Laplacian-level kinetic energy approximations based on the fourth-order gradient expansion: Global assessment and application to the subsystem formulation of density functional theory

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We test Laplacian-level meta-generalized gradient approximation (meta-GGA) non-interacting kinetic energy functionals based on the fourth-order gradient expansion (GE4). We consider several well known Laplacian-level meta-GGAs from literature (bare GE4, modified GE4, and the MGGA functional of Perdew and Constantin [Phys. Rev. B 75,155109 (2007)], as well as two newly designed Laplacian-level kinetic energy functionals (named L0.4 and L0.6).

First, a general assessment of the different functionals is performed, testing them for model systems (one-electron densities, Hooke’s atom and different jellium systems), atomic and molecular kinetic energies as well as for their behavior with respect to density-scaling transformations. Finally, we assess, for the first time, the performance of the different functionals for Subsystem Density Functional Theory (DFT) calculations on non-covalently interacting systems.

We find that the different Laplacian-level meta-GGA kinetic functionals may improve the description of different properties of electronic systems but no clear overall advantage is found over the best GGA functionals. Concerning Subsystem DFT calculations, the here proposed L0.4 kinetic energy functional is competitive with state-of-the-art GGAs, whereas all other Laplacian-level functionals fail badly. The performance of the Laplacian-level functionals is rationalized thanks to a two-dimensional reduced-gradient and reduced-Laplacian decomposition of the non-additive kinetic energy density.

I. INTRODUCTION

The non-interacting kinetic energy (KE) density-functional has been the subject of intense research in electronic structure theory for almost one century, since the introduction of the Thomas-Fermi model. This interest has been further motivated, and theoretically justified, by the introduction of density functional theory (DFT) which has the non-interacting KE as a main ingredient. In the orbital-free formulation of DFT the non-interacting KE density-functional is, in fact, the main contribution to the electronic energy which has no explicit analytical expression in terms of the density (the other term being the exchange-correlation (XC) functional). Despite the effort spent on it, however, the development of accurate non-interacting KE functionals resulted to be an extremely difficult task. As a consequence, orbital-free DFT is still of limited practical utility, showing reasonable accuracy only for some solid-state applications. Recently, the density-decomposed orbital-free DFT, that treats differently the localized and delocalized densities, seems to bring further progress in the field. We also acknowledge that the possibility of reaching chemical accuracy within orbital-free DFT calculations was proved recently, at least for one-dimensional systems, by calculations employing machine learning techniques to approximate the non-interacting kinetic energy (with $\sim 10^5$ parameters). This study showed however also the extreme difficulty of this problem.

On the other hand, in recent years, the interest in KE functionals was strongly renewed by the development of density embedding methods, where a many electron system with electron density $n(r)$ is partitioned into two subsystems A and B, such that the total electron density is $n = n_A + n_B$, and the mutual interaction is accounted for by an appropriate embedding potential. Of special relevance in this context is the subsystem formulation of DFT within the Kohn-Sham formalism. In the standard formulation of this method the system is described by two coupled sets of Kohn-Sham equations with constrained electron density (KSCED). Hence, the density is constrained to satisfy the condition $n = n_A + n_B$ by the inclusion of an external embedding potential of the form (here the embedding potential for subsystem A is reported; a similar expression holds for
\[ v_{\text{emb}}^A[n_A; n_B](\mathbf{r}) = v_{\text{ext}}^A(\mathbf{r}) + v_J[n_B](\mathbf{r}) + \frac{\delta E_{\text{exc}}^{\text{add}}[n_A; n_B]}{\delta n_A(\mathbf{r})} + \frac{\delta T_s^{\text{add}}[n_A; n_B]}{\delta n_A(\mathbf{r})}, \]

where \( v_{\text{ext}}^A \) and \( v_J[n_B](\mathbf{r}) \) are the external (i.e. nuclear) and the Coulomb potentials due to subsystem B, while the non-additive XC and kinetic energy terms are defined as

\[ E_{\text{xc}}^{\text{add}}[n_A; n_B] = E_{\text{xc}}[n_A + n_B] - E_{\text{xc}}[n_A] - E_{\text{xc}}[n_B], \]

\[ T_s^{\text{add}}[n_A; n_B] = T_s[n_A + n_B] - T_s[n_A] - T_s[n_B]. \]

Using an iterative freeze-and-thaw procedure\(^{83,92}\) the full variational solution for the total system can be obtained, which is equivalent to the usual Kohn-Sham solution, except for approximations included in the non-additive kinetic interaction term and eventually in the non-additive XC contribution, if hybrid or orbital-dependent functionals are employed in the subsystem formalism.\(^{89,90,93,94}\) Henceforth, the acronym FDE will be used to refer to this fully variational approach.

