

Performance Improvement of Deorbitalized Exchange-Correlation Functionals

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Deorbitalization of a conventional meta-generalized-gradient exchange-correlation approximation replaces its dependence upon the Kohn-Sham kinetic energy density with a dependence on the density gradient and Laplacian. In principle, that simplification should provide improved computational performance relative to the original meta-GGA form because of the shift from an orbital-dependent generalized Kohn-Sham potential to a true KS local potential. Often that prospective gain is lost because of problematic roughness in the density caused by the density Laplacian and consequent roughness in the exchange-correlation potential from the resulting higher-order spatial derivatives of the density in it. We address the problem by constructing a deorbitalizer based on the RPP deorbitalizer [Phys. Rev. Mater. **6**, 083803 (2022)] with comparative smoothness of the potential along with retention of constraint satisfaction as design goals. Applied to the r²SCAN exchange-correlation functional [J. Phys. Chem. Lett. **11**, 8208 (2020)], we find substantial timing improvements for solid-state calculations over both r²SCAN and its earlier deorbitalization for high precision calculations of structural properties, while improving upon the accuracy of RPP deorbitalization for both solids and molecules.

I. INTRODUCTION

The enormous impact of Hohenberg-Kohn-Sham density functional theory (DFT) upon computational study of molecular and materials properties is a consequence of the remarkable cost-accuracy balance of DFT when implemented with modern approximate exchange-correlation (XC) functionals. The cost-accuracy trade-off is determined by the intrinsic limits and transferability of the specific XC approximation used. That compromise clearly differs by problem class. A small set of very large molecules can be treated with an XC approximation far up the Perdew-Schmidt “Jacob’s ladder” complexity hierarchy [1]. In contrast, the computational costs of high-throughput screening calculations on large data sets of such molecules or ab initio molecular dynamics (AIMD)

on counterpart condensed phases compel use of lower-rung approximations. See, for example, the discussion in Sections 2.2.21-23 of Ref. 2.

The SCAN meta generalized-gradient approximation (metaGGA) [3, 4] and its variants [5–7], have proven to be effective compromises for accuracy and efficiency in structural and energetic calculations for both solids and molecules [4, 8] albeit with some limitations [9, 10]. A design advantage of such meta-GGA functionals is the introduction of inherently nonlocal density information through the use of the non-interacting (Kohn-Sham, KS) kinetic energy density [11, 12],

$$\tau_s := \frac{1}{2} \sum_i f_i |\nabla \phi_i(\mathbf{r})|^2 \quad (1)$$

with orbitals ϕ_i and occupation numbers f_i . A resultant design advantage is satisfaction of many more exact constraints than possible with a GGA, particularly for the iso-orbital limit.

Despite the successes, there are difficulties of imple-

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mentation with these metaGGAs that limit their full exploitation. The explicit orbital-dependence introduced by τ_s causes typical practical application to rely on non-local XC potentials in generalized KS (gKS) equations. Solutions to those can be slow relative to pure KS implementations [13]. This slowness also originates with the complicated form of the functionals and their potentials, their consequent sensitivity to the density [14], and concomitant need for dense grids for converged integrals [15, 16]. Calculations sometimes are done non-self-consistently, [8, 17] and self-consistent pseudopotentials for these functionals have proven difficult to develop. [14]

Effort to resolve these issues has led to the redesign of SCAN, first by streamlining performance at the cost of significant constraint noncompliance, [6] then by iso-orbital indicator redefinition, [5] and, most recently, in r²SCAN, [7] by restoration of most of the constraints met by SCAN, while retaining the improvements in numerical performance of prior variants. Nonetheless, some numerical problems remain. [14, 16]

Deorbitalization [13, 18, 19] has been developed as a strategy both to obviate these issues and to retain the interpretive power of the pure KS equation. In deorbitalization, the τ_s dependence used in meta-GGA XC functionals as a local indicator of bond character is replaced with a function of the density $n(\mathbf{r})$, its gradient ∇n , and Laplacian $\nabla^2 n$. The result is a pure XC density functional and a pure (local) KS potential.

The obvious deorbitalization requirement is a sufficiently accurate approximate orbital-free kinetic energy density (KED) functional. That is an active area of research. [20] Practical deorbitalization has come through the methodology of Mejía-Rodríguez and Trickey [13, 18, 19] (M-RT). They reparametrized existing KED functionals, e.g. Perdew-Constantin (PC) [21] and others from orbital-free DFT [20], to match KED-dependent indicator functions of the first 18 isolated atoms. The most generally successful case of M-RT deorbitalization involves the reparametrization (PC_{opt}) of original PC. That gave successful deorbitalizations of both SCAN [13, 18] and r²SCAN [19], in the sense of providing what was deemed to be a faithful reproduction of the parent metaGGA error patterns for structural and energetic properties evaluated on standard molecular and crystalline benchmarks.

That faithfulness also has held up in application to materials and outside the original test sets. [22, 23] Other KEDs, e.g. Cancio-Redd (CR) [24] actually may lead

to improved error patterns relative to use of the exact KED for some metaGGAs, but may not for best-in-breed functionals.

A limitation of the M-RT error-pattern criterion of deorbitalization faithfulness is that it can lead to acceptance of deorbitalized XC functionals that deviate noticeably from the exact form in the limit of slowly varying density, *i.e.*, constraint violation. In the context of rethinking the PC functional form, the recent OFR2 study [25] emphasized constraint satisfaction, including restoration of correct slowly-varying limit behavior. That recovers fourth-order gradient expansion compliance lost in going from SCAN to r²SCAN [6]. The resulting deorbitalization does well in calculations for solids (retaining the numerical advantages of rSCAN) but not as well as r²SCAN-L (r²SCAN deorbitalized by PC_{opt}, the reparametrized version of PC used by M-RT[13, 18]) for molecules.

As an aside for clarity, note that a second modification of the original M-RT deorbitalization has been developed[26] to treat metaGGAs that rely on more than one iso-orbital indicator, particularly the Tao Mo functional (TM) [27]. That is not relevant here.

Alternative strategies to Laplacian-level deorbitalization include use of higher-order spatial derivatives [28–30] and deorbitalization of the exchange potential only [17]. Those are of only contextual interest here, as they may have more significant numerical challenges than those arising from the M-RT procedure.

Given success at reproducing the test-case performance of existing orbital-dependent metaGGAs, the challenge is the effective numerical implementation of such a deorbitalized functional. The well-known sensitivity to density Laplacians serves as a caution. See but for one example, the discussion in Ref. 13. Nonetheless, some of the M-RT functionals [13, 18] have shown marked performance improvement over their gKS parents, with time per self-consistent-field (SCF) cycle of the Kohn-Sham solution as much as a factor of three shorter with the VASP code [31, 32]. But recent work indicates [26] that it is quite typical for deorbitalized functionals to require many more self-consistent cycles to achieve SCF convergence than for their parent functionals with gKS, thus offsetting the gains made per cycle.

A usefully accurate description of the KED for deorbitalizing a meta-GGA indicator functional by a simple generalized gradient approximation (GGA) alone does not seem possible [13]. But, like Laplacian-based KEDFs,

Laplacian-based XC deorbitalizers can be ill-behaved with regards to integral convergence and require relatively dense grids. [16]. Moreover, they also can have large, spiky variations in the magnitude of the XC potential. Such variations arise from the high-order spatial derivatives in the X potential that follow from $\nabla^2 n$. [25]

The natural, critical question raised by these issues is this: can deorbitalization, in fact, succeed in achieving faster, more reliable metaGGA calculations? At a more detailed level, how does the difficulty of using $\nabla^2 n$ compare with the difficulties of gKS calculations? For which run conditions, and for which applications, if any, does deorbitalization provide a clear advantage? With respect to computational efficiency, where and how can M-RT deorbitalization be improved?

One clear opportunity is to find ways to reduce numerical instabilities induced by the density-Laplacian. In this work, we take a specific step in that amelioration by intervening in those aspects of the deorbitalized functional that are most problematic, while retaining (or even eventually restoring) important constraints along the way. Our scheme is based on exploitation of the Cancio-Redd [24] and Cancio-Stuart-Kuna [33] (CR hereafter) approach of minimizing the structural complexity of KED functionals. We combine that with the constraint satisfaction of the “r²SCAN piece-wise polynomial” (hereafter RPP) deorbitalized indicator function [25] used in the OFR2 functional to produce a deorbitalizer with a smoother X potential than OFR2. We show that this functional performs well compared to r²SCAN and to r²SCAN-L (PC_{opt}) and as well or better than OFR2 for solid test cases. It improves significantly on OFR2 for molecular test sets but still is not as good for those as r²SCAN-L(PC_{opt}). We leave until later a more systematic approach to constructing smoothed functionals along with a measure to assess the result.

Secondly, we report a study of timing characteristics, using a single compute node. We show that deorbitalization can be an attractive option for metaGGA calculations, reproducing similar if not better results than original metaGGA for solids and for equation of state calculations, being up to twice as fast in total compute time as gKS calculations. At the same time, we find a number of situations in which the two approaches are about equal in performance and slow compared to GGAs, indicating need for future work.

In the remaining presentation, Section II details the deorbitalization approach and the various KED models

utilized. Sec. III describes the methods for performing structural and timing benchmarks. Sec. IV reports the structural bench-marking and detailed timing analysis for solids, including preliminary *ab initio* molecular dynamics (AIMD) outcomes. Sec. V presents our conclusions and prospects for future work.

II. BACKGROUND

A. MetaGGA Structure

A conventional metaGGA XC functional (ignoring spin-decomposition) is defined by

$$E_{\text{xc}}[n] := \int d\mathbf{r} e_{\text{xc}}[n(\mathbf{r}), \nabla n(\mathbf{r}), \tau_s(\mathbf{r})] \quad (2)$$

where both the density n and noninteracting kinetic energy density τ_s are expressed in terms of orbitals ϕ_i and occupation numbers f_i , with

$$n(\mathbf{r}) := \sum_i f_i |\phi_i(\mathbf{r})|^2 \quad (3)$$

and τ_s given by Eq. (1). To cope with the explicit orbital dependence in the XC energy (and therefore, implicit density dependence), customarily the ground state energy is found via the gKS procedure. In it, the XC potential is calculated as the functional derivative of the energy with respect to the individual orbitals. The result is the gKS equation in which the kinetic energy operator becomes

$$-\frac{1}{2}\nabla^2 + \nabla \cdot \left(\frac{\partial e_{\text{xc}}}{\partial \tau_s} \right) \nabla, \quad (4)$$

thus including a dependence upon the XC energy beyond that of the multiplicative potential of the pure KS formulation. (In practical implementation, that term can be rendered as an orbital-dependent potential, hence still requiring a generalization of the KS procedure).

