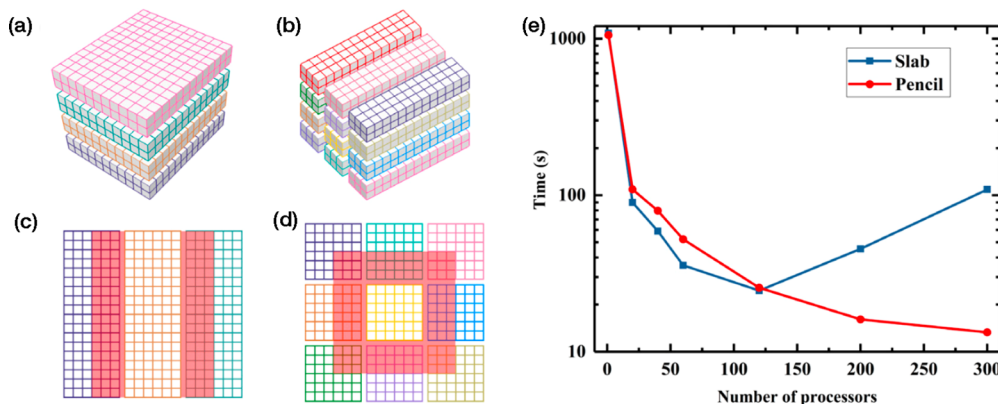


Figure 14. 2D-FFT algorithm. (a) Slab domain decomposition using four processors. (b) Pencil domain decomposition using a 4×3 processor grid. The buffer region (in red) is used to store information for the points near a domain boundary for the center processor in slab (c) and pencil (d) decompositions. (e) Comparison of wall time for a density optimization of 32,000 atoms of bulk Al with slab and pencil domain decompositions. Reprinted with permission from ref 520. Copyright (2018) Elsevier.

N_z) and $N/\min(N_x, N_y, N_z)$ for the slab and pencil decompositions, respectively. As illustrated in Figure 14(c) and (d), an additional buffer region is needed to store information for the points near a domain boundary. That region is used to perform finite difference schemes for each processor and can reduce the massive communication frequency.

In the form implemented in the ATLAS code, these approaches were benchmarked by comparison of total (wall clock) time for density optimization of Al with a 32,000 atom supercell. Both slab and pencil domain decompositions were used. Those results also are shown in Figure 14(e). Below 200 processors, the performance of the two approaches is comparable. Above that, the pencil outperforms the slab decomposition significantly. Shao et al.⁵²⁰ showed that the parallel efficiency of the pencil decomposition reached 0.92 for a 2,048,000 atom simulation distributed on up to 4,096 processors.⁵²⁰ Another impressive achievement by those authors is the recent successful optimization of the electron density of cubic aluminum using the WT KEDF and LDA XC functional for a sample of 100,271,632 atoms.⁵⁴⁹

3.5.3. Real-Space Algorithms for OFDFT Simulations of Million-Atom Systems. The FE and FD real-space representations introduced in Subsection 3.3.2 are generally efficient and scalable. However, when multimillion-atom systems are considered, real-space methods have the opportunity to use a lower spatial resolution for the space far away from notable features (e.g., vacancies in bulk systems and clusters in vacuum) and reduce computational complexity thereby.^{518,530,550} A realization of this idea has been implemented by Gavini et al., who proposed the quasi-continuum OFDFT scheme.^{533,550} In it, three feature-centered grids of different resolutions are adopted. We refer the interested reader to refs 518, 530, and 550 for additional information about the quasi-continuum approach as well as other FE methods. We also refer the reader to additional references describing other scalable real-space methods that have found success in DFT codes, namely refs 11 and 551–557.

3.6. Software for OFDFT Simulations of Materials

As discussed in Section 1, the TF and TFD approximations applied to atoms yield one-dimensional differential equations that have analytical solutions. They attracted interest in no

small measure because of the extremely limited computational capacity of that era. To our knowledge, the first computational solution to the TFD equations was reported in 1950 by Reitz.²⁸ Other numerical solutions followed.^{37,38,116,558–561} Given limitations of both the KEDFs and the available computational resources back then, those efforts targeted small systems, e.g. single atoms or small molecules.

Beginning in the 1980s, OFDFT software became more general and capable of treating wider system classes, including condensed phases,^{43,44,451} surfaces,⁵⁶² and metal clusters.⁵⁶³ It seems fair to say, however, that those calculations were largely proof-of-principle efforts with limited impact outside the OFDFT method development community.

With the development of more accurate KEDFs and, concurrently, the betterment of pseudopotentials, since 2007 several groups have developed general-purpose codes, aimed at large-scale (up to hundreds of millions of atoms⁵⁴⁹) OFDFT simulations.^{53–55,162,486,520,533,564} A summary of available and recently reported OFDFT software follows.

1. PROFESS:

The PRinceton Orbital-Free Electronic Structure Software (PROFESS)^{55,486,564} package is an open source code released under a CPC license.⁵⁶⁵ It appears to be the most comprehensive OFDFT code currently available, as it can perform a wide range of simulations under diverse conditions. It employs a PW basis set; hence, Coulomb interactions, gradient, and Laplacian operations are evaluated in reciprocal space by FFT. The electron density is found by direct minimization of the total energy functional. Several minimization algorithms are available (SD, CG, TN, BFGS, and more). Besides electron density optimization, with use of forces and stresses, PROFESS can run molecular dynamics and structure relaxations with most GGA and two-point KEDFs. In terms of pseudopotentials, PROFESS not only supports LPP but also supports recent experimental schemes that go beyond LPP^{161,169} (see Subsection 2.7). Recently, libKEDF, a library for the efficient and accelerated evaluation of local, semilocal, and nonlocal functionals, was released.⁵⁶⁶ The library will be integrated in PROFESS. PROFESS also has been coupled to Quantum Espresso and augmented with finite-temperature KEDFs and XC DFAs to provide a

general platform for even-handed comparison of OFDFT-based and conventional KSDFT-based AIMD calculations.¹⁶²

2. ATLAS:

Ab-initio orbiTaL-free density functionAl theory Software (ATLAS) is a FD implementation (both the Laplacian operator and the Poisson equation) of OFDFT. PW basis sets and associated FFTs also are provided for the evaluation of two-point KEDFs and use as Poisson solvers. The primary electron density optimization algorithm is direct energy minimization. It can be used for both periodic and nonperiodic systems. For two-point functionals with density-dependent kernels, ATLAS uses OE-SCF instead.

The source code can be obtained from the ATLAS Web site (atlas-ch.cn⁵⁴⁹).

3. GPAW:

GPAW⁵¹ is released under the GNU Public License version 3 or any later version.⁵⁶⁷ In GPAW, there is an OFDFT implementation¹²⁰ consisting of a modified KSDFT solver adapted for tackling indirect solution of the OFDFT Euler equation via an SCF algorithm (recall Subsection 3.2.1). The OFDFT implementation in GPAW features local and semilocal KEDFs. Two-point KEDFs are yet to be implemented.

4. DFTpy:

DFTpy⁵⁴ is open source, released under an MIT license.⁵⁶⁸ It relies on Fourier space for the treatment of Coulomb interactions, for the computation of gradient and Laplacian operations, and for evaluation of two-point KEDF kernels. Various density optimization algorithms are provided, including OE-SCF.⁴⁶² DFTpy is a flexible, object-oriented OFDFT implementation with friendly APIs for Jupyter Notebooks. It thus is adapted to prototyping of new features. It can be used as a module for computing energy, forces, and stress. It also provides an API to the Atomic Simulation Environment^{569,570} for AIMD and other functionalities. In addition, it provides a hydrodynamic time-dependent OFDFT implementation.⁵⁴

5. CONUNDrum:

CONUNDrum⁵² is released under the GNU GPL v3 license.⁵⁶⁷ Both a high-order central FD method and FFT are used for calculation of various energy and potential terms. Density optimization is via direct energy minimization with SD and nonlinear CG algorithms. In addition to one- and two-point KEDFs, machine-learnt KEDFs also are available.