As shown by Eqs. (1) and (2), the accuracy of the FDE method relies on the availability of accurate kinetic energy approximations. However, unlike for orbital-free DFT, the FDE approach makes use not of the bare non-interacting KE, but rather of the non-additive KE contribution (the remaining part of the kinetic energy, i.e. the subsystems’ KE, is treated in a Kohn-Sham fashion within the KS-CED equations\(^{43,44}\)). For non-bonded interactions, the non-additive KE is quite small and well-behaved, so that, in analogy to the XC energy, it can be efficiently described by semilocal approximations. Thus, for a broad range of problems (e.g. hydrogen bonds, dipole-dipole, and dispersion complexes) the FDE method can reach a high performance, often below the chemical accuracy.\(^{62,64}\)

Motivated by the practical appeal of the FDE method, in the last years different semilocal KE approximations were proposed to describe the non-additive KE bifunctional.\(^{36,37,61,63,65}\) However, all these approximations are making use only of the simplest semilocal ingredients, i.e. the density \( n \) and its gradient \( \nabla n \), being based on the generalized gradient approximation (GGA). Comparison with the experience accumulated for the much widely investigated XC functional, shows nevertheless, that the GGA level shows some inevitable limitations due to its intrinsic simplicity, and in particular cannot properly distinguish between different density regimes.\(^{63,64}\)

Thus, the investigation of the performance of more sophisticated functionals beyond the GGA level (meta-GGA functionals) in the context of the FDE method is of high interest. To date, however, to our knowledge, no such study has ever been performed. For this reason this work has as principal goal to perform a general assessment of some existing\(^{23,55}\) and new Laplacian-level meta-GGA KE functionals in the context of the FDE method. This study will be performed by first testing the general quality of the KE functionals on a wide set of systems and properties. Then, direct application of the KE functionals in FDE calculations will be considered.

At this point it is important, however, to note that most meta-GGA XC functionals are implemented using as additional ingredient to the GGA ones, the positive-defined kinetic energy density \( \tau = \frac{1}{2} \sum_i |\nabla \phi_i|^2 \), where \( \phi_i \) are the occupied Kohn-Sham orbitals. The Laplacian of the density instead is not used directly, but mimicked in the atomic core through a function depending on \( n \), \( \nabla n \), and \( \tau \).\(^{96,97}\) This choice is convenient, because the positive-defined kinetic energy density has a more regular behavior than \( \nabla^2 n \), which oscillates and diverges near the atomic nucleus. At the same time the use of \( \tau \) causes the so constructed meta-GGA functionals to have a non-local dependence on the density (via the orbital-dependent \( \tau \)).

The focus in this paper will be on meta-GGA KE functionals using the Laplacian of the density as meta-GGA ingredient. In this way, despite some possible limitations due to the behavior of \( \nabla^2 n \), it is possible to construct a truly semilocal KE functional, suitable to be used in the FDE formalism, and having meta-GGA quality. We recall in fact that \( \nabla^2 n \) is an important ingredient for the construction of functionals and enters in the definition of the fourth- and higher-order gradient expansion of the exact kinetic energy.\(^{64}\)

II. KINETIC ENERGY FUNCTIONALS

A Laplacian-level semilocal KE functional has the general form

\[ T_s[n] = \int d\mathbf{r} \, \tau^{TF} F_s(n, \nabla n, \nabla^2 n), \]

where \( \tau^{TF} = \frac{3}{10} (3\pi^2)^{2/3} n^{5/3} \) is the Thomas-Fermi kinetic energy density\(^{23} \) and \( F_s \) is a suitable kinetic enhancement factor. Under a uniform scaling of the density \( (n_A(\mathbf{r}) = \lambda^3 n(\lambda \mathbf{r}), \lambda \geq 0) \), the exact non-interacting kinetic energy behaves as \( T_s[n_A] = \lambda^2 T_s[n] \), i.e. as the Thomas-Fermi KE. Therefore, to have Eq. (4) satisfying this constraint, \( F_s(n, \nabla n, \nabla^2 n, ...) \) must be invariant under the uniform density scaling. Such a goal can be achieved by considering the following dimensionless reduced gradient and Laplacian

\[ p = \frac{|\nabla n|^2}{4(3\pi^2)^{2/3} n^{8/3}} ; \quad q = \frac{\nabla^2 n}{4(3\pi^2)^{2/3} n^{5/3}}. \]