In SCAN and some of its descendants, τ_s enters the XC functional via the iso-orbital indicator α ,

$$\alpha(n, \nabla n, \tau_s) := \frac{\tau_s - \tau_w}{\tau_{\text{TF}}} . \quad (5)$$

Here τ_w is the von Weizsäcker KED, namely that of a system with a single occupied orbital and density n ,

$$\tau_w = \frac{1}{8} \frac{|\nabla n|^2}{n} \quad (6)$$

and τ_{TF} is the KED of the homogeneous electron gas of density n

$$\tau_{\text{TF}} = \frac{3}{10}(3\pi^2)^{2/3}n^{5/3}. \quad (7)$$

Akin with other refinements of SCAN, r²SCAN uses a regularized version of α ,

$$\bar{\alpha} := \frac{\tau_{\text{S}} - \tau_{\text{W}}}{\tau_{\text{TF}} + 0.001\tau_{\text{W}}}. \quad (8)$$

This regularization is primarily to handle undesirable behavior for exponentially small densities. Nevertheless α serves as a convenient point of reference from which to define orbital-free KE density models.

The central roles of α and $\bar{\alpha}$ make it helpful to summarize, for context, the key structural elements of r²SCAN exchange. Following Eq. (2) with $e_{\text{XC}} = n(\mathbf{r})(\epsilon_{\text{x}} + \epsilon_{\text{c}})$, for exchange we have

$$\epsilon_{\text{x}}^{\text{r}^2\text{SCAN}}(p, \bar{\alpha}) = \epsilon_{\text{x}}^{\text{LDA}}[n]F_{\text{x}}^{\text{r}^2\text{SCAN}}(p, \bar{\alpha}) \quad (9)$$

$$\epsilon_{\text{x}}^{\text{LDA}}[n] = -(3/4)(3/\pi)^{1/3}n^{1/3} \quad (10)$$

$$F_{\text{x}}^{\text{r}^2\text{SCAN}}(p, \bar{\alpha}) = \{h_{\text{x}}^1(p) + f_{\text{x}}(\bar{\alpha})[h_{\text{x}}^0 - h_{\text{x}}^1(p)]\}g_{\text{x}}(p) \quad (11)$$

Here, the dimensionless reduced gradient (squared) is

$$p := s^2 \equiv \frac{|\nabla n|^2}{4(3\pi^2)^{2/3}n^{8/3}}. \quad (12)$$

The $h_{\text{x}}^1(p)$ is a GGA form that is a rather complicated function of p . It enforces the gradient expansion of exchange through second order for $\alpha \approx 1$. Switching between GGA forms for $\alpha \approx 1$ and the iso-orbital limit ($\alpha \approx 0$), and extrapolation to $\alpha \rightarrow \infty$ is accomplished with the function $f_{\text{x}}(\bar{\alpha})$. It has two exponential regions ($\bar{\alpha} < 0$ and $\bar{\alpha} > 2.5$) joined by a seventh-order polynomial in $\bar{\alpha}$. $g_{\text{x}}(p)$ is a damping function that goes to unity in the limit $p \rightarrow 0$ and goes to zero as p grows arbitrarily large. Detailed expressions are in the Supplemental Material for Ref. 7.

An important bound, given the centrality of α , is

$$\tau_{\text{S}} \geq \tau_{\text{W}} \Rightarrow \alpha \geq 0. \quad (13)$$

This may be recognized as the non-negativity of the contribution to the Kohn-Sham kinetic energy derived from Pauli exclusion. Note that the r²SCAN switching function $f_{\text{x}}(\bar{\alpha})$ just described is set up to enforce this requirement even if numerical precision errors otherwise would violate it. The indicator α is related trivially to the Pauli

kinetic energy density which is the subject of much of orbital-free DFT (OF-DFT)[20],

$$\tau_{\theta} := \tau_{\text{S}} - \tau_{\text{W}} \quad (14)$$

or

$$\alpha = \tau_{\theta}/\tau_{\text{TF}} \equiv F_{\theta}. \quad (15)$$

Here F_{θ} is the Pauli enhancement factor of OF-DFT.

To deorbitalize a conventional metaGGA according to M-RT, the exact KS KED is replaced by an approximate orbital-free (OF) semi-local density functional

$$\tau_{\text{OF}} = \tau(n, \nabla n, \nabla^2 n). \quad (16)$$

The deorbitalized metaGGA then is

$$E_{\text{XC}}[n] = \int e_{\text{XC}}\{n(\mathbf{r}), \nabla n(\mathbf{r}), \tau_{\text{OF}}[n(\mathbf{r}), \nabla n(\mathbf{r}), \nabla^2 n(\mathbf{r})]\} d\mathbf{r}. \quad (17)$$

Minimization of the total energy with respect to n then yields a pure KS equation with a local XC potential of the form

$$v_{\text{XC}} = \frac{\partial e_{\text{XC}}}{\partial n} + \frac{\partial e_{\text{XC}}}{\partial \tau_{\text{OF}}} \frac{\partial \tau_{\text{OF}}}{\partial n} - \nabla \cdot \left(\frac{\partial e_{\text{XC}}}{\partial \nabla n} + \frac{\partial e_{\text{XC}}}{\partial \tau_{\text{OF}}} \frac{\partial \tau_{\text{OF}}}{\partial \nabla n} \right) + \nabla^2 \left(\frac{\partial e_{\text{XC}}}{\partial \tau_{\text{OF}}} \frac{\partial \tau_{\text{OF}}}{\partial \nabla^2 n} \right), \quad (18)$$

In it, the first and third terms treat the explicit dependence of e_{XC} on n and ∇n respectively, while the others treat implicit dependence through τ_{OF} . A closely related quantity is the OF kinetic energy potential

$$v_{\text{S}} := \left(\frac{\partial \tau_{\text{OF}}}{\partial n} \right) - \nabla \cdot \left(\frac{\partial \tau_{\text{OF}}}{\partial \nabla n} \right) + \nabla^2 \left(\frac{\partial \tau_{\text{OF}}}{\partial \nabla^2 n} \right). \quad (19)$$

In the OF-DFT literature this is more commonly expressed as taking the functional derivative of a model Pauli KED, τ_{θ} . That yields the Pauli potential via

$$v_{\theta} = v_{\text{S}} - v_{\text{W}}, \quad (20)$$

with v_{W} the KE potential for τ_{W} .

Comparison of Eqs. (4) and (18) illuminates the distinct, key challenges for ordinary versus deorbitalized metaGGA XC numerical implementations. A conventional, orbital-dependent metaGGA introduces a complicated kinetic energy operator in addition to the ordinary

local Kohn-Sham potential. A deorbitalized counterpart produces a purely local potential whose construction involves up to fourth spatial derivatives of the density, with concomitant possible numerical difficulties.

B. Deorbitalization models

An orbital-free KED functional of the general form Eq. (16) obviously can be decomposed as in Eq. (14). The result may be reduced by scaling constraints to

$$\tau_{\text{OF}}(n, \nabla n, \nabla^2 n) = F_{\theta}^{\text{OF}}(p, q) \tau_{\text{TF}}(n) + \tau_{\text{W}}(n, p). \quad (21)$$

Here F_{θ} is as in Eq. (15) and

$$q := \frac{\nabla^2 n}{4(3\pi^2)^{2/3} n^{5/3}} \quad (22)$$

is the dimensionless reduced density Laplacian, partner to the reduced density gradient p defined in Eq. (12). In addition to scaling, other important constraints include α non-negativity [Eq. (13)] and the gradient expansion of F_{θ} for slowly-varying density. [34, 35]

KED functionals that have proven particularly useful for deorbitalization share several characteristics. They are proper metaGGAs, in that they depend upon n , p , and q . The design of most has followed, to some degree at least, the strategy employed in the PC KED functional [21], that is to build a model for the KED in the slowly-varying density limit. Then, because models based on the formally correct gradient expansion in that limit vary much more slowly with p than the iso-orbital limiting case (the von-Weizsäcker functional, τ_{W}), eventually such models fail the non-negativity constraint [36], $\tau_s \geq \tau_{\text{W}}$. That failure is taken to indicate the onset of an iso-orbital or nearly iso-orbital spatial region, and the slowly varying KED is replaced by τ_{W} by means of a suitable switching function.

For PC and PC_{opt} (the reparametrized PC used by MRT [13, 18]), the KED for the slowly-varying limit is a modified fourth-order gradient expansion for which the Pauli enhancement function is

$$F_{\text{PC}}^{\text{SV}} = \frac{1 + \Delta F_{\theta}^{(2)} + \Delta F_{\theta}^{(4)}}{\sqrt{1 + [\Delta F_{\theta}^{(4)} / (1 + 5p/3)]^2}}. \quad (23)$$

The superscript “SV” denotes “slowly varying”. The ingredient quantities are the second and fourth-order gra-

dient expansion corrections

$$\Delta F_{\theta}^{(2)} := -\frac{40}{27}p + \frac{20}{9}q \quad (24)$$

$$\Delta F_{\theta}^{(4)} := \frac{8}{81}q^2 - \frac{1}{9}pq + \frac{8}{243}p^2 \quad (25)$$

The second-order gradient correction $\Delta F_{\theta}^{(2)}$ consists of a $5p/27$ contribution from the kinetic energy gradient expansion minus a $5p/3$ factor from the von Weizsäcker KED ($\tau_{\text{W}} = (5p/3)\tau_{\text{TF}}$). That yields an overall negative slope with respect to p . Together with the possibility of negative q , this leads to eventual incipient violation of KED non-negativity which forces the switch from slowly-varying to the iso-orbital model forms.