Besides the packages just listed, other software implementing real-space OFDFT algorithms has surfaced but apparently is not publicly available.^{484,533,571,572} There have been various materials simulations with these.^{573–576}

We conclude by remarking that OFDFT software currently is under development in several research groups around the world. The overall trend seems to favor material science applications and pseudopotential implementations. As multiscale software continues to improve and new multiscale schemes evolve (such as embedding^{577–579}), the role of OFDFT software will grow in importance. The future likely will see OFDFT solvers and conventional KSDFT solvers being used together in comparatively unconventional multiscale computational protocols. To achieve this goal, OFDFT

software seems to be following the trend of other materials science software, namely, transition from stand-alone codes to modules and libraries that can be used independently (nearly “black box”) by other software in flexible ways.

4. APPLICATIONS

In most cases, the prediction of structure–function relationships requires ab initio molecular dynamics (AIMD) simulations to represent configurational sampling and evaluate thermodynamic functions. Thus, the complexity of the requisite simulations can become intractable even for comparatively computationally efficient methods, such as semilocal conventional KSDFT. Doing AIMD with OFDFT is, in contrast, expected to be cost-effective for geometry optimizations and molecular dynamics of clusters, surfaces, and bulks. Aside from MD, OFDFT simply allows tackling of much larger systems than possible with conventional KSDFT, thereby helping to bridge the divide between few-atom system results and reality.

Note also that it is possible to accelerate OFDFT-based AIMD by applying techniques akin to what is done in conventional KSDFT.⁵⁸⁰ Even though acceleration methods such as the extended-Lagrangian Born–Oppenheimer molecular dynamics techniques have been proposed and benchmarked for model systems,⁵⁸¹ they have not yet been employed in OFDFT simulations. However, we expect they will be.

Here we give examples of what already has been achieved. In the subsections that follow, we review several applications of OFDFT that enabled modeling of systems that clearly would be outside the applicability range of conventional KSDFT. These include models of the mechanical properties of bulk metallic systems (including alloys), metallic clusters, and nanostructures. Even though OFDFT currently is limited by the systems it can approach (because of limitations of present-day approximate KEDFs and local pseudopotentials), the set of systems benefiting from OFDFT-based modeling is large and growing as approximate KEDFs become more refined, transferable, and accurate.

We review bulk solid studies first and then move on to nanostructures and clusters, liquids and matter under extreme conditions. Then we survey applications of those subsystem DFT methods that rely on KEDFs in their formulation.

4.1. Solids

Since the bronze age (3000 BC) human beings have understood that alloying metals modifies their bulk properties: mechanical, thermal, and optical. Even today, metallurgy is an active field of research and engineering. OFDFT has played a key role in this context, providing accurate models for alloys and defective metals.

Thematically in this review, we have noted that the $O(N^3)$ computational scaling of conventional KSDFT poses a severe incremental cost barrier regarding treatable system sizes. Yet system size matters. For example, to control finite-size effects, predictive models of alloys and defective metals require large system sizes (hundreds or even millions of atoms in the simulation cell). Radhakrishnan and Gavini⁵⁷⁶ found “remarkable cell-size effects that are present in computational domains consisting of up to 10^3 – 10^4 atoms, even in simple defects such as vacancies” either in bulk Al⁵⁷⁶ or in alloys.⁵⁷² Strong finite-size effects required Das et al.⁵⁷² to use a 879-atom cell to converge the OFDFT electronic structure of a monovacancy in

bulk Al. Additionally, Qiu et al.⁵⁸² reported that vacancy migration and formation energies display strong finite-size effects (fcc-Al needed supercells with up to 10,976 atoms). They employed the WT and WGC KEDFs and LDA and PBE XC functionals. Validation was carried out against conventional KSDFT for feasible system sizes (up to 256 atoms). While they found little difference between predictions with LDA and PBE XC functionals, the WT KEDF overestimated the vacancy formation energy by about 0.7 eV (out of an experimental value of 0.61 eV). WGC gave much improved vacancy formation energies (0.80–0.72 eV depending on the XC functional employed). Their study shows that practical implementations of OFDFT are predictive for studying vacancy formation, such as the collision cascade process in irradiated materials⁵⁸² or materials under loading shock conditions.⁵⁸³

Zhuang et al.¹⁷¹ employed the WT KEDF and PBE XC functional implemented in PROFESS to investigate the elastic and thermodynamic properties of complex Mg–Al intermetallic compounds. First they benchmarked the performance of OFDFT as compared with both conventional KSDFT and experimental results and found good agreement. For example, the lattice constant of Mg₁₃Al₁₄ is 10.314, 10.437, and 10.183 Å from experiment, OFDFT, and conventional KSDFT, respectively. They also found similarly good agreement for phonon spectra of hcp Mg and fcc Al. In a second step, they evaluated the dynamical stability of alloys by computing phonon spectra on 3 × 3 × 3 supercells for all the compounds considered, including Mg₂₃Al₃₀. This resulted in a 1431-atom system. Their computations enabled the determination of thermodynamic properties, such as the temperature-dependent formation energy, heat capacity, and thermal expansion coefficient. The simulation yielded linear thermal expansion coefficients of 22.8 and 22.6 × 10⁻⁶ K⁻¹ at $T = 298$ K for Mg and Al, respectively. Those results compare well with the experimental values, 24.8 and 23.1 × 10⁻⁶ K⁻¹, an indication that OFDFT is quantitatively reliable for computing properties of Mg–Al intermetallic compounds.

A detailed account regarding various kinds of defects in metals and metal alloys (such as vacancies and vacancy clusters, interstitials, dislocations, and grain boundaries) can be found in the recent review by Witt et al., ref 145. It is worth highlighting a few specific examples to showcase the usefulness of OFDFT to materials science. For one, Xia et al. employed the WGC-decomposition approach (see Section 2.7.1) and HC KEDFs, the PBE XC functional, and BLPS pseudopotentials implemented in PROFESS to study several Li–Si alloys (LiSi, Li₁₂Si₇, Li₇Si₃, Li₁₃Si₄, Li₁₅Si₄, and Li₂₂Si₅). They found that over 80% of the OFDFT results had errors in the equilibrium lattice constants within 1% of the conventional KSDFT reference values, with the largest deviation less than 2%. The bulk moduli errors lie within 5 GPa (or approximately 10%) of the conventional KSDFT values.

Atomic-scale characterization and understanding of dislocations in bulk systems is another topic of importance for materials science. Because dislocation formation is difficult to monitor *in situ*, theory and modeling potentially are of very high value. Shin and Carter⁵⁸⁴ tackled the problem by employing the WT and WGC KEDFs, the LDA XC functional, and BLPS pseudopotentials with the PROFESS code. They benchmarked against conventional KSDFT with the same pseudopotentials and XC functional. The differences are quite small, within 25.8 (5.8) meV/atom for the total energy from

the WT (respectively WGC) KEDF, and 5.5 (5.2) GPa for elastic constants, and 16.3 (28.9) mJ m⁻² for stacking fault energies. The same authors also presented a thorough analysis of screw and edge dislocations in magnesium. They analyzed dislocation width and dynamics for the first time with an ab initio method which is much more accurate than the embedded-atom force fields⁵⁸⁵ typically employed in the engineering community. In another study, Hung and Carter⁵⁸⁶ analyzed the ductile properties of aluminum and showed that Al metal is much more resistant to plastic deformation than is predicted by commonplace engineering methods such as continuum models.

4.2. Metal Clusters, Quantum Dots, and Nanostructures

As mentioned, OFDFT-based AIMD (OFDFT-AIMD) can be quite useful for modeling metallic systems, including metal clusters. Multiple OFDFT-AIMD studies of metal clusters have been done dating to the early 1990s. We summarize a representative selection in what follows.