The enhancement factor becomes therefore

\[ F_s(n, \nabla n, \nabla^2 n) = F_s(p, q). \]

In this paper we consider the following approximations for the kinetic enhancement factor:

(i) Thomas-Fermi (TF), defined as

\[ F_s^{TF} = 1 + aq, \]
with $a$ a parameter. This is the simplest approximation and becomes exact for the uniform electron gas, as well for any region of space where the density is constant. The term $aq$ integrates to zero and is unimportant for the kinetic energy and its functional derivative. Thus, usually the parameter $a$ is set to zero. However, it was shown that $a = 5/3$ improves the quality of the TF KE density (not relevant for the present work).

(ii) **Second order gradient expansion** (GE2)\(^{15}\), defined as

$$F_s^{GE2} = 1 + \frac{5}{7} p + \frac{20}{3} q.$$  

(8)

As for the TF case, the last term in Eq. (7) integrates to zero and does not contribute to the kinetic energy and potential. Therefore, it is usually disregarded in most applications.

(iii) **Fourth-order gradient expansion** (GE4)\(^{23}\). This is written as

$$F_s^{GE4} = F_s^{GE2} + \Delta,$$  

(9)

with

$$\Delta = \frac{8}{71} q - \frac{1}{7} p q + \frac{8}{27} p^2 \geq 0.$$  

(10)

This enhancement factor is a simplified version, obtained via the Green’s theorem integration of terms comprising higher order derivatives of the density. It holds for finite systems under the assumption that $n(\mathbf{r})$ and $\nabla n(\mathbf{r})$ vanish as $r \rightarrow \infty$. For the full GE4 expression, see Refs. 55.\(^{12}\) We note that the GE4 KE displays a serious drawback for finite systems, as it shows the wrong behavior in the tail of the density of a finite system. In this region in fact the density decays exponentially as $n \sim e^{-\alpha r}$ and the von Weizsäcker KE is almost exact. Hence, the exact kinetic energy density behaves as $\tau \sim \tau^W \sim \tau^{GE2} \sim n$. On the other hand, we have that $\tau^{GE4} \sim n^{1/3}$, being much worse than GE2. Moreover, the corresponding potential diverges under the same conditions. This behavior is not surprising if we consider that higher-order gradient expansion terms are derived from small perturbations of the uniform electron gas, so that they contain the right physics for a slowly-varying density regime, but fail in rapidly-varying regions, such as in the tail of finite systems, or near the nucleus. For this reason GE4 usually worsens the atomic KE with respect to GE2. Similarly, the sixth-order gradient expansion (GE6)\(^{12}\), that contains terms of order $O(p^4, q^4, q^2 p, \ldots)$, has a kinetic energy density which even diverges in the tail of the density of a finite system ($\tau^{GE6} \sim n^{-1/3}$). Consequently, the GE6 KE diverges for any finite system.

(iv) **Modified fourth-order gradient expansion** (MGE4), defined by

$$F_s^{MGE4} = F_s^{GE4} \sqrt{1 + \left(\frac{\Delta}{1 + \frac{5}{7} p}\right)^2}.$$  

(11)

This construction was proposed in Ref. \(^{55}\) and recovers GE4 for a slowly-varying density. Instead, near the nucleus and in the tail of an atom (when $|q| \to \infty$) $F_s^{MGE4} \to 1 + F_s^{W}$, with $F_s^{W} = \frac{2}{7} p$ the von Weizsäcker (W) enhancement factor. This latter is a much reasonable limit for rapidly-varying density regions and is also the correct limit for a uniform density perturbed by a small-amplitude short-wavelength density wave\(^{22}\).

(v) The **Laplacian-level meta-GGA** (MGGA)\(^{55,56}\), with the following expression

$$F_s^{MGGA} = F_s^{W} + (F_s^{MGE4} - F_s^{W}) f_{ab}(F_s^{MGE4} - F_s^{W}),$$  

(12)

where

$$f_{ab}(z) = \begin{cases}
0 & z \leq 0 \\
\left(\frac{1+e^{a/(a-z)}}{e^{a/2}+e^{a/(a-z)}}\right)^b & 0 < z < a \\
1 & z \geq a,
\end{cases}$$  