For PC, the interpolation function between the slowly-varying form of the Pauli enhancement factor and the von Weizsäcker lower bound $F_{\theta}^{\text{W}} = 0$ takes the form

$$\Theta_{\text{PC}}(x) = \begin{cases} 0, & x \leq 0 \\ f_{\text{PC}}(x/x_0) & 0 < x < x_0 \\ 1, & x \geq x_0 \end{cases} \quad (26)$$

where

$$f_{\text{PC}}(t) = \left[\frac{1 + e^{1/(1-t)}}{e^{1/t} + e^{1/(1-t)}} \right]^b, \quad 0 < t < 1. \quad (27)$$

Putting things together, the final form is

$$F_{\theta}^{\text{PC}} = \alpha_{\text{PC}} = F_{\text{PC}}^{\text{SV}} \Theta_{\text{PC}}(F_{\text{PC}}^{\text{SV}}) \quad (28)$$

The original PC parameter values are $x_0 = 0.5389$, $b = 3$. The PC_{opt} values are $x_0 = 1.784720$, $b = 0.258304$ [13], determined by fitting to the KED α values of small- Z atoms. That reparametrization helps to produce faithful deorbitalization of some metaGGAs but comes at the cost of a somewhat inaccurate form for the slowly-varying limit ($p, q \rightarrow 0$).

Some drawbacks of the PC model were uncovered in Ref. 33, which provided a modified form to fix them. The switching model Θ_{PC} was found to cause unphysical features in the KED in covalent bonds, particularly for systems treated with pseudopotentials. Concurrently, the PC limit for $q \rightarrow \infty$ and p finite causes an unphysical treatment of the exponentially decaying density in the asymptotic region of a molecule, characterized by $q \rightarrow \infty$ and $p/q \rightarrow 1$. The first problem can be resolved by use of a switching factor that obeys $\tau > \tau_{\text{GE}}$ and is as smooth as feasible. The second problem can be resolved by recognizing that the second-order gradient expansion for the

KED has the scaling behavior of the exact KED for an exponentially decaying density and also is valid for small p and q . Thus it can suffice as a bare-bones model for the slowly-varying limit.

Those choices, with further refinement [24], lead to the CR model, namely

$$\alpha_{\text{CR}} := 1 + \Delta F_{\theta}^{(2)} \Theta_{\text{CR}} \left(\Delta F_{\theta}^{(2)} \right) \quad (29)$$

with

$$\Theta_{\text{CR}}(z) = \left[1 - \exp \left(-\frac{1}{|z|} \right)^a (1 - H(z)) \right]^{1/a} . \quad (30)$$

$H(z)$ is the Heaviside unit step function. The exponent $a = 4.0$ produces reasonably close estimates for the total KE of atoms[24], while $a = 2.0$ produces somewhat smaller Pauli potentials.

The RPP deorbitalizer [25] is a PC variant designed specifically for deorbitalizing r²SCAN whilst retaining constraint satisfaction. As such it starts with the same form as Eq. (28),

$$\alpha_{\text{RPP}} = F_{\text{RPP}}^{\text{SV}} \Theta_{\text{RPP}}(F_{\text{RPP}}^{\text{SV}}) \quad (31)$$

where $F_{\text{RPP}}^{\text{SV}}$ is, as before, a metaGGA suitable for the slowly varying limit, to wit

$$F_{\text{RPP}}^{\text{SV}} = 1 + \Delta F_{\theta}^{(2)} + \Delta F_{\theta}^{\prime(4)} + \Delta F_{\theta}^{\text{asy}} . \quad (32)$$

In this expression, the fourth-order term $\Delta F_{\theta}^{\prime(4)}$ has the form of Eq. (25) but with dramatically altered coefficients: $b_{qq} = 1.801019$, $b_{pq} = -1.850497$, and $b_{pp} = 0.974002$. Those correct a corresponding error in the r²SCAN exchange functional to restore gradient expansion compliance to fourth order in the slowly-varying limit. The last term of Eq. (32) is higher than fourth-order and defines the asymptotic large p, q behavior. It is given by

$$\begin{aligned} \Delta F_{\theta}^{\text{asy}} = & c_3 p^2 \left(e^{-|c_3|p} - 1 \right) + \left(\Delta F_{\theta}^{\prime(4)} - c_3 p^2 \right) \\ & \times \left\{ \exp \left[-\left(\frac{p}{c_1} \right)^2 - \left(\frac{q}{c_2} \right)^2 \right] - 1 \right\} . \end{aligned} \quad (33)$$

This imposes the second-order gradient expansion as the limit for $p, q \rightarrow \infty$, p/q finite, as in CR. Optimal coefficient values were determined against appropriate norms

[25] to be

$$c_1 = 0.202352 \quad (34)$$

$$c_2 = 0.185020 \quad (35)$$

$$c_3 = 1.53804 . \quad (36)$$

The switching functional Θ_{RPP} involves the same piecewise logic as in PC, Eq. (26), but with the nonanalytic switching function f_{PC} , Eq. (27) replaced by the polynomial

$$f_{\text{RPP}}(t) = 20t^3 - 45t^4 + 36t^5 - 10t^6 . \quad (37)$$

The switching constant $x_0 = 0.819411$. (There is no b constant to set.)

In our context, a rather obvious step is to seek a smoothed RPP (SRPP) functional by replacing the RPP switching functional with the CR form,

$$\alpha_{\text{SRPP}} = 1 + (F_{\text{RPP}}^{\text{SV}} - 1) \Theta_{\text{CR}}(F_{\text{RPP}}^{\text{SV}} - 1) . \quad (38)$$

A somewhat smoother functional, denoted SRPP2, arises from employing the exponent $a = 2$ in Θ_{CR} .

Fig. 1 shows how these design choices affect the behavior of the Pauli enhancement factors, *i.e.*, the α to be used in deorbitalization. That figure presents F_{θ} versus p for the specific value $q = 0$. The von-Weizsäcker lower bound value of $F_{\theta}^{\text{W}} = 0$ appears as a black solid line. GEA2 denotes the gradient expansion model $1 + \Delta F_{\theta}^{(2)}$ that characterizes the slowly varying limit, $p, q \rightarrow 0$. The slowly-varying limit for the PC metaGGA starts at the homogeneous electron gas value of 1 at $p = 0$ and at first follows along the GEA2 trajectory. But GEA2 transgresses the von-Weizsäcker bound before $p = 0.8$. (Recall the negative slope in the p -coefficient of the gradient expansion discussed earlier.) The switching function Θ_{PC} avoids that crossing by inducing the sudden dip in PC off the GEA2 and subsequent zero value, thus complying with the non-negativity constraint for F_{θ} . That dip is a fairly clear candidate to suspect as the cause of fluctuations in the PC Pauli potential and in the potential of a XC functional deorbitalized with PC. The dip leads to rapid variations in the PC KED in regions of near iso-orbital density, such as covalent bonds, [33] which would be exacerbated in calculating the potential. This supposition is shown to be borne out in the following section. In contrast, the CR model (blue-dashed) and SRPP (black dot-dashed) make the transition (from GEA2 to von Weizsäcker bound) in a much smoother

fashion, while RPP is an intermediate case. Finally F_θ for PC_{opt} fails to satisfy the homogeneous electron gas limit ($F \rightarrow 1, p, q \rightarrow 0$) but does have a reasonably smooth transition.

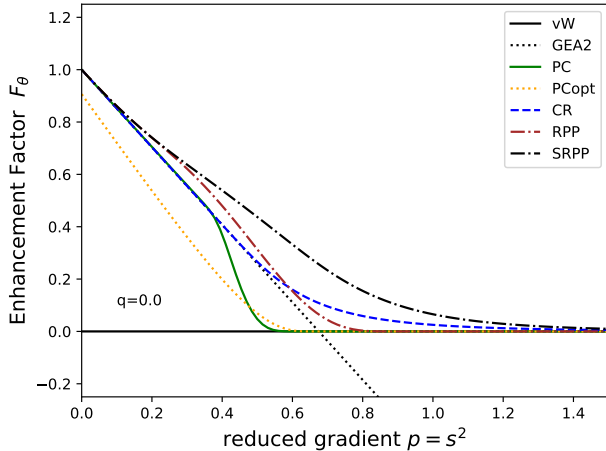


FIG. 1: Pauli enhancement factor F_θ for various KED functionals as a function of reduced density gradient p with reduced Laplacian $q = 0$.

C. Kinetic and exchange potentials

The smoothing problems that arise in the deorbitalizers are well illustrated by a plot of the kinetic potential v_θ (Eq. (19)) for the Hydrogen atom for various Laplacian-dependent KED functionals. See Fig. 2.

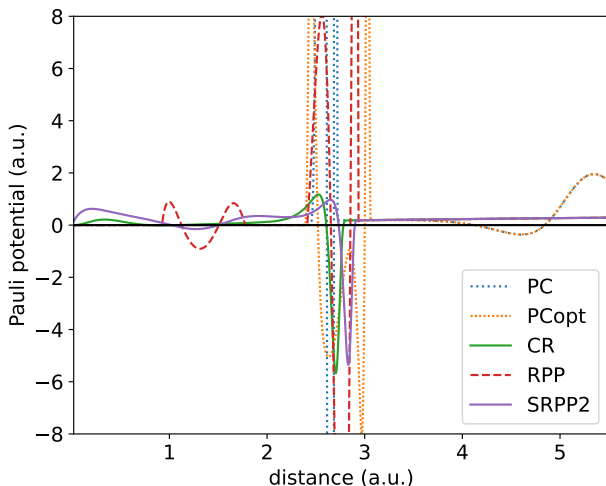


FIG. 2: Pauli kinetic potential $v_\theta = v_\tau - v_w$ for various Laplacian-level KED functionals for the hydrogen atom, evaluated at the exact atomic density. The exact value for the H atom, $v_\theta = 0$, is at the solid black line.

Most of the approximate KED functionals deliver potentials that are reasonably close to the exact value (zero) except in the vicinity of 2.5 to 3.0 a.u. There violent fluctuations in the potential are evident. Those can be traced to a spurious diagnosis of density behavior. At high densities, the KED functionals diagnose hydrogen to be an iso-orbital system, which is correct. But at low density they treat it like a slowly varying electron gas. The fluctuations occur for the values of r at which the transition between the two limits occurs. The problem is worst for PC which suffers oscillations larger than 100 hartrees. The problem is somewhat less severe for PC_{opt} and RPP with oscillations in the tens of hartrees. Only the CR model [24] and its extension to the RPP form, SRPP2, show more reasonable variations, on the order of a few hartrees. In fact, the size of those variations is correlated closely (with the exception of PC_{opt}) with the abruptness of the function used to describe the transition (from iso-orbital to slowly varying electron gas) in the enhancement factor, as seen in Fig. 1. As a caution, we note that the plot shows potentials evaluated with the exact ground-state density for hydrogen. The unreasonable oscillatory behavior almost certainly would be worse for self-consistent densities from the associated potentials.