Despite its fundamental flaws (recall Section 2), historically the $T_{TF} + \lambda T_{vW}$ (TF λ vW) KEDF has been used in ways that encouraged further exploration of OFDFT treatment (with LDA XC) of clusters. Kanhere et al. treated simple Na and Mg dimers and trimers⁵⁸⁷ with the $\lambda = 1/9$ version. It also was used to study the most probable low-energy configurations of diverse clusters including the ground-state geometries and energetics of small Li clusters and the Li_nAl_m cluster,^{588–591} as well as melting of Al₁₃.^{592,593} Despite its deficiencies, the TF λ vW functional yielded bond lengths within 5% of the conventional KSDFT values,⁵⁸⁹ an encouraging indication of the promising nature of OFDFT-AIMD for study of non-periodic metallic systems. Thus, for example, Blaise et al.⁵⁹⁴ used OFDFT-AIMD with TF λ vW, $\lambda = 1/9$ to study the structural and dynamical properties of neutral and charged Na clusters up to sizes of several hundred atoms. Their ab initio calculations confirmed the validity of the so-called “liquid drop” model (often used in the nuclear physics community) as a function of the total cluster charge.⁵⁹⁵ Moreover, the Guo group⁵⁶² adopted OFDFT to study the equilibrium structure and bonding of small silicon clusters.⁵⁶³ They also used TF λ vW KEDF, but with $\lambda = 1/5$. Although the values of cohesive energy they obtained were only semiquantitative, the bond lengths and angles compared very well to values from conventional KSDFT.⁵⁶³ The OFDFT-AIMD approach was exploited by Aguado and coauthors as well in a number of studies of alkali clusters and alkali metal alloy clusters.^{126,127,453,596–599}

The first application of two-point KEDFs to clusters is credited to the Carter group.¹⁷⁴ They used the WGC²⁹⁷ two-point KEDF and LDA XC functional to study two-dimensional arrays of Al quantum dots. The issue investigated was the induction of a metal–insulator transition by increase of interdot separation. The only input to their model was a measure of the electron density decay as a function of intercluster separation. Their study was important because it showed that two-point functionals, even if developed for applications to bulk systems, could be applied to systems with remarkable inhomogeneity. (Note that the authors of ref 174 were careful to clarify that very low density values are probably unreliable because the Taylor expansion in WGC, mentioned in Subsection 2.4.2, may not be able to recover an appropriate kernel in low-density regions.)

Applications of OFDFT ventured to investigating tensile yielding of metal nanowires, behavior that is important for predicting how the metal behaves under extreme stress. In ref 173, Hung and Carter adopted the WGC KEDF, LDA XC functional, and BLPS pseudopotentials in the PROFESS code to model large-sized Al (111) nanowires (up to 20 nm in length). Interestingly, they found that such nanowires develop specific slip planes or amorphization of the crystal structure depending on wire size (see Figure 15). Employing the same

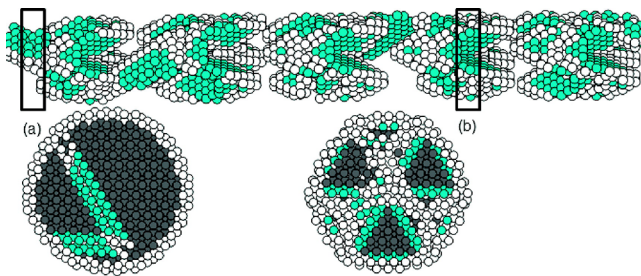


Figure 15. Aluminum wires under tensile stress. The wires break in specific ways (slip planes are made visible by the color coding) predicted by OFDFT with the two-point KEDF WGC. Green atoms correspond to hcp structure, gray to fcc structure, and white to an unknown structure. Reprinted with permission from ref 173. Copyright (2011) American Chemical Society.

approach, the Carter group went further and studied plastic properties of body-centered-cubic (bcc) Mg–Li alloys as potential lightweight metals. In the model, they used simulation cells containing up to 8640 atoms.⁴⁵⁴

Aguado et al. adopted a two-point functional developed in the group of Stott by simplifying the CAT KEDF³¹⁶ (similar in spirit to the WT functional¹⁴⁴) in conjunction with the LDA XC functional to study melting of various-sized sodium clusters.^{600–604} Quite interestingly, calorimetry experiment results⁶⁰⁵ were reproduced quantitatively by the simulations,⁶⁰¹ as shown in Figure 16. Detailed information is in the review by Aguado et al.^{606,607}

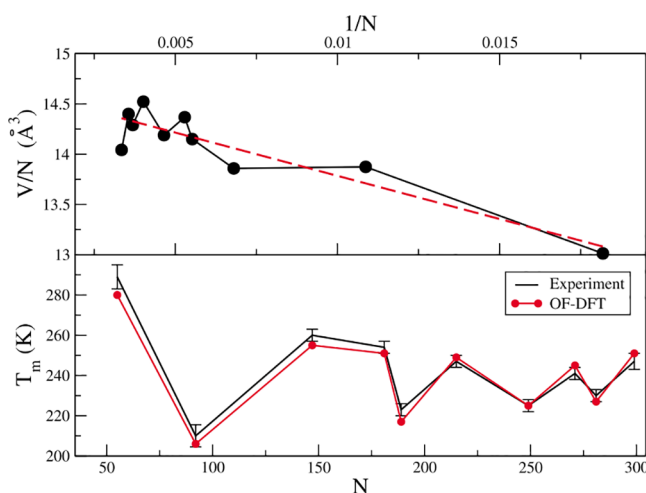


Figure 16. Size variation of the volume per atom (top) and melting temperature (bottom) in Na clusters from OFDFT simulations and experiment. The dashed line in the upper panel is the best linear fit to the data. Reprinted with permission from ref 601. Copyright (2005) American Physical Society.

Recall from Subsection 2.4.2 that the LMGP KEDF⁸⁸ recently developed is applicable to quite inhomogeneous systems, including clusters.

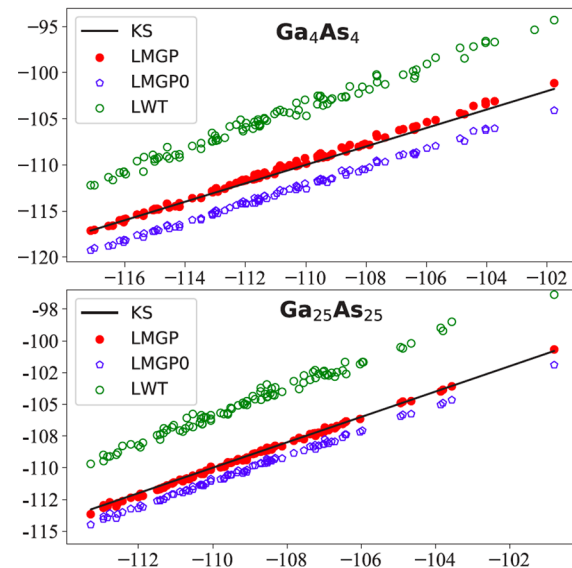


Figure 17. Energies (eV/atom) of 100 random structures of Ga_4As_4 and $\text{Ga}_{25}\text{As}_{25}$ compared to conventional KSDFT results (on the x axis) as well those from other KEDFs. LMGPO is LMGP with the corrective $1/q^2$ as the $q \rightarrow 0$ term removed. LWT is the result of using the integrand in eq 106 simply evaluated at $t = 1$; hence, it is LMGPO with the kernel terms arising from the functional integration removed. Adapted with permission from ref 88. Copyright (2019) American Physical Society.

Figure 17 and Table 5 show the results of benchmark studies for several hundred random-structured metal and semiconducting clusters. The LDA XC functional was adopted in all these OFDFT calculations, which were done with the ATLAS code. The outcomes indicate that two-point functionals with fully density-dependent kernels (such as LMGP) improve the performance of OFDFT considerably in modeling finite metal clusters and quantum dots compared with values from the semilocal functionals mentioned above. This is true for both total energy and electron density. The results also indicate that OFDFT can treat systems with highly inhomogeneous electron density, such as clusters and quantum dots, as accurately as conventional KSDFT at a fraction of the computational cost.