(13)

is an analytical real-valued sharp-interpolating function, with $a = 0.5389$ and $b = 3$. Note that $F_s^{MGE4} - F_s^{W}$ is adimensional\(^{55}\). MGGA is one of the best models for the exact kinetic energy density, fulfilling many exact conditions, as the rigorous lower bound\(^{55,56}\):

$$\tau^W(\mathbf{r}) \leq \tau(\mathbf{r}).$$  

(14)

(vi) In addition to the models listed above, in this work we consider also a new Laplacian-level meta-GGA KE approximation defined by the simple ansatz

$$F_s^{L\kappa} = 1 + 2\kappa - \left(\frac{\kappa}{1 + \frac{2}{\pi}} + \frac{\kappa}{1 + \frac{4}{\pi}}\right),$$  

(15)

with

$$x_1 = \frac{5}{27} p + \Delta + \left(\frac{5}{27} p\right)^2,$$  

(16)

$$x_2 = 2 \frac{\left(\frac{5}{27} p\right) \Delta}{\kappa} + \left(\frac{5}{27} p\right)^3.$$  

(17)

This functional is designed to respect the following limits:

1. In the slowly-varying density limit ($s$ and $q \to 0$) it behaves as

$$F_s^{L\kappa} \to 1 + \frac{5}{27} p + \Delta + O(p^4, q^4, q^3 p, \ldots),$$  

(18)

therefore it recovers GE4, except for the unimportant term $(20/9) q$. Note also that Eq. (18) does not contain terms of 6-th order (e.g. $(\nabla n)^6$), so that it recovers GE4 quite closely for a wider range of (small) values of $s$ and $q$.

2. In the rapidly-varying density-limit ($s$ and $|q| \to \infty$) it behaves as

$$F_s^{L\kappa} \to 1 + 2\kappa.$$  

(19)

Thus, it can be made to recover the behavior of the APBEK\(^{63,64}\) or revAPBEK\(^{63,64}\) functionals in the rapidly-varying density-limit. Consequently, we define two variants of the functional:
L0.4 where $\kappa = \frac{1}{2} \kappa^{APBEK} = 0.402$

L0.6 where $\kappa = \frac{1}{2} \kappa^{revAPBEK} = 0.623$.

We constructed the L0.4/L0.6 functionals in order to recover the APBEK/revAPBEK limit because the latter functionals have been found to yield very accurate embedding energies still keeping good accuracy for other properties (total and relative kinetic energies). On the other hand, recent works at GGA level found that functionals diverging at large $s$ are very poor for the embedding theory.

We remark that the MGE4, MGGA, L0.4 and L0.6 functionals recover the GE4 limit, whereas other Laplacian-level meta-GGA KE functionals, e.g. those in Refs. 67,68,100, do not. Thus the latter functionals are not considered in this work.

Figure 1 shows several enhancement factors as functions of the reduced gradient $s = \sqrt{q}$. The plots are reported for two values of $q$ in the range appropriate to physical densities ($q = 0$ and $q = 2$). For GE2, W (von Weizsäcker), GE4, MGE4, and MGGA we subtracted in the plot the term $20/9q$ to have a more direct comparison with L0.4 and L0.6 that do not include such a term.

For $q = 0$ (upper panel), all functionals behave similarly in the small-$s$ region ($s \lesssim 0.7$), recovering the modified GE2 (in case of APBEK and revAPBEK GGAs), and the GE4 respectively (in case of Laplacian-based functionals). (See the inset in the upper panel of Fig. 1). At medium values of $s$ ($0.7 \lesssim s \lesssim 1.7$) MGE4, L0.4, and L0.6 still recover, by construction, the GE4 behavior. On the other hand, MGGA shows an unphysical strong oscillation, due to the sharp interpolation function (Eq. (13)). Finally, at large values of $s$, GE4, MGE4, and MGGA diverge, whereas the L0.4 and L0.6 functionals show a saturation towards the APBEK and revAPBEK limits. Note however that these limits are only reached at very large values of the reduced gradient. Moreover, all the functionals, but MGGA, fail to respect the exact constraint $F_s \geq F_s^W$.

For $q = 2$ (lower panel), a moderately-varying density regime is considered. The picture is not much changing for large $s$ values, but it is drastically modified for small values of the reduced gradient. Note however, that for the GGA functionals (APBEK and revAPBEK) we have of course the same plot as for the previous case. In this case, at medium values of $s$ all Laplacian-level meta-GGA functionals agree rather well, except for the unphysical oscillations displayed by MGGA. For small $s$-values ($s \to 0$) instead different behaviors are observed: MGE4 and MGGA have the same trend and move towards Thomas-Fermi: GE4 tends to $1 + (8/81)q^2 \approx 1.395$ displaying its well known divergent behavior in this regime; and finally, the L0.4 and L0.6 functionals tend to

$$F^L_{sK} \to 1 + 2\kappa - \frac{\kappa}{1 + \frac{S}{8r^2} + \kappa},$$

that is $F^{L0.4} \to 1.20$ and $F^{L0.6} \to 1.24$.