Figures 3 and 4 show plots of the local part of the exchange potential (Eq. 18) for the parent meta-GGA functional ($r^2\text{SCAN}$) and the full X potential for three deorbitalized versions of it (PC_{opt} , RPP, and SRPP). The systems are the H atom (at two different length scales) and the Si atom. (Aside: $r^2\text{SCAN}$ deorbitalized with the RPP form has the formal functional name OFR2, but for simplicity in making comparisons with other deorbitalization options, we will continue to refer to it as RPP.) Note that the X potential in Fig. 3 is multiplied by the radial coordinate to bring out features nearer the valence edge where most variation in potentials occurs. For $r^2\text{SCAN}$, the contribution from the functional derivative with respect to the kinetic energy density is omitted, as it is incorporated in the physics in the gKS scheme via Eq. 4. These potentials are computed using a modified version of the Python code *densities* [37] which employs analytic spherical Hartree-Fock densities. They were evaluated on a double-exponential grid with 300 to 10,000 radial points.

Though an unequivocal comparison of the gKS local part and ordinary KS potentials is not possible, it is clear that deorbitalization of $r^2\text{SCAN}$ introduces unphysical oscillations. In particular, we see large noise in the re-

gion $r \approx 3$ au for the H atom, where the Pauli potential of Fig. 2 is seen to fail. However, the SRPP deorbitalization of r^2 SCAN results in modestly less noise in the potential compared to the PC_{opt} or RPP deorbitalizations. A similar trend occurs for Si, with SRPP slightly smoother than RPP and both smoother than PC_{opt} . Interestingly, the RPP and SRPP produce potentials noticeably smoother overall than the local part of the parent metaGGA, despite use of the density Laplacian.

The noticeable bump in the r^2 SCAN H atom potential at large r [Fig. 3(a)] may be a remnant of a major issue in the original SCAN. In a region of exponentially decaying density, v_x for SCAN goes to a large positive value rather than zero [6, 7]. That unphysical behavior forces the use of extremely fine grids and leads to difficulties in obtaining pseudopotentials.[14] (Note that the exact KS exchange potential for the H atom must cancel the self-Hartree potential, hence rv_x must go asymptotically to -1 , which no potential considered here does properly.)

The short-range behavior for any atom is exemplified by the H atom case. All potentials for it fail by diverging at extremely small distances, rather than trending to a small negative value. [38]. Overall v_x from PC_{opt} and SRPP are rather close. Unlike v_x from RPP, they are roughly similar quantitatively to the parent v_x .

D. Noise problem quantification

The unphysically noisy features in the Pauli kinetic potentials shown in Fig. 2 can be diagnosed readily as arising primarily from the density Laplacian term, i.e., the last term of Eq. (19). Clearly it is closely related to the last term in Eq. (18) for the exchange-correlation potential. This makes sense since both have the Laplacian of a functional of the Laplacian of the density. Thus fourth spatial derivatives of the density arise in both as well as third derivatives of e_{xc} in the latter case. Those high-order spatial derivatives intrinsically accentuate oscillations in a function. Reduction of the magnitude of this term should lead to smoother potentials. The reduction in v_θ oscillations from PC to CR shown in Fig. 2 is commensurate with that supposition. Such reduction should yield more efficient numerical performance.

For a quantitative assessment of the extent to which that supposition is correct, we borrow a measure from electrostatics. Observe that satisfaction of Laplace's

equation

$$\nabla^2 \psi = 0 \quad (39)$$

by some function ψ within a specified spatial region is the condition for minimizing the action defined by

$$I = \frac{1}{2} \int |\nabla \psi|^2 dV \quad (40)$$

with respect to variations in ψ , with the value of ψ on the boundary of the solution region fixed.

The impact of the Laplacian term in the kinetic potential thus can be measured by calculating the action for an appropriate ψ . Here we choose

$$I^{\text{deorb}} = \frac{1}{2} \int \left| \nabla \left(\frac{\partial \tau}{\partial \nabla^2 n} \right) \right|^2 dV. \quad (41)$$

This technique has been applied previously to minimize the Laplacian term of the overall exchange-correlation potential. See Ref. 39 for details. Our choice to focus on the kinetic potential reflects the fact that the Laplacian term of the X potential actually generates two plausibly problematic contributions. One, $\nabla^2(\partial e_{\text{xc}}/\partial \tau)$, arises from the functional dependence of e_{xc} on τ . The other, $\nabla^2(\partial \tau/\partial \nabla^2 n)$, arises from the dependence of the deorbitalization of τ on $\nabla^2 n$. The effects of the form of $e_{\text{xc}}(\tau)$ upon performance characteristics of both conventional and deorbitalized meta-GGAs has been the subject of recent attention. [6, 7, 25] However, comparison of Fig. 2 to Fig. 3 shows that the location of major error in the hydrogen X potential (the bump in deorbitalized potentials around 2 to 3 au) coincides with that of dramatic noise in the Pauli potential. This implicates, strongly, the second term as a culprit in generating X potential noise, hence that term is a focus of concern. A second benefit of this choice is that it focuses on an aspect of the potential that is independent of the choice of XC functional being deorbitalized.

Minimization of oscillations in that term on a useful test density or densities would have the potential of minimizing its impact on numerical performance in general. Perhaps that should be a design goal for a deorbitalized functional. We will explore that idea in a followup to the present study. For now, to get an efficient, useful estimate of the impact of deorbitalizer-generated noise, we simply evaluate Eq. (41) for the H atom as calculated from a given KED functional. The crucial point is that the exact Pauli potential (Eq. (19) for this case is zero.

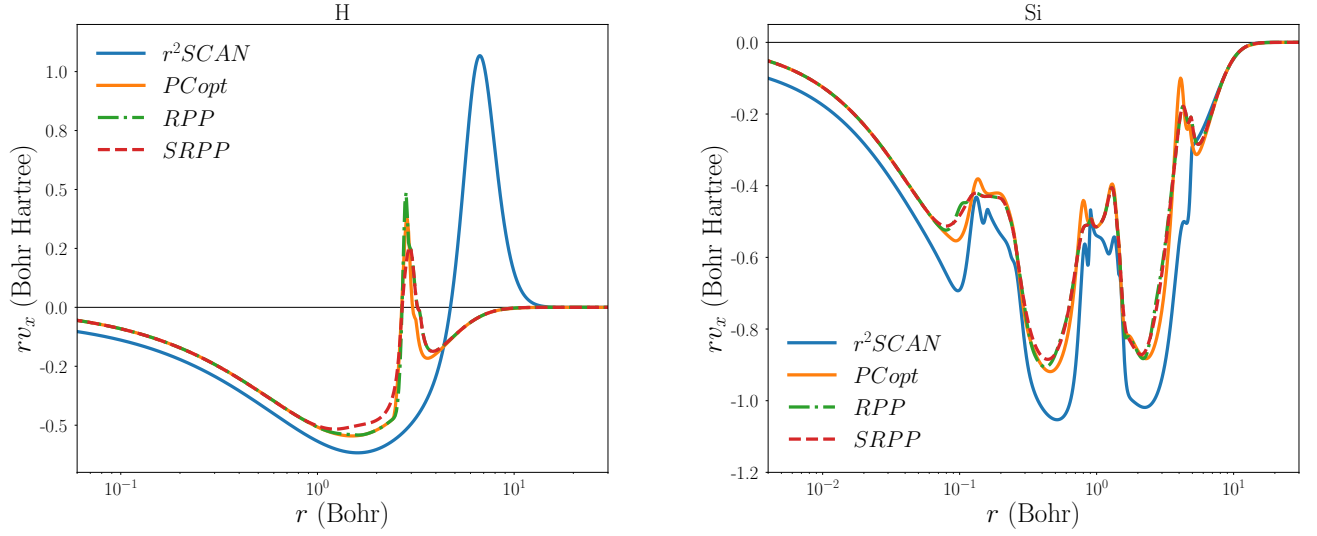


FIG. 3: Local part of the $r^2\text{SCAN}$ exchange potential for the H and Si atoms (scaled by radius r) compared to the exchange potentials of several deorbitalized versions of $r^2\text{SCAN}$ for the same atoms.

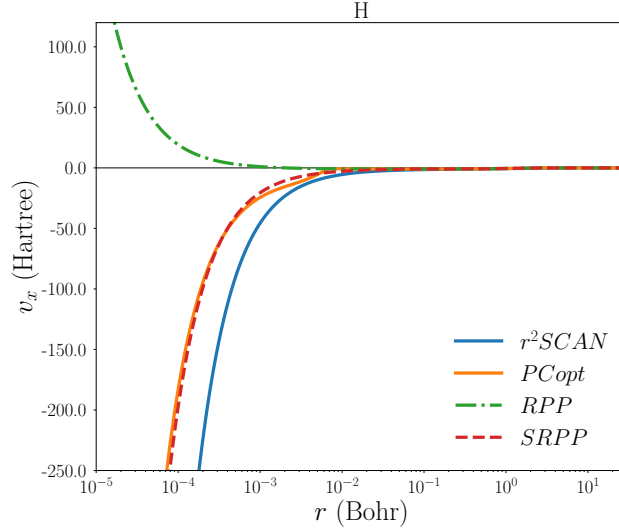


FIG. 4: The local part of the $r^2\text{SCAN}$ X potential for atomic H compared to deorbitalized X potentials, plotted over a much larger range in length and energy.