Because of the computational cost, until recently there have been no applications of two-point functionals with density-dependent kernels to nanoscale systems⁷⁹ such as materials interfaces and nanoparticles. That state of affairs changed with the introduction of the OE-SCF method (Section 3.2.1). It requires only ≈ 15 nonlocal KEDF evaluations for each geometry (depending weakly on system size and type). As an example, Figure 18 shows results for a nanoscale metal–semiconductor interface and a semiconductor nanoparticle, each containing hundreds of thousands of atoms. Both were studied with the LMGP KEDF.

We have mentioned already the avoidance of finite-size effects via OFDFT. For obvious computational cost reasons, it is rather common for conventional KSDFT calculations to be done on system sizes that are small compared to the scale of

Table 5. Mean Unsigned Error (MUE) for the Total Energy in eV/atom and, in Parentheses, Percentage Mean Unsigned Relative Error (MURE) for the Electron Density^a

Systems	LMGP	LMGP0	LWT	TF+15vW	WT
Mg ₈	0.18(3.79)	0.63(4.12)	1.19(4.05)	1.09(11.36)	8.79(16.0)
Si ₈	0.22(4.84)	2.17(4.90)	4.86(4.74)	1.46(8.28)	41.7(17.5)
Ga ₄ As ₄	0.34(5.40)	2.21(5.43)	6.15(4.89)	1.55(8.94)	51.8(19.3)
Mg ₅₀	0.05(3.31)	0.35(3.42)	0.84(2.38)	0.95(9.56)	3.23(10.3)
Si ₅₀	0.11(4.59)	0.95(4.65)	3.73(3.60)	1.53(7.24)	16.4(14.2)
Ga ₂₅ As ₂₅	0.13(5.21)	1.06(5.26)	4.29(3.19)	1.67(7.79)	22.7(16.8)
Mg ₈ ^S	0.28(5.20)	1.16(5.34)	2.66(5.29)	0.27(7.63)	19.4(18.6)
Mg ₈ ^{V S}	0.09(3.94)	1.67(4.10)	3.88(4.87)	0.10(5.60)	24.0(17.5)

^aMUE is defined as $\frac{100\%}{2N} \int |n_{\text{OFDFT}}(\mathbf{r}) - n_{\text{KSDFT}}(\mathbf{r})| d\mathbf{r}$ relative to conventional KSDFT values. Superscripts S and V S stand for “strained” and “very strained”, where the minimal interatomic distances between atom pairs are 1.2 and 0.8 Å, respectively. Both are much shorter than the equilibrium bond distance, 2.4 Å. Reproduced from Mi *et al.*⁸⁸.

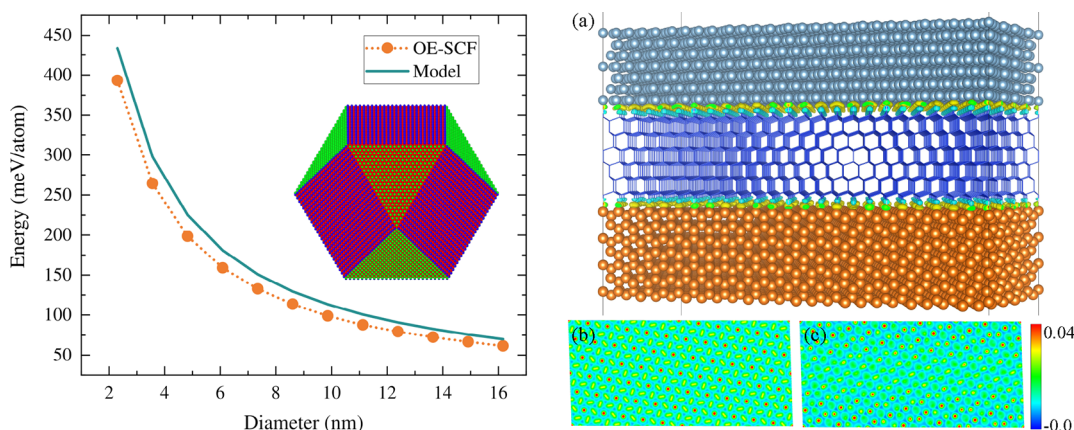


Figure 18. Left: Energy per atom of large icosahedral Si nanoparticles containing up to 102,501 atoms (dashed orange curve with dots is for the LMGP functional with OE-SCF solver; solid green curve is the empirical formula taken from ref 608). Right: Inset (a): depiction of a 3-layer interface, Al(111) (top), Si(111) (middle), and Mg(100) (bottom). Slabs of large dimensions were required in order to match the lattices of the materials at the interfaces. The charge flow across the Si–Al and Mg–Si interfaces (computed as the difference between the electron density of the 3-layer interface and the electron densities of the isolated slabs) is displayed with an isosurface volume plot visible in inset (a). Insets (b) and (c): Two-dimensional cut of the charge flow across the interface for the Si–Al and Mg–Si interfaces, respectively, which is seen to range between −0.01 and +0.04 electrons. Adapted with permission from ref 462. Copyright (2021) American Chemical Society.

experimental samples. That is particularly the case for modeling of nanoparticles and materials interfaces. For the latter systems, models often are unable to reproduce the correct crystal lattices at the interface because resolution of the differing lattice constants would require excessively large simulation cell sizes to reach a commensurate length. Reference 462 showed that with accurate KEDFs it is possible to overcome this difficulty and use appropriate system sizes. Shao *et al.*⁴⁶² in particular found that LMGP reproduced, quantitatively, the energy as a trend of nanoparticle radius (culminating in the largest nanoparticle depicted in the figure). For the metal–semiconductor interface in Figure 18 (right panels), the authors showed that the charge rearrangement across the interface reproduces known trends for passivated and unpassivated Si surfaces. (The charge rearrangement was computed by subtracting from the electron density of the interface the electron densities of the isolated slabs.)

4.3. Liquid Metals and Alloys

We turn next to the applications of OFDFT to liquid metals (including alloys) and their surfaces. We refer the reader to the reviews by Aguado *et al.* in ref 607 and by Witt *et al.* in ref 145 for additional and complementary details.

OFDFT-AIMD was used with success for studying the structure of simple liquid metals (such as Li, Na, Mg, Al, and

Cs) beginning in the 1990s.^{452,469,609,610} Typically that was with the LDA XC functional. That progress exploited the relevance of the HEG to such metals as embodied in use of the linear response formulation of two-point functionals beginning with WT⁴⁴ and continuing with SM,⁴⁶ WGC,²⁹⁷ and other KEDFs.^{43,299,320} Recall details in Section 2.4.2. In those simulations, first-principles LPPs (such as NPA and BLPS; see Section 3.1.2 for details), as well as empirical or model LPPs (such as the Topp-Hopfield LPP⁴⁴⁷ for Na and the Fiolhair model potential⁴⁴⁴ for Cs), were used. Note that empirical LPPs still are useful, as there exist parametrizations for those ions that are not approachable, so far, by first-principles LPPs, such as transition metals.⁶¹¹ We also note that the KEDFs mentioned are broadly useful for simulating complex liquids, including their surfaces.⁶¹²

An early success from Madden *et al.* was calculation of the dynamic structure factor of liquid Na⁶⁰⁹ using the LDA XC, their SM KEDF,⁴⁶ and the Topp-Hopfield LPPs.⁴⁴⁷ They obtained excellent agreement with experimental results for static and dynamic structure factors.⁶⁰⁹ In subsequent work that used first-principles LPPs,⁴⁵⁷ the liquid phases of Na, Mg, and Al⁶¹⁰ were studied at temperatures close to their melting points. It was found that the calculated average coordination numbers and radial distribution functions (RDFs) agreed well

with both conventional KSDFT and experimental outcomes. Recently, the same KEDF and XC functional were used with the PROFESS code to study the dynamic properties of liquid Sn. The simulations reproduced the experimental results for the collective fluid dynamic behavior quite well.⁶¹³