The consequences of that evaluation are shown, along with the kinetic energy obtained by integrating τ_{OF} , in Table I with the deorbitalizers PC, PC_{opt} , CR, RPP, SRPP, and SRPP2. There is a close correlation between the smoothness of transition between slowly-varying and iso-orbital limits exhibited by the enhancement factors seen in Fig. 1 and the quantified measure I^{deorb} . The switching behavior of PC clearly is the most abrupt. Thus it has an enormous I^{deorb} value. PC_{opt} and RPP do much better, with a reduction in I^{deorb} by a factor of five and an order of magnitude respectively. Clearly, the functionals with the CR switch (CR, SRPP, SRPP2)

are the best. That pattern is manifest to a considerable extent in the corresponding kinetic potentials (Figs. 2) and to a lesser but still significant effect for X potentials, 3). The next issue, obviously, is to address to what extent (if any) this smoothing endeavor and its quantitative assessment has consequences in calculations on real systems.

TABLE I: Kinetic energy and integrated noise measure I^{deorb} evaluated on the Hydrogen atom for several deorbitalized models of the KE density.

Model	Exact	PC	PC _{opt}	RPP	CR	SRPP	SRPP2
KE (ha)	1/2	0.507	0.506	0.504	0.514	0.527	0.550
I (au)	0	220	44.0	19.0	1.705	1.755	1.555

III. COMPUTATIONAL METHODS

For molecular system tests, calculations were done with the NWChem 7.0.2 code [40] using the def2-TZVPP basis set and `xfine` grid settings. Test comparisons are to heats of formation computed according to the now-standard procedure of Curtiss *et al.* [41, 42] for the 223 molecules of the G3X/99 test, to bond lengths for the T96-R test set [43, 44], and to harmonic vibrational frequencies using the T82-F test set [43, 44].

Solid structural properties were calculated with the VASP-5.4.4 code [45], following the same methodology and protocol as in Ref. 46. For hexagonal close-packed structures the ideal c/a ratio was used. For cohesive energies, isolated atom energies were calculated using a $14 \times 15 \times 16$ unit cell and Γ -point sampling. Static-crystal lattice constants and cohesive energies are compared to published results for 55 solids [47] and, correspondingly, for bulk moduli of 44 solids [48]. In addition, band gaps of 21 insulators and semiconductors [49] were computed. Equilibrium lattice constants a_0 and bulk moduli B_0 at $T = 0^\circ K$ were determined by calculating the total energy per unit cell at twelve points in the volume range $V_0 \pm 10\%$, where V_0 is the calculated equilibrium unit cell volume, followed by fitting to the stabilized jellium equation of state (SJEOS) [50]. All the error values are from comparison with tabulated experimental values.

For timing calculations, the same methodology was used except that for the 55 solids [47] test set, the calculations were redone at the experimental lattice constants. In that way only single-point energy evaluations were needed for timings. Calculations used a single node on the University of Florida HiperGator system (Gen. 3) with an AMD EPYC 75F3 32-core processor with 4GB of memory per core. They were executed in parallel across all 32 cores.

IV. RESULTS

A. Structural properties

Evidently the comparison of interest is primarily for r^2 SCAN versus its deorbitalized versions, r^2 SCAN-L, built with the PC_{opt} deorbitalizer [19], with RPP [25], and with SRPP and SRPP2. An underlying issue, discussed in detail above, is that PC_{opt} breaks some constraints that those other three preserve. This motivates further comparison, between PC_{opt} and the three more constraint-compliant deorbitalizers.

Tables II and III compile mean errors (ME) and mean absolute deviations (MADs) for r^2 SCAN, and deorbitalized r^2 SCAN, for the four deorbitalizers, along with results from PBE [51] as a baseline. In addition to these standard measures of functional performance, we include the spread of errors as defined by the maximum (most positive) error minus the minimum (most negative.) This last measure is motivated by the wide variations in timing performance that we discuss below. That variability raises the question whether there might be similar variability in predictive performance.

For molecular systems, the results presented in Table II show that SRPP is a substantial improvement (40% reduction) over RPP for heat of formation MAD. SRPP2 also provides an improvement over RPP, albeit a significantly smaller one. On bond lengths and vibrational frequencies, the three (RPP, SRPP, SRPP2) perform about the same; none proves competitive with PC_{opt}. All three RPP-derived deorbitalizations show a disappointing degradation of performance, especially for heats of formation and frequencies, compared to PC_{opt} deorbitalization, and none can be considered a completely faithful deorbitalization of r^2 SCAN. Also note the near equality of the magnitude of MEs and MADs for SRPP2 and to a lesser extent, SRPP. This, and the fact that the MEs for both are negative, indicate that they tend to underbind consistently, leading to larger MADs.

For solids, Table III shows quite different outcomes. SRPP, SRPP2, and RPP all have distinctively better

TABLE II: Comparison of molecular structural property results for the r^2 SCAN XC functional and various deorbitalized-(r^2 SCAN-L) variants derived from PC_{opt} , RPP, SRPP, and SRPP2 KED functionals. PBE results included for context. Heat of formation mean errors (ME), mean absolute deviation (MADs), and spread in kcal/mol, bond length errors in Å, and frequency errors in cm^{-1} .

		PBE	r^2 SCAN	PC_{opt}	r^2 SCAN-L		
					RPP	$SRPP$	$SRPP2$
Heats of Formation	ME	-20.878	-3.145	1.845	8.796	-5.918	-9.470
	MAD	21.385	4.488	5.300	13.109	7.804	10.405
	Spread	88.091	29.818	42.938	72.117	48.329	46.036
Bonds	ME	0.018	0.005	0.008	0.014	0.011	0.013
	MAD	0.018	0.010	0.011	0.014	0.012	0.014
	Spread	0.168	0.183	0.197	0.182	0.069	0.180
Frequencies	ME	-33.781	11.336	-7.248	-26.743	-22.875	-22.945
	MAD	43.613	30.899	25.709	36.711	36.134	36.800
	Spread	261.16	212.42	201.73	266.59	276.09	268.84

TABLE III: As in Table II for solid-state structural properties. Mean errors (MEs), mean absolute deviations (MADs) and spreads for equilibrium lattice constants in Å, cohesive energies in eV/atom, bulk moduli in GPa, and band gaps in eV.

		PBE	r^2 SCAN	PC_{opt}	r^2 SCAN-L		
					RPP	$SRPP$	$SRPP2$
Lattice constants	ME	0.046	0.026	0.022	0.003	-0.003	-0.002
	MAD	0.053	0.037	0.038	0.029	0.028	0.028
	Spread	0.222	0.311	0.212	0.154	0.143	0.142
Cohesive energies	ME	-0.070	-0.134	-0.327	-0.017	0.051	0.064
	MAD	0.252	0.238	0.349	0.217	0.224	0.227
	Spread	2.096	2.343	1.841	1.726	1.773	1.801
Bulk modulus	ME	9.704	1.367	-4.249	1.084	1.928	2.228
	MAD	11.022	5.963	10.115	8.542	7.866	7.851
	Spread	64.09	63.67	96.90	81.64	76.25	76.25
Band Gaps	ME	-1.69	-1.20	-1.38	-1.60	-1.58	-1.57
	MAD	1.69	1.20	1.38	1.60	1.58	1.57
	Spread	4.89	4.32	4.72	4.99	4.93	4.92

MADs than PC_{opt} for lattice constants, cohesive energies, and bulk moduli. SRPP and SRPP2 are a tiny bit better than RPP on lattice constant MAD, slightly worse on cohesive energy, and about 9% better on bulk modulus. The bulk modulus spread for SRPP and SRPP2 is better than for RPP as well and substantially better than that from PC_{opt} .

Overall, SRPP and SRPP2 preserve the formal properties of RPP and either improve on its performance or maintain it. RPP is already known to be better on metallic solids than PC_{opt} , (and better in some measures than r^2 SCAN) but worse on molecules [25]. That behavior is

confirmed by our results (again, see Tables II and III). Among the three, SRPP and SRPP2 are slightly better than RPP for solids, while SRPP is the best of the three over all systems though not competitive with PC_{opt} for molecules. These outcomes reinforce the finding by Kaplan and Perdew [25] that re-introduction of compliance with constraints broken by PC_{opt} actually can reduce the breadth of applicability of the deorbitalization.

The error spread reduction provided by the RPP-derived models was unexpected. Those spreads are quite a bit better than the original r^2 SCAN for both cohesive energies and lattice constants and not much worse for

bulk moduli. In that regard, PC_{opt} shows some degradation. Similar results are found if we measure standard deviations. That is to say, deorbitalization can, in some cases, lead to a significant reduction in outliers compared to the gKS calculations with the parent metaGGA XC functional, an unexpected bonus. The inverse correlation of spread to our measure of potential smoothness is suggestive. If some of that spread is due to numerical instability rather than functional accuracy, these results would be a trade-off between removing numerical problems with gKS and introducing problems with the Laplacian. The relative absence of outliers leads us to speculate that the noise reduction in smoothed deorbitalized potentials may reduce the risk that geometry optimization procedures will discover spurious local minima.

B. Timing results

Table IV presents timing statistics for the 223 molecule G3X/99 test set. Table V displays the timing data for the six-molecule subset, denoted AE6, of the G3 test set. Each table shows the total time for the run of a test set, and the average time taken per system. In addition, each table shows the average number of SCF cycles needed to converge to self-consistency, (defined as the total number of SCF cycles divided by the number of systems) and the average time per SCF cycle, taken as the average time divided by the average number of cycles. In Table IV, spreads in these last two measures are shown also. Those are defined, as before, as the difference between the maximum and minimum values of a quantity observed across a data set.

Table IV shows that both SRPP and SRPP2 are substantially faster than RPP for the molecules. Similar to the PC_{opt} deorbitalization, neither of them delivers a speed advantage over the parent, orbital-dependent functional. The effects of test-set sampling are shown in Table V. The AE6 timings give a much more favorable comparison of SRPP2 over RPP but no meaningful gain for SRPP2 versus either PC_{opt} or the parent r^2SCAN functional.

Methodological effects are shown in Table VI. It gives the timings for the G3 test with the VASP 5.4.4 calculations done as isolated systems in a large orthorhombic box. Distinct from the solid calculations, the default cut-off energy was set to 600 eV and Γ point sampling was used. Unlike the G3 calculations in NWChem, the VASP tests exhibit a large degradation in performance by both

RPP and SRPP deorbitalizations with respect to the parent functional. Interestingly, the time per SCF cycle for the SRPP deorbitalizer is 1.29 s/cycle in VASP, while in NWChem it is 1.28 s/cycle. However, for the parent functional r^2SCAN , VASP takes 2.87 s/cycle compared to 1.34 s/cycle in NWChem. Unfortunately, the number of SCF cycles is much higher for the deorbitalized forms, roughly a factor of 2.5 for SRPP and a factor of 3 for RPP. In NWChem the average number of SCF cycles is roughly the same for all three. The speed advantage per cycle of the deorbitalized forms is lost thereby in the quasi-molecular case. This brings to light the drastic effect basis set methodology (plane-wave PAW versus gaussian) can have on the performance of the deorbitalized forms.

For solids, Table VII presents a clearly different story. Both SRPP and SRPP2 outperform the parent, orbital-dependent functional on total time. SRPP2 in particular needs only about 75% of the time of r^2SCAN . Notably, SRPP2 also requires only about 72% more time than PBE. This overall gain compared to the PC_{opt} deorbitalization comes from a drastic reduction in the number of SCF cycles needed by SRPP and SRPP2, about half of the number required by PC_{opt} . Importantly, the times per SCF cycle for SRPP and SRPP2 are essentially the same as for RPP, less than half that for r^2SCAN , and actually faster than even for PBE. The combined result is an overall speedup.

The required number of cycles follows, reasonably closely, the measure of smoothness in the Laplacian contributions in the exchange potential (Table I), hence strongly implicates those terms in the slow SCF convergence performance seen in some of the deorbitalizations. We surmise that the sensitivity of such terms to rather small changes in the density makes achievement of a self-consistent density harder than in gKS or GGA calculations. This diagnosis is supported in the extreme spread of numbers of cycles for deorbitalized functionals compared to the parent functional as evaluated via gKS. The spread in observed cycle counts for r^2SCAN (gKS) is only slightly more than twice the average number of cycles, while for the deorbitalized forms (KS), the spreads range from 4.4 to 6.0, with SRPP2 the best performer.

More insight can be found by visualizing the timings for the individual members of the solid test set. These are shown as average time per SCF cycle in Fig. 5a and number of SCF cycles in Fig. 5b. Results for r^2SCAN itself as well as versions deorbitalized with PC_{opt} , with

TABLE IV: Timings and number of SCF cycles required for computing the 223 molecules of the G3X/99 test set. Time measurements are expressed in seconds (*s*).

223 molecules	r ² SCAN	PC_{opt}	r ² SCAN-L			PBE
			<i>RPP</i>	<i>SRPP</i>	<i>SRPP2</i>	
Total time	2432.00	2410.90	3036.90	2433.60	2373.70	1417.70
Average time	10.91	10.81	13.62	10.91	10.64	6.36
Avg. number SCF cycles	7.79	8.07	8.42	8.19	8.28	8.45
Spread	23	24	24	24	24	22
Avg. time per SCF cycle	1.34	1.28	1.56	1.28	1.23	0.72
Spread	6.27	6.25	7.38	6.14	6.13	4.10

TABLE V: As in Table IV for the AE6 test set.

AE6	r ² SCAN	PC_{opt}	r ² SCAN-L		
			<i>RPP</i>	<i>SRPP</i>	<i>SRPP2</i>
Total time	25.40	25.40	35.00	27.20	26.00
Average time	4.23	4.23	5.83	4.53	4.33
Avg. number SCF cycles	6.50	6.33	8.33	7.17	7.17
Avg. time per SCF cycle	0.66	0.67	0.76	0.65	0.62

RPP, and with SRPP are shown. The distribution for SRPP2 is only modestly better than SRPP, so it was omitted for clarity.

Fig. 5a shows the clear SCF cycle-time advantage of the deorbitalized functionals versus their orbital-dependent parent, including the nearly 3 times faster performance of PC_{opt} originally reported by M-RT. The other deorbitalizers are slightly less swift but still well above twice as fast as r²SCAN per cycle. Notably, the issue is one of outliers. For about half the data set, the r²SCAN (gKS) time is under 5 s and not quite two times slower than the deorbitalized forms (with KS potentials). But there are multiple bad actors in the orbital-dependent case. For them the slowdown is much worse. Deorbitalized functionals do poorly on a much smaller data set of nine transition metals, all with nearly empty *d* shells, e.g. Y or Sc, or nearly empty spin subshells, like Fe.

Fig. 5b shows the number of SCF cycles needed for r²SCAN and the three deorbitalized versions considered in the preceding plot. One sees clearly how the SCF process for original r²SCAN converges in consistently fewer cycles than for any of the three deorbitalized variants. The RPP-type deorbitalizers perform quite reasonably but with a number of outliers. In contrast PC_{opt} has

a rather larger number of outliers which cancel its edge in the time-per-cycles metric. Note also the scatter plot of SCF cycles needed versus the r²SCAN requirement in Fig. 6. The smoothed SRPP and SRPP2 do provide consistent improvement upon the performance of RPP, with a few exceptions among the semiconductors and simple metals.

In sum, while there will be remaining issues in achieving self-consistency with Laplacian-dependent functionals, it is clear that such issues are ameliorated significantly by having a smoother potential.

C. Detailed analysis of computational parameter dependence

Given the importance of materials calculations with plane-wave basis codes such as VASP and the significant performance differences between molecules and solids for such codes, investigation of computational technique and parameter choice effects is imperative. We pursue that in two ways. In this subsection we examine the dependence of timing results on various run parameter choices. In the following subsection we consider the equation of state (cold curve) of Al and a short AIMD calculation for it.

Table VIII provides timing information for r²SCAN

TABLE VI: As in Table IV for the 223 molecules, but with the calculations done in VASP 5.4.4. for the deorbitalized functionals RPP and SRPP as compared to the parent functional.

223 molecules	r ² SCAN	r ² SCAN-L	
		<i>RPP</i>	<i>SRPP</i>
Total time	11100.38	13573.66	13030.01
Average time	49.78	60.87	58.43
Average SCF cycles	17.22	51.90	43.14
Average time per SCF cycle	2.87	1.10	1.29

TABLE VII: Timings and number of SCF cycles required for computing the 55 solids. Times are expressed in seconds (s).

55 solids	r ² SCAN	<i>PC</i> _{opt}	r ² SCAN-L			PBE
			<i>RPP</i>	<i>SRPP</i>	<i>SRPP2</i>	
Total time	6052.81	7812.30	6024.63	4892.41	4553.67	2650.44
Average time	110.05	142.04	109.54	88.95	82.79	48.19
Avg. number SCF cycles	15.64	60.31	40.35	30.49	31.42	14.20
Spread	32	341	242	176	138	19
Ave. time per SCF cycle	6.93	2.56	2.91	2.97	2.89	3.26
Spread	22.39	7.87	7.30	6.97	7.46	7.77

versus r²SCAN-L deorbitalized via RPP and SRPP for the 55 solid test set. It shows the effects of differing choices of the energy cutoff, convergence parameter, and minimization algorithm. Cases labeled “0” through “3” successively reduce the convergence tolerances while keeping the SCF minimization procedure fixed as the preconditioned conjugate gradient method. That is the recommended option for meta-GGA exchange-correlation functionals. Cases 4, 5, and 6 explore effects of the use of less-expensive diagonalization algorithms.

Note that for all cases there is a significant advantage for the deorbitalized forms with respect to the parent functional for the time taken by a single calculation of electronic orbitals. The speed-up stays a little over a factor of two, maybe better for the fastest eigensolver.

In contrast, the advantage in terms of the number of cycles needed to reach self-consistency for the gKS procedure used with r²SCAN over the deorbitalized forms and their pure KS potential does vary quite a bit depending on the tightness of convergence criteria. gKS loses its advantage relative to deorbitalized KS as convergence is loosened and SRPP loses its advantage relative to RPP. Ultimately, for Case 3, the sloppiest one for preconditioned conjugate gradients and Case 6, the sloppi-

est overall, there is basically no difference in performance among the three strategies in this regard. Therefore the total time speed-up of deorbitalization is quite dramatic, almost a factor of 2. (It may seem peculiar that all functionals in Case 6 take substantially more cycles to achieve self-consistency than Case 3, with both larger plane-wave cutoff and stricter convergence tolerance. This is a result of the instability of the RMS-DIIS, which makes sacrifices in stability to achieve faster overall times [52].

There is a very plausible, simple explanation for such a finding. Reducing the cut-off energy amounts to introducing a low-pass filter that eliminates a good deal of noise in the potential. Though the RPP and SRPP potentials have very different noise characteristics, it is reasonable to suppose that those differences are substantially removed by smoothing with a low energy cutoff. Similarly, a less demanding tolerance means greater acceptance of effects of noise in the potential, reducing their consequences. We hypothesize that the smooth atomic-like basis used in molecular calculations is coarse relative to individual plane waves, therefore causing a similar noise reduction in the potential and yielding the “egalitarian” timing results seen for these calculations.

TABLE VIII: Parameter and SCF algorithm dependence of timing for computing the set of 55 solids, for r²SCAN itself, and r²SCAN-L deorbitalized with RPP or SRPP. E_{cut} is the plane-wave cut-off energy; E_{diff} is the energy tolerance used to end the self-consistency cycle. The algorithms to solve the KS or gKS equation, using the keyword terminology for the ALGO input parameter in VASP, are: (All) preconditioned conjugate gradient, (Normal) blocked Davidson iteration, (Fast) hybrid of block-Davidson and RMM-DIIS (Residual Minimization Method with Direct Inversion in the Iterative Subspace), and (Very Fast) RMM-DIIS.