Significant improvements over the SM functional were introduced with the simplified CAT functional³¹⁶ (recall Section 2.4.2). It was developed to study the structure and dynamics of diverse liquids. In the simulations listed next, the LDA XC DFA was used and the temperatures were chosen to be close to experimental values. The studies were as follows:

1. Simple and alkaline earth liquid metals: Cs,⁴⁵² Li,⁴⁶⁹ Na,⁶¹⁴ Mg,⁶¹⁵ and Al³¹⁶ plus liquid alkaline earth metals (Be, Ga, and Ba);⁴⁷¹

2. Liquid alloys: liquid Na–Cs and Na–Cs alloys,⁶¹⁶ Li–Na alloy,⁶¹⁷ Ga, In, and binary alloy Ga–In;⁶¹⁸

3. Liquid surfaces and interfaces: surface of liquid Li and Na,⁶¹⁹ liquid–vapor interface in liquid binary alloys,⁶²⁰ and liquid surface of Sn,⁶²¹ interface of the liquid metals Al,⁶²² liquid–vapor interfaces of Ga and In,⁶¹⁸ liquid–vapor interfaces of liquid binary alloys (NaK, NaCs, and LiNa),⁶²³ and the liquid In surface.⁶¹²

4. Nonmetals and transition metals: liquid Si at near melting point⁶²⁴ and high-pressure conditions,⁶²⁵ bulk liquid Ga,⁶²⁶ liquid noble metals,⁶²⁷ transition metals,⁶¹¹ and liquid Zn, Cd, and Hg divalent metals.⁶²⁸

Most of the studies in the foregoing list employed the simplified CAT KEDF. The earlier works presented comparisons among results obtained with the simplified CAT and the one-point functional $T\lambda\text{vW}$ with $\lambda = 1$. Perhaps unsurprisingly, the simplified CAT KEDF was found to be more accurate and to provide a better description of the structure of the liquid metals considered on the basis of comparison with experimentally determined structure factors.⁴⁵² An additional take-home message from the list of studies above is that the simplified CAT KEDFs in conjunction with pseudopotentials provide a nearly quantitative description of the structure and dynamics of bulk as well as liquid metal surfaces without incurring limitations from the finite-size effects⁶¹⁹ which had plagued conventional KSDFT simulations of similar types of systems.⁶²⁹

In a related application, the Carter group adopted the two-point WGC KEDF in conjunction with the PBE XC functional and BLPS LPPs to study the melting point of Li using the PROFESS code.⁴⁶¹ Figure 19 shows their OFDFT-AIMD results for both the RDF and static structure factors at ion temperatures 470 and 725 K as compared with X-ray diffraction and neutron diffraction data, respectively. The agreement is evident. These results on liquid metals are encouraging indicators of the value of OFDFT-AIMD simulations. Moreover, the OFDFT-AIMD predictions of structure factors predicted from ab initio dynamics for liquid Li are in better agreement with experimental data than the predictions from classical force fields.⁴⁶⁰

4.4. Matter under Extreme Conditions

Most materials physics is done in the same pressure–temperature (P, T) domain as the chemistry of the molecular constituents of those materials, roughly $T \lesssim 2000$ K, $P \lesssim 1$ Kbar. Electronic structure calculations at $T = 0$ K generally suffice therefore, so zero-temperature AIMD is used successfully to calculate the equation of state (EOS) and various properties of the condensed systems. Immediately, the

Radial Distribution Function

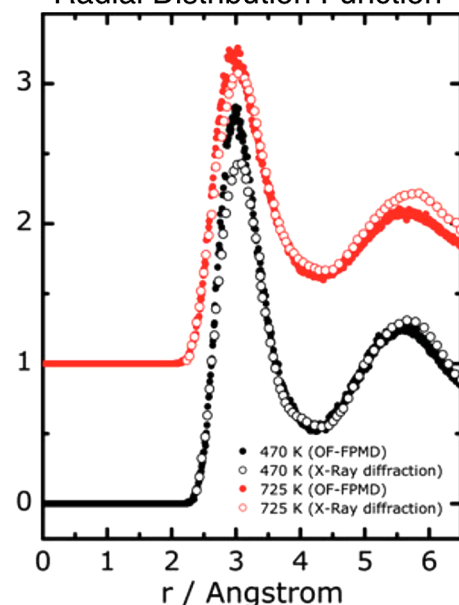


Figure 19. Radial distribution functions (RDFs) of liquid Li at 470 and 725 K. The simulations used 128 independent atoms in the simulation cell in the canonical ensemble. The results at 725 K are shifted upward by one unit for ease of viewing. The legend “OF-FPMD” denotes what we term OFDFT-AIMD. Adapted with permission from ref 461. Copyright (2013) Taylor & Francis.

computational cost-scaling advantage of zero-temperature OFDFT and the KEDFs we have discussed is relevant for such studies.

However, there are systems in which both the chemistry and the condensed phase physics occur at much higher T and P . Especially spectacular examples are found in the interiors of giant planets and exoplanets.⁶³⁰ Other examples of this so-called warm dense matter (WDM) regime occur in inertial confinement fusion.⁶³¹ For WDM, one can have $T \lesssim 10\text{--}15$ eV (with 1 eV = 11,604.5 K) and $P \lesssim 1$ Mbar or even higher.

Systems in the WDM $P\text{--}T$ domain have electron populations that are highly excited. But they are not so thoroughly excited as to make the system into a two-component plasma (electrons, ions) with small quantum corrections. Because quantum mechanics still is dominant, it has proven efficacious to use finite- T AIMD. (Remark: Chemists may be surprised (or bemused) to learn that it is conventional in the physics literature to refer to nonzero T as “finite T ” even though zero is, of course, finite.) Immediately that implicates free-energy DFT.⁶³² For present purposes, it suffices to say that free-energy DFT differs from ground-state DFT in three ways. First, the non-interacting kinetic energy T_s , eq 23, becomes the non-interacting free energy $\mathcal{T}_s = T_s[n] - TS_s[n]$ (where $S_s[n]$ is the non-interacting entropy). Second, the exchange-correlation energy, eq 27, becomes the exchange-correlation free energy $\mathcal{F}_{xc}[n; T]$. And third, the occupation numbers, eq 22, take on Fermi–Dirac T -dependence. Observe that the KS equation is unchanged in form.

The relevance of OFDFT is plain. The simple presence of another physical parameter, T , means that there are more simulations to do than at $T = 0$ K. Moreover, at least in principle all the KS levels are occupied. In practice, many more occupation numbers can be numerically significant than at $T =$

0 K, so the benefit of OFDFT $O(N \ln N)$ scaling relative to conventional KSDFT cubic scaling is amplified compared to the ground state ($T = 0$ K).

We do not review free-energy OFDFT here in detail, but only sketch the key ingredients and give a few examples as a guide to the recent literature and a connection to the ground-state OFDFT methods and KEDFs that are our focus. In parallel with the ground-state case, the critical ingredient is \mathcal{T}_s . For brevity, we refer to approximations for it as KEDFs, even though they have an entropic contribution. The most widely used have been finite- T Thomas–Fermi²⁹ or TFvW. The finite- T von Weizsäcker term was given by Perrot⁶³³ more than 40 years ago, while TF-based finite- T AIMD studies first appeared almost 30 years ago.^{634,635} Despite the fact that TF and TFvW obviously are far from satisfactory on fundamental grounds, they continue to be used.^{636–639} (Remark: It is a misleading peculiarity of the literature that many of the finite- T TF or TFvW-based papers state, incorrectly, that OFDFT is *defined* by the use of those functionals.)

Connection with modern ground-state OFDFT was made in 2012 when Karasiev et al.⁶⁴⁰ presented a method for generalizing a ground-state gradient-dependent (GGA) KEDF into a gradient-dependent \mathcal{F}_s . They demonstrated the workings and effectiveness of that scheme by use on their semiempirical PBE2 GGA KEDF (recall discussion in Subsection 2.4.1 about “modified conjoint” KEDFs). A bit over a year later, the first nonempirical finite- T GGA KEDF was presented, namely VT84F.⁸⁵ Constraint-based, it was constructed with both proper $T = 0$ K limiting behavior and large- T behavior.

As we discussed in Subsection 2.4.1, the constraint satisfaction in VT84F (as a $T = 0$ KEDF) is predicated on physical densities with nuclear-site cusps that satisfy the Kato condition. That property carries over to $T > 0$ K. Recall, from that same discussion, that the LKT KEDF⁸⁶ was constructed to satisfy positivity constraints for typical pseudodensities. By means of the procedure in ref 640. LKT recently also was extended to being a free-energy KEDF,⁶⁴¹ LKTF.

So far as we are aware, those are the only two nonempirical one-point free-energy KEDFs to date. Two-point \mathcal{F}_s approximations are equally rare. To our knowledge, there are only two, both the work of Sjostrom and Daligault.^{166,318,642} They have the structure of eqs 13 with kernel (eq 14), but of course with T -dependent densities and kernels. The first of the Sjostrom–Daligault functionals is straightforwardly the Wang–Teter kernel⁴⁴ extended by calibration to the finite- T Lindhard response. It has a deficiency, namely that for large wavevectors (small distances), the kernel $\omega(\mathbf{k}, T)$ goes strongly negative (see Figure 1 in ref 44). That behavior can yield “unphysical densities with infinite negative energy”.³¹⁸ Reference 318 cured that problem by changing the T_{vw} term to scaled two-point form (temperature-dependence suppressed for clarity),

$$\begin{aligned} \mathcal{F}_{\beta_{vw}} := & \frac{1}{2} \int d\mathbf{r} d\mathbf{r}' [\nabla n^{1/2}(\mathbf{r}) \cdot \nabla n^{1/2}(\mathbf{r}')] [\delta(\mathbf{r} - \mathbf{r}') \\ & + \beta(|\mathbf{r} - \mathbf{r}'|)] \end{aligned} \quad (188)$$

The nonlocal scaling function $\beta(|\mathbf{r} - \mathbf{r}'|)$ and kernel ω then are determined by enforcing the Lindhard response in the HEG limit while requiring that the kernel vanish for large k . In ref 641, this \mathcal{T}_s approximation was denoted “sd β -vW14F”.

Regarding the XC free energy, ground-state E_{xc} forms evaluated with finite- T densities often have been used. This “ground-state approximation” is reasonable up to about $T = 1$ eV in many but not all cases and certainly is not valid generally.^{643–645} Until 2014⁶⁴⁶ there was no strict \mathcal{F}_{xc} counterpart of the Perdew–Zunger LDA E_{xc} .²⁸¹ A partial counterpart to the PBE GGA E_{xc} ²²⁵ was presented in 2014,⁶⁴⁷ but a strict counterpart did not appear until 2018.⁶⁴⁸ While there is a hybrid \mathcal{F}_{xc} ,⁶⁴⁹ that is not relevant to OFDFT, the history of prior development of \mathcal{F}_{xc} approximations is irrelevant here.

With this brief introduction, we turn to a few illustrative examples of calculations. First are two examples of Thomas–Fermi-based calculations. The simpler of the two was by Lambert et al.⁶³⁶ It used simple Thomas–Fermi for \mathcal{F}_s and had no exchange-correlation free-energy contribution in the OFDFT-AIMD. Nevertheless, the prediction for the principal Hugoniot (the locus in P, T of all final states of a material at density ρ reachable by passage of a single shock wave) for iron is rather good for $T \gtrsim 100$ eV when compared to the SESAME table EOS 2140. See Figure 20. Below that temperature the

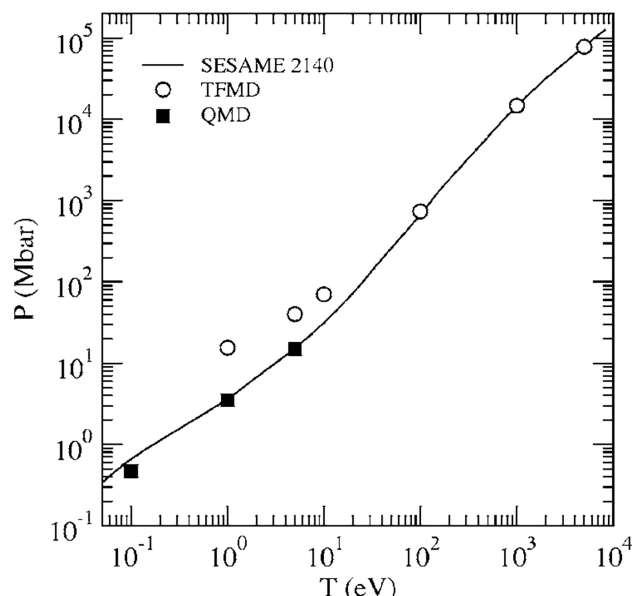


Figure 20. Principal Hugoniot for Fe from free-energy OFDFT with only finite- T TF and no \mathcal{F}_{xc} compared to the SESAME library EOS 2140 and, for lower temperatures, conventional free energy KSDFT AIMD (denoted here as “KSMD”, widely called “QMD”) done with LDA XC. Reprinted with permission from ref 636. Copyright (2006) American Physical Society.

deviations with respect to conventional free-energy KSDFT-AIMD are a clear sign of both the deficiencies of TF and the lack of a \mathcal{F}_{xc} term. This diagnosis is confirmed by a more recent study that includes \mathcal{F}_{xc} contributions (at the level of the ground-state approximation); see ref 651.

The second example is the calculation of the EOS and dynamic structure factor of aluminum by White et al.¹⁶⁷ for $0.1 \leq T \leq 10$ eV. The finite- T OFDFT simulations used 864 atoms in the simulation cell, compared to 108 for the conventional KSDFT finite- T validation calculations, an illustration of the computational cost scaling advantage of OFDFT. Figure 21 shows the resulting EOSs. Once again, one

sees the low-temperature deficiency of the TF approximation. Also obvious is the importance of the inclusion of the XC free energy.

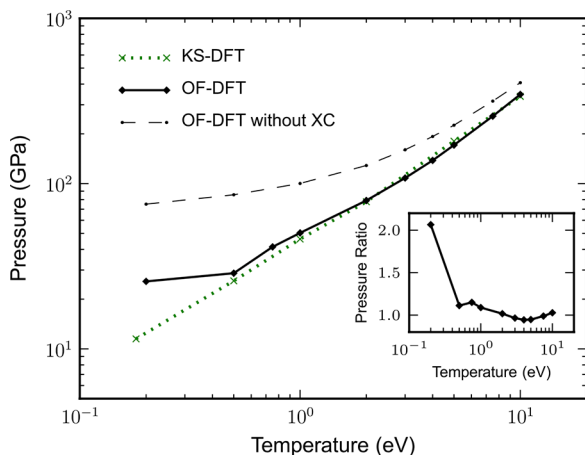


Figure 21. Al EOS at $\rho = 2.7 \text{ g/cm}^3$ from free-energy OFDFT with finite- T TF and with (and without) LDA \mathcal{F}_{xc} (ground-state approximation) compared with conventional free-energy KSDFT results. The inset shows the pressure ratio between OFDFT and conventional KS values. Reprinted with permission from ref 167. Copyright (2013) American Physical Society.