	Case 0	Case 1	Case 2	Case 3	Case 4	Case 5	Case 6
Algorithm	All	All	All	All	Normal	Fast	Very Fast
E_{diff} (eV)	1×10^{-6}	2.72×10^{-5}	1×10^{-6}	2.72×10^{-5}	1×10^{-6}	1×10^{-6}	1×10^{-4}
E_{cut} (eV)	800	800	600	600	800	800	500
<i>Average time</i>							
r ² SCAN	110.05	101.28	95.51	81.47	112.26	104.26	58.37
RPP	109.54	63.78	55.23	42.02	103.97	63.11	22.66
SRPP	88.95	59.11	55.56	45.56	111.93	79.25	23.62
<i>Average SCF cycles</i>							
r ² SCAN	15.64	12.67	15.78	12.78	12.91	12.93	17.36
RPP	40.35	20.93	20.11	13.75	29.96	27.33	18.51
SRPP	30.49	17.55	17.49	13.69	29.09	30.80	18.75
<i>Average time/cycle</i>							
r ² SCAN	6.93	7.74	5.96	6.16	8.59	7.91	3.34
RPP	2.91	3.14	2.77	2.98	3.73	2.45	1.21
SRPP	2.97	3.32	3.03	3.21	4.11	2.83	1.24

D. Molecular dynamics of Aluminum

For context, in Table IX we provide the timing and SCF count for calculating the static lattice fcc Al equation of state at zero temperature. The data are averages over runs at 12 lattice parameters, $3.8484 \text{ \AA} \leq a_0 \leq 4.1146 \text{ \AA}$. (At each lattice constant, the calculation was started from the same density, not from the equilibrium density of the preceding lattice constant. This technical aspect is of importance later.)

Solid Al is a particularly advantageous case for deorbitalization. Consistent with the discussion above about the 55 solid test set (recall Table VII), all three deorbitalizations (PC_{opt}, RPP, SRPP) are substantially faster in total time than the parent functional. The spreads are even better for RPP and PC_{opt}, and as good as the parent for SRPP. Also as expected, RPP and SRPP outperform PC_{opt} substantially with regard to the required number of SCF cycles.

Those rather encouraging timing and cycle count results would lead to the expectation that the SRPP and RPP functionals would outperform the parent in driving AIMD. To investigate that, we obtained timing statistics for a short AIMD simulation of liquid-phase aluminum.

The simulation system consisted of 108 atoms, at bulk density $\rho = 2.34 \text{ g/cm}^3$ in a $12.7239 \times 12.7239 \times 12.7239 \text{ \AA}^3$ cell. The Γ -point was used to sample the Brill-

ouin zone. In VASP 5.4.4, the system was treated with a three-electron pseudopotential ($3s^2 3p^1$ valence, 10 electrons in the core; denoted PAW_PBE Al_GW). Non-spherical contributions within the PAW spheres were included self-consistently. Minimization used the RMM-DIIS algorithm with energy cutoff set to 500 eV, as in Case 6 in Table VIII. The self-consistent energy convergence tolerance E_{diff} was $1e^{-4}$ eV. Approximate Fermi-type thermal smearing with a width of 0.0881555 eV was applied, and the initial temperature set to 1023 K. Molecular dynamics used the Verlet algorithm and a Nosé thermostat. 6501 steps of time-step 0.941 femtoseconds were taken for each simulation. Each calculation utilized 5 nodes on the University of Florida HiperGator cluster (Gen. 3), with 64 cores per node (total of 320 cores) and 4 GB of memory per core (1280 GB total).

Fig. 7 gives the resulting timing statistics. Total times in hours are in the upper panel, Fig. 7a, along with the total number of SCF cycles. The time per SCF cycle is shown in the lower panel Fig. 7b. The outcome is that all of the deorbitalized functionals are at a disadvantage with respect to the parent functional r²SCAN for total run time. This is in contrast to what one would have expected from the fcc Al equation of state results just discussed.

Clearly, the problem is not in the time per SCF cycle. As with the Al equation of state, the best performers on

TABLE IX: Timings (in seconds, s) and number of SCF cycles required for computing the zero-temperature fcc Al static lattice equation of state for r²SCAN and three deorbitalizations, with PBE values for context.

fcc Al	r ² SCAN	r ² SCAN-L			PBE
		PC_{opt}	RPP	$SRPP$	
Total time	990.89	611.87	366.93	415.89	296.04
Average time	82.57	50.99	30.58	34.66	24.67
Avg. number SCF cycles	26.75	45.00	17.58	20.83	12.42
spread	23.00	51.00	9.00	19.00	4.00
Avg. time per SCF cycle	3.12	1.16	1.75	1.68	2.00
spread	0.42	0.32	0.21	0.41	0.30

a per-cycle basis are the deorbitalized functionals, particularly RPP and SRPP. They out-perform even PBE. But for the AIMD, the number of SCF cycles needed for those deorbitalized functionals is almost prohibitively high, over four times the number needed for the parent functional, and over five times that needed for PBE.

This excessive SCF cycle count also is in marked contradiction to our findings for the 55 solids. Recall Table VIII. It shows that for the lowest accuracy tolerances in cutoff energy and SCF convergence (Case 6), the same as employed in the AIMD, the number of cycles needed for the deorbitalized functionals and the parent functional are roughly the same. That clearly is not the case in the AIMD driven by any of the deorbitalizers.

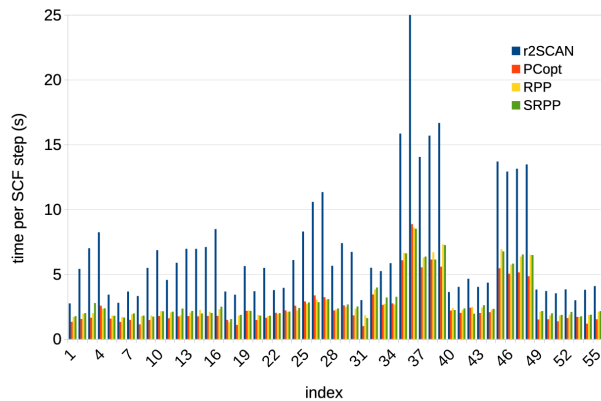
One way to characterize this problematic cycle count is that convergence to self-consistency from the fully converged density of the previous MD step is not that much quicker than convergence from an LCAO density (12 cycles average per MD step for SRPP, in contrast to 14 cycles average over the test set and 18 to 21 for the equation of state calculation for Al). The behavior also is in contrast with the dramatic reduction of PBE and r²SCAN requirements to less than 3 cycles average per step. This may be an unpleasant consequence of the inclusion of the density Laplacian in the X functional and associated sensitivity to seemingly small changes in the density. A peculiarity is that a tolerance of $E_{\text{diff}} = 2.75 \times 10^{-5}$ needs an average of 14 cycles to converge in the case of the 55 element test set (see Table VIII) while, on the other hand, use of $E_{\text{diff}} = 1 \times 10^{-4}$ in the MD needs 18 cycles. The difficulty is that though RMS-DIIS is extremely fast it is less stable than the other solvers.

We have pursued two diagnostic follow-ups which indicate that the issue is quite complex and not readily resolvable. First, altering the MD timestep should pro-

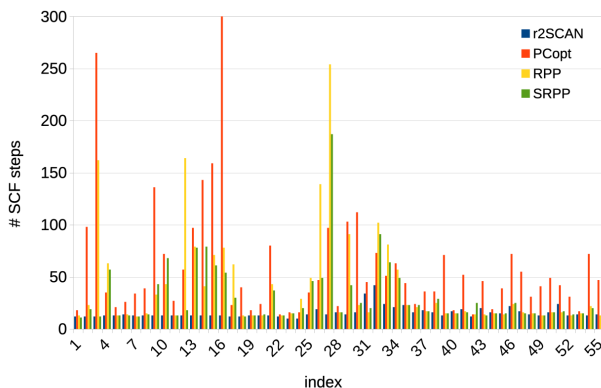
vide a quick consistency check regarding the behavior. We find remarkably that the convergence issues, if anything, *increase* as the timestep is decreased. See Table X. Since this obviously cannot be the case in the limit of zero time-step, something peculiar is happening but we have been unable to identify the cause.

Secondly, investigation of the number of SCF cycles needed as a function of simulation time shows a rather intriguing trend. The RPP and SRPP deorbitalized functionals start out with convergence times that are quite competitive to r²SCAN and PBE. Only gradually, over about 10 MD steps, do they settle down to the more slowly converging behavior. The slowdown in convergence is correlated to increased fluctuations in convergence performance, symptomatic of an unresolved stability issue. This behavior is documented in Fig. 8. Perhaps it is noteworthy that, of the three deorbitalizers, the one with smoothest potential, SRPP has the smallest fluctuations in cycles-per-step. We have sought diagnostic insight from calculating mean-square-displacements and radial distribution functions for both T=1093K and 298K AIMD but without much gain in understanding. Plots of the results are in the Supplemental Information.

The behavior suggests that tendencies to instability in the deorbitalized forms take some SCF cycles to show up and might be ameliorated by taking an occasional step using, e.g. PBE, to find the density. We tried one such strategy. At each MD step, the first five SCF cycles were done with PBE, then continued with the selected deorbitalized functional. Unfortunately, this strategy actually increased the number of SCF cycles instead of reducing them.



(a) Average time per SCF cycle in seconds.



(b) Number of SCF cycles.

FIG. 5: Bar charts showing timing performance for r^2 SCAN, and r^2 SCAN-L with the PC_{opt} , RPP, and SRPP deorbitalizers for the entries in the 55-solid test set. Indexing of materials: 1-4 elemental semiconductors; 5-16: compound semiconductors; 17-22: ionic compounds; 23-31: simple metals; 32-55 transition metals.

V. DISCUSSION AND CONCLUDING REMARKS

We have constructed a measure, I^{deorb} , to control instability sources in the XC potential of deorbitalized metaGGAs, in terms of a metric derived from the Pauli potential (the functional derivative of the deorbitalized kinetic energy density with respect to the density). This measure clearly diagnoses the relative instability or reliability of performance of standard deorbitalizers and helps motivate a modified form of RPP, the SRPP and SRPP2.