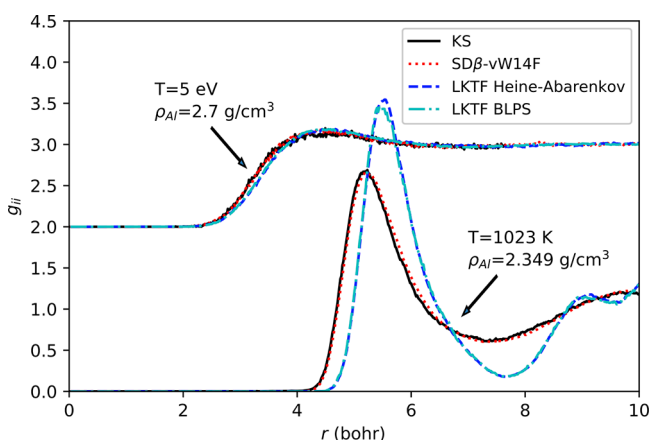


Figure 22. Comparison of calculated RDFs for Al at two state conditions. Conventional free-energy KSDFT results are the black solid curves. Free-energy OFDFT results from the one-point LKTF \mathcal{T}_s are the blue dashed curves; those from the two-point $sd\beta$ -vW14F functional are the red dotted curves. The RDFs for the case $T = 5 \text{ eV}$, $\rho = 2.7 \text{ g/cm}^3$ are shifted upward by two units for clarity (the caption in the original is incorrect). Reprinted with permission from ref 641. Copyright (2020) American Physical Society.

Figure 22 shows a comparison between the best current one- and two-point approximations for \mathcal{T}_s , namely LKTF and $sd\beta$ -vW14F. The physical quantity is the RDF. What is apparent is that the one-point functional is quite competitive at larger T and ρ but that the two-point functional clearly represents the low- T , low- ρ regime better. The calculations were done for two different pseudopotentials, BLPS^{163,164,458} and Heine-Abarenkov.^{445,652} Reassuringly, the two yield very similar results. The isothermal EOS also is interesting. Figure 23 shows that

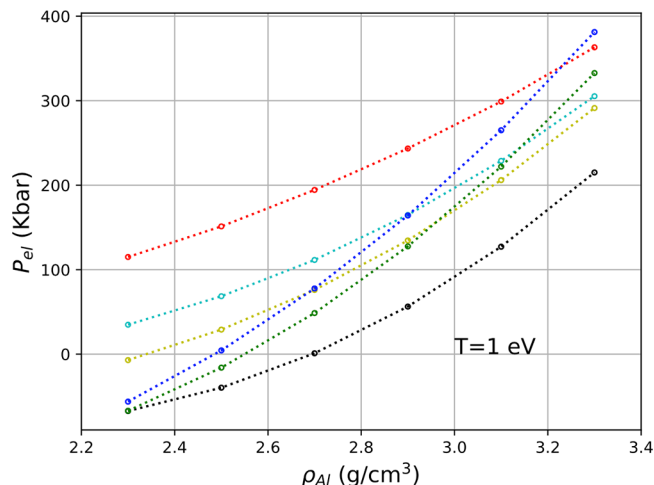


Figure 23. Al EOS at $T = 1 \text{ eV}$ from various \mathcal{F}_s KEDFs. Top to bottom curves at lowest ρ are TF (red), Perrot (teal), TF^1_s -vW (olive), VT84F (blue), and LKTF (green). Conventional free energy KSDFT curve is black. Reprinted with permission from ref 641. Copyright (2020) American Physical Society.

LKTF actually is closest to conventional KSDFT in the region of low bulk density, unlike the comparison for the RDF. Finally, Figure 24 shows the scaling of the wall time per AIMD

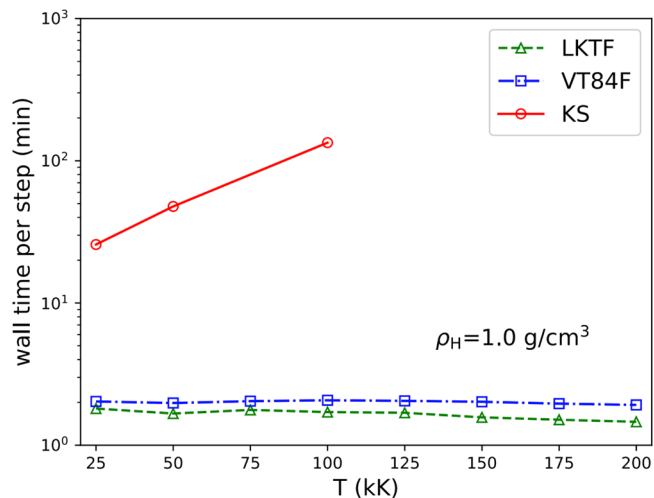


Figure 24. Wall-clock time per MD step for 108 H atoms for conventional free-energy KSDFT vs OFDFT with the LKTF and VT84F KEDFs. Reprinted with permission from ref 641. Copyright (2020) American Physical Society.

step as a function of T from about 25 kK to 200 kK for LKTF and VT84F compared to conventional free-energy KSDFT. The system is 108 H atoms. The promised OFDFT cost scaling, independent of thermal occupation, is obvious. Note that the conventional free-energy KSDFT calculations end at the temperature shown because they were unaffordable above that T . A similar plot is given in ref 315 for the $sd\beta$ -vW14F \mathcal{T}_s KEDF as a function of T over 2 orders of magnitude for the case of 128 H atoms placed randomly.

4.5. Applications with Orbital-Free Embedding Schemes

In Subsection 2.7.5 we discussed the central importance of the so-called nonadditive KEDF eq 158 in subsystem DFT (sDFT). A high-fidelity nonadditive KEDF approximation is

essential; hence, we provide a brief review of LDA, GGA, and two-point nonadditive KEDFs with respect to the type of simulations they can support. This is not a comprehensive survey; rather our intent is to sketch what sDFT can do nowadays as it is enabled by progress in KEDF development. Additional details can be obtained from embedded method reviews.^{180,181,190,408,409,430,653} We also remark that, unless specified differently, all the examples given in this section regard subsystem DFT simulations wherein all subsystem densities are driven to self-consistency.

Obviously, the Thomas–Fermi KEDF is the most rudimentary functional that can be used as a nonadditive KEDF. As originally determined by Wesolowski and Weber,⁴²¹ because the nonadditive KEDF magnitude is small for most weakly interacting subsystems, TF may be a utilitarian choice in some situations⁴²¹ even though it generally gives overly repulsive nonadditive KEDF values and incorrect energy curves.^{296,654,655}

GGA nonadditive KEDFs generally are superior to TF for reproducing ground-state densities and energies.^{181,296,656,657} For those systems in which the interaction between the constitutive subsystems is weak, GGA nonadditive KEDFs have been studied thoroughly. Overall, it was found that GGA nonadditive KEDFs reach sub-5-kcal/mol accuracy in the prediction of most weak intermolecular interaction energies as defined by standard test sets such as S22 or S66.^{658,659} Systems studied include van der Waals complexes,^{660,661} hydrogen-bonded complexes,^{654,656,662,663} and their combinations.^{60,425,426,655} In particular, recent work on partition DFT (related to sDFT but formulated in such a way that the density partitioning is unique while allowing self-consistent solutions for all subsystem densities⁴³⁴) shows that GGA nonadditive KEDFs even can provide a quantitative description of the energetics and electron density for selected diatomics connected by a single covalent bond compared to conventional KSDFT of the full system.^{293,664}

Semilocal nonadditive KEDFs employed in sDFT have enabled AIMD simulations of liquids such as water,^{296,665} and supercritical CO₂,²⁹⁵ as well as solvated systems.⁶⁶⁶ In them, GGA nonadditive KEDFs produced accurate results while providing dramatic computational cost reduction via improved scaling (from cubic to linear). Interestingly, an intrinsic error cancellation between the repulsive nonadditive KEDF and the attractive nonadditive XC led to semilocal sDFT giving improved results compared to those from semilocal conventional KSDFT.²⁹⁶

OFDFT can play an even greater role in sDFT simulations. For example, the combination, in a single subsystem DFT simulation, of conventional KSDFT solvers for molecular subsystems and OFDFT solvers for metallic surfaces recently was reported by Shao et al.⁶⁶⁷ In that work, the binding energies of molecules at Al(111) surfaces were reported to be within a few meV of the conventional KSDFT values. Additionally, thanks to the advantageous computational scaling of OFDFT, the hybrid OFDFT/KSDFT subsystem DFT method yielded the electronic structure of large water/Al(111) interfaces in record wall times.