In consequence, SRPP and SRPP2 produce noticeably smoother kinetic and exchange potentials than other deorbitalizers. Both modified forms preserve or modestly enhance the predictive performance of RPP in the de-

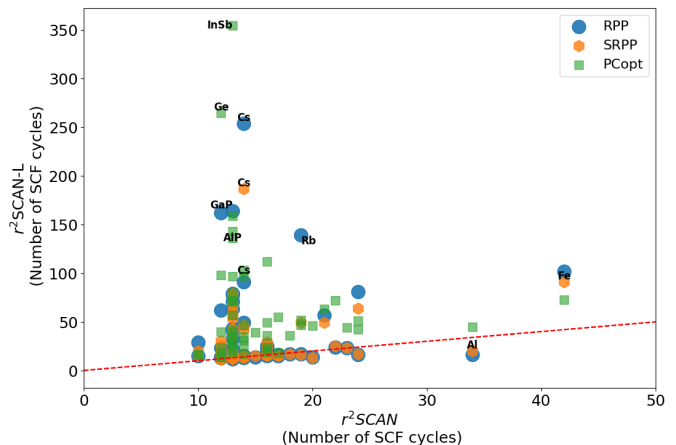
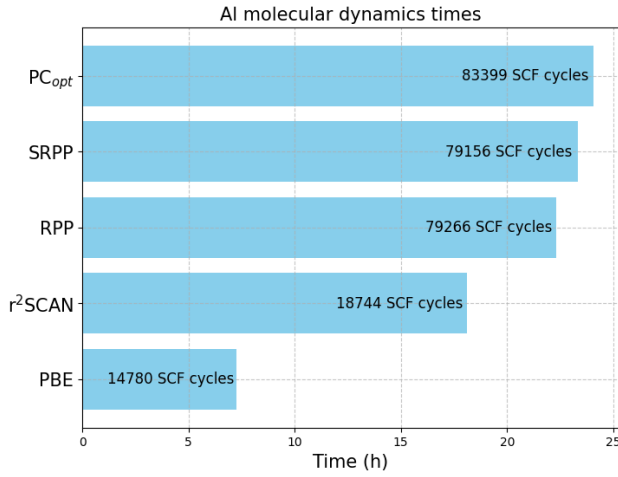


FIG. 6: Scatter plot showing the number of SCF cycles for 55 solids calculated using the deorbitalized functionals relative to the number of cycles required with the r^2 SCAN functional. The red dotted line is the locus of values for the deorbitalized functionals that are equal to the r^2 SCAN values.

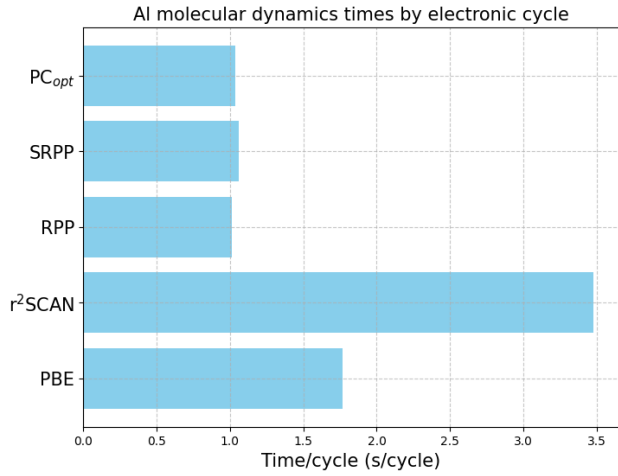
orbitalization of the r^2 SCAN functional on a standard test set for the structural properties of solids. But for the smoothed functionals, this constraint compliance for slowly varying densities comes with a much smaller penalty in predictive performance for highly inhomogeneous systems than one might expect from the behavior of the RPP. The smoothed functionals improve upon RPP markedly for the G3 molecular test set, though the deorbitalized form still is not competitive with r^2 SCAN-L(PC_{opt}) for heats of formation. The SRPP2 variant, though less effective than SRPP on the molecular cases, is equal or better on the solids and delivers better timing performance. It is plausible that this gain is a consequence of the SRPP2 X potential being smoother than the SRPP one.

The implications for timing performance are complicated and unfortunately somewhat obscure. For molecules, both the time per SCF cycle and the number of cycles needed to reach convergence are essentially the same for both the parent and deorbitalized functionals.

For fixed (static) solid geometries, the story is different. Unlike earlier deorbitalized functionals, these particular deorbitalized metaGGAs significantly outperform the explicitly orbital dependent, metaGGA parent (used in gKS form) so far as time per SCF cycle is concerned. While the deorbitalized forms (r^2 SCAN-L with L either SRPP or SRPP2) need more cycles to reach self-consistent convergence, the increment is not so large as to offset the much shorter cycle time. As a result, SRPP and SRPP2



(a) Bar chart showing total run times in hours; total number of SCF cycles taken also noted.



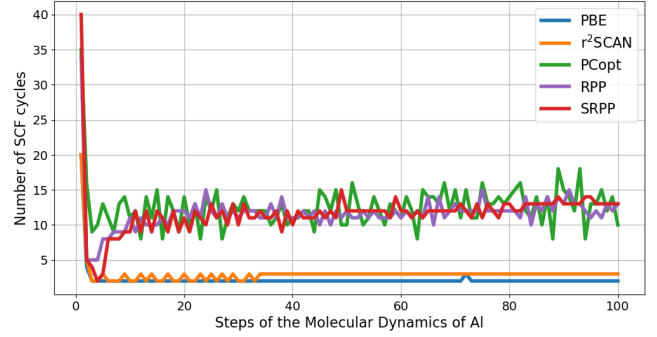
(b) Time per electronic cycle, in seconds.

FIG. 7: Timing statistics for 6,501 step AIMD simulation of liquid Al, using PBE, r²SCAN, and r²SCAN-L with several deorbitalization strategies.

are much faster than r²SCAN for static lattice solid calculations, as seen in Tables VII, VIII and IX.

That advantage does not propagate into the one AIMD example we have tried. It appears that the density Laplacian dependence in the deorbitalized functionals introduces instabilities that greatly increase the number of SCF cycles needed in the AIMD context. Our speculation has been that since the XC potential is sensitive to small perturbations in the density because of the Laplacian, potentials for successive AIMD steps may occasionally be different enough to lead to unusual variation in the density, and thus slower SCF convergence. However, the limited testing we have done with reduced AIMD step-time is not consistent with this argument. Similarly, one

FIG. 8: Number of SCF cycles to convergence per Molecular Dynamics step for different approximations, plotted for the first 100 MD time steps.



might think that there could be a clue as to the cause of this behavior in the average SCF cycle time versus its spread. But Table VII shows that SRPP, SRPP2, and RPP itself all have similar ratios of spread of time per cycle to average time per cycle. In fact, the ratio for SRPP (2.35) is slightly below that for RPP (2.51).

In the case of plane-wave pseudopotential calculations of solids, we see two outcomes for equation-of-state calculations:

1. For high accuracy calculations, the performance of deorbitalized metaGGA XC functionals relative to the parent is markedly dependent on the smoothness of the the XC potential, with smoothed functionals (SRPP and SRPP2) significantly outperforming not only the earlier deorbitalizers (e.g. PC_{opt}), but the parent orbital-dependent functional itself. In that regard, this work has achieved one of the long-standing goals of the deorbitalization endeavor.
2. Reducing accuracy requirements via either lowered energy cut-off for orbital representations or the energy tolerance for achieving self-consistency, causes all the metaGGA forms (orbital or deorbitalized) to reach rough parity of the number of cycles needed to achieving self-consistency. The orbital-dependent gKS calculations continue to be inferior compared to pure KS from the deorbitalized functionals on a per-cycle basis. But the relative advantage of SRPP and SRPP2 versus the other deorbitalizers is largely lost because lowering the cutoff effectively smooths all potentials to the same degree, while reducing the SCF tolerance reduces the sensitivity of the calculation to instabilities from

the potential, so the wall-clock times for all tend to converge.

Because Gaussian basis functions are themselves smooth and are equivalent to fixed combinations of many plane waves, molecules calculated with them behave, oddly enough, like the cruder and faster plane-wave calculations, with little distinction in timing among forms of deorbitalizers. Apparently, the nature of limiting the basis has a counterpart smoothing, hence stabilizing effect as does reducing the energy cutoff in a plane-wave code.

AIMD calculations provide a third, somewhat discomfiting pattern of behavior. As for the case of lower accuracy calculations for the equation of state, deorbitalized functions outperform the orbital-dependent parent on a per-cycle basis, while smoothed and unsmoothed deorbitalizers have similar SCF convergence rates. But PBE and r²SCAN perform the task specific to MD, namely finding orbitals for a new set of nuclear positions starting from the density for a modestly different set of positions, in two or three SCF cycles. In contrast, the deorbitalized forms used here take almost the same number of cycles as they do when starting from, say, an LCAO density. An attempt to start at each position with a PBE density did not help.

One ansatz for AIMD that remains to be investigated would be to take the density from some numerically stable method (PBE to keep it simple), and evaluate the total energy and forces with a single *non*-self-consistent calculation with the metaGGA-L functional of interest. For the comparatively coarse convergence levels required, that should give nearly the same forces (and energies) as from the conventional self-consistent metaGGA-L, and better than those from self-consistent PBE, at a fraction of the timing cost of self-consistent metaGGA-L.) Doing so would combine best-time performance vis-a-vis finding the density at any step with something that is fast (for a single SCF step) and accurate to calculate forces.

As another line of pursuit, it may be advantageous to take the measure of noise defined in this paper and use it as an aid to construct deorbitalizers with yet smoother potentials, either by optimizing the switching function between iso-orbital and slowly-varying limits or strategically modifying "appropriate norms" in the deorbitalizer or parent functional. This is the focus of ongoing research.

SUPPLEMENTARY MATERIAL

The following file is available free of charge.

- SRPP_Timing.SuppInfo: In this file, we provide a detailed, system-by-system tabulation of the numerical results of the test calculations against standard molecular and solid test sets. We also provide system-by-system SCF timing and SCF cycle count tabulations, corresponding to the plot of SCF cycle count for each of the first 100 AIMD steps in Fig. 8 above.
- Plots of mean-squared displacements and radial distribution functions of Al from the various XC functionals driving AIMD at T=298K and 1023K.

ACKNOWLEDGMENTS

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TABLE X: Total times for the molecular dynamics of Al using the SRPP deorbitalizer for different values of timestep.

XC functional	timestep (POTIM)	Total time (s)	Number of SCF cycles	Time by cycle (s)	Average Temperature (K)
r ² SCAN-L(SRPP)	1.4	87897.82	78872	1.11	1023.021
	0.940793	83935.23	79156	1.06	1022.869
	0.5	95314.36	86477	1.10	1022.973
r ² SCAN-L(RPP)	0.940793	80395.09	79266	1.01	1022.995
	0.5	86879.67	87786	0.99	1023.052
r ² SCAN	0.940793	65212.99	18744	3.48	1023.058
	0.5	54600.70	13554	4.03	1023.056