Perhaps the most spectacular application to date of sDFT with semilocal nonadditive KEDFs is credited to the VandeVondele group.⁶⁶⁸ They tackled a tobacco virus completely *ab initio* by combining sDFT with linear-scaling conventional KSDFT techniques for the subsystems. The virus is effectively a protein with quaternary structure. It was

geometry optimized in a simulation cell that was filled by water molecules. In Figure 25 we show the electronic structure of the

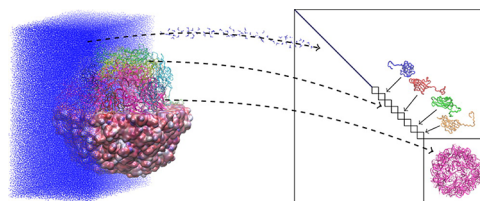


Figure 25. Electronic structure of a tobacco virus computed with sDFT combined with linear-scaling conventional KSDFT for the monomers making up the protein and the solvent. The figure maps portions of the virus onto its density matrix as defined by sDFT, with each diagonal block corresponding to one subsystem. The off-diagonal blocks are zero because orbitals of different subsystems belong to different KS systems which have independent Hilbert spaces and therefore need not be orthogonal. The nonadditive KEDF used is Lee–Lee–Parr.⁵⁸ Reprinted with permission from ref 668. Copyright (2016) American Chemical Society.

tobacco virus computed by them colored by features of the electrostatic potential. While there is no particular reason for carrying out the geometry optimization of a virus, the simulation of ref 668 is a source of excitement and makes one realize that the all-electron, *ab initio* description of biological systems is an achievable goal and not science fiction.

Beside these applications, a vast array of chemical problems, for instance, solvation free energies,⁶⁶⁶ solvent effects on different types of spectroscopies,^{669,670} magnetic properties,^{671–675} charge-transfer states,^{676–680} open-shell systems,⁶⁸¹ and excited states,^{657,682–687} have been explored by semilocal sDFT and its time-dependent extension.^{653,657,684,688} The fact that research continues to yield improved KEDFs suggests strongly that nonadditive KEDFs also will improve and, with that, the applicability of sDFT also will expand.

In addition, ways to expand the applicability of nonadditive KEDFs arise by coupling them with innovative embedding schemes. For example, the 3FDE method^{689,690} handles strong interactions, for which nonadditive KEDFs would fail, by using a constraining potential derived from capping groups. That constraining potential also maintains a correct electron density in the regions of strong inter-subsystem density overlap. The recent adoption of sDFT to improve many-body expansions also is an interesting way to expand the applicability of sDFT.^{691,692}

Along these lines of thought, Laricchia et al. tested various Laplacian-level meta-GGA nonadditive KEDFs for sDFT calculations.²⁸⁵ They found improved sDFT results compared to GGA nonadditive KEDFs when testing interaction energies of small diatomics.

A step forward in accuracy and applicability has been achieved recently by employing two-point KEDFs. The reasons for the comparatively late employment of two-point KEDFs as nonadditive KEDFs are twofold. First, most such KEDFs are primarily suitable for weakly inhomogeneous systems (such as bulk metals). This limitation is at odds with sDFT. In it, the subsystem densities decay at the boundaries. Second, while there exist two-point KEDFs suitable for non-homogeneous systems, most either suffer from numerical instability, can only handle weak non-homogeneity,²⁹⁷ or are too expensive to be employed in large-scale sDFT simulations.⁷⁷ However, the LMGP functionals discussed in Subsection 2.4.2 can be used as

nonadditive KEDFs because they are particularly well suited to describe finite systems.⁸⁹ LMGP provides considerable performance improvement relative to the best GGA functionals in terms of both interaction energies and predicted electron densities.⁸⁹ Evaluation of the full significance of two-point nonadditive KEDFs for sDFT awaits comparable large-scale sDFT calculations. Note, however, that recently Shao et al.⁴²⁵ constructed a GGA functional that mimics the behavior of two-point functionals (i.e., on one hand to cut off the von Weizsäcker functional in high-density regions and on the other hand to cut off the TF functional in low-density, asymptotic regions). Their functional improved upon existing GGA nonadditive KEDF performance by matching the accuracy of two-point nonadditive KEDFs for interaction energies and electron densities for the S22-5 and S66^{658,659} test sets.

Thus far, we have discussed the accuracy of sDFT simulations by citing their ability to reproduce interaction energies (and energy curves) between molecules, and the structure and dynamics of molecular condensed phases (hence, we inferred also the quality of the computed atomic forces). There are, however, other properties that can be adjudged, such as dipole moments, NMR properties, optical spectra, etc. Clearly, the quality of the predicted molecular properties depends on the quality of the electron density, which in turn depends on the quality of the potential employed. In sDFT this translates to the quality of the embedding potential and, as in all KS-DFT calculations, the quality of the XC functional. While we have not dwelled on some of these important details, we refer the interested reader to excellent and comprehensive reviews on the subject (see, e.g., pp 335–337 and 345–346 of ref 181 and sections 5.2–5.6 of ref 190).

For more details on high-efficiency implementations, parallelization schemes, and other details of sDFT implementations, we refer the interested reader to refs 577–579, 665, 667, 668, and 693–695.

5. CONCLUSION

With this review, we have attempted to provide the reader with systematic access to the emerging value and quality of orbital-free density functional theory (OFDFT), both in advances of understanding the theory itself and in improved kinetic energy density functionals (KEDFs). Our claim is that because of steady and strong advances in the development of approximate KEDFs, OFDFT has already played a significant role in understanding materials and cluster chemistry and physics. We expect that role to grow and expand. That expectation is rooted in the growing number of investigators of OFDFT and codes implementing OFDFT. Given promising recent developments surveyed here, applications to catalysis, for example, as well as other parts of chemistry and materials science are expected to surface soon, either with pure OFDFT simulations or with other, related methods such as embedding and decomposition schemes.

In the modeling of chemical processes, atom dynamics and finite-size effects constitute the biggest bottlenecks in current simulation protocols. As we showed in Section 4, OFDFT already resolves those issues for a large class of systems. The challenge now is to widen its applicability to embrace an even larger set of system types. One of the persistent roadblocks to universal applicability of OFDFT has been the limited accuracy of available KEDFs. Another is the lack of local pseudopotentials for second-row elements and transition metals. While such

roadblocks affect pure OFDFT, we have described several hybrid schemes that are less affected by these shortcomings.

The historical account of the development of OFDFT given in Section 1 and the theoretical background presentation in Section 2 show clearly that only by following a rigorous, constraint-satisfaction strategy can one formulate generally applicable (transferable) KEDFs. Thus, we hope this review will help guide OFDFT developers as well as spark interest in employing OFDFT among those research groups that customarily have employed only conventional KSDFT in their research. This is particularly relevant, as OFDFT is now accessible through several software suites (see Section 3).

Obviously, we are indebted to the OFDFT research community that came before us and until recently was rather small. Spanning chemistry, physics, and materials science, that hardy band has inspired us and our research groups over the years, with healthy and constructive debate aimed at the goal of serving the broader scientific community with useful, predictive, swift, ab initio electronic structure methods.

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Author Contributions

[†]All authors contributed equally to this work. CRediT: **Wenhui Mi** funding acquisition, writing-original draft, writing-review & editing; **Kai Luo** funding acquisition, writing-original draft, writing-review & editing; **Samuel B. Trickey** conceptualization, funding acquisition, project administration, writing-original draft, writing-review & editing; **Michele Pavanello** conceptualization, funding acquisition, project administration, writing-original draft, writing-review & editing.

Notes

The authors declare no competing financial interest.

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