The density regime characterized by $s = 0$ and $q$ finite, is important in the middle of molecular bonds. In order to understand better this density regime, we report in Fig. 2, $\partial^2 F_s(s = 0, q)/\partial q^2$ versus $q$ for several enhancement factors. Note that $\partial^2 F_s(s, q)/\partial q^2$ is an intrinsic quantity of the enhancement factor, being independent on the linear $20q$ term. As reference, we consider the exact Kohn-Sham KE enhancement factor derivative ($\partial^2 F_{exact}(s = 0, q)/\partial q^2$) (computed numerically) in the middle of the bond between two interacting jellium slabs of $r_s = 3$, and thickness $2\lambda_F$ each (with $\lambda_F = 2\pi/k_F$ being the Fermi wavelength). When the jellium slabs are close to each other ($z \ll \lambda_F$), $q$ is small (and positive) at the bond, and when the jellium slabs are far to each other ($z \geq \lambda_F$), $q$ is large (and positive). While MGE4 and MGGA show strong oscillations, the LK functionals perform remarkably smooth, and close to the reference. Note that in case $s = 0$ and $q \to 0$, the reference curve contains higher order terms that are out of reach for GE4-based functionals.
III. DENSITY-SCALING BEHAVIOR

Consider the family of density scalings

\[ n_\lambda(r) = \lambda^{3\beta+1} n(\lambda^\beta r), \quad \lambda > 0, \]

where \( \beta \) is a parameter, which is changing not only the external potential associated with the density \( n \) (as in the uniform density scaling), but also the particle number \( (N \rightarrow \lambda N) \). Under this scaling, the KE gradient expansion terms behave as

\[ T_s^{TF}[n_\lambda] = \lambda^{2\beta+5/3} T_s^{TF}[n], \]

\[ T_s^{(2)}[n_\lambda] = \frac{1}{2} T_s^{W}[n_\lambda] = \frac{1}{2} \lambda^{2\beta+1} T_s^{W}[n], \]

\[ T_s^{(4)}[n_\lambda] = \lambda^{2\beta+1/3} T_s^{(4)}[n], \]

where \( T_s^{(2)} = \int \, dr \, \tau^{TF} [\frac{\partial T_s}{\partial \tau} + \frac{2\lambda}{3} q] \) and \( T_s^{(4)} = \int \, dr \, \tau^{TF} \Delta(p, q) \) are the second- and fourth-order terms of the KE gradient expansion, respectively. On the other hand, the reduced gradient \( s \) and the reduced Laplacian \( q \) scale according to \( s_\lambda(r) = \lambda^{-1/3} s(\lambda^2 r) \) and \( q_\lambda(r) = \lambda^{-2/3} q(\lambda^2 r) \), respectively. Thus, the reduced gradient and Laplacian are independent on \( \beta \) and the slowly varying density limit \( (s, q \rightarrow 0) \) is reached whenever \( \lambda \rightarrow \infty \). For \( \lambda \ll 1 \), instead, a rapidly varying density regime is always set up.

For different values of the parameter \( \beta \), Eq. (21) spans an impressive number of physical properties. Some of them are analyzed in next subsections.

A. Thomas-Fermi scaling

For \( \beta = \frac{1}{3} \) the Thomas-Fermi scaling is obtained. In the limit \( \lambda \rightarrow \infty \) any system resembles the features of the Thomas-Fermi density. The Thomas-Fermi scaling is the basis for the asymptotic expansion of the kinetic energy

\[ T_s = c_0 \lambda^{7/3} + c_1 \lambda^2 + c_2 \lambda^{5/3} + \ldots , \]

with the coefficients \( c_0 = 0.768745 \), \( c_1 = -0.5 \), and \( c_2 = 0.2699 \), fixed from the same semiclassical theory of the neutral atom. The expansion of Eq. (21) is very accurate even for real atoms with errors of the order of 0.5%-0.2%.

Using the method proposed in Ref. 101, we extracted the coefficients \( c_1 \) and \( c_2 \) for all the functionals considered in this work. We assumed that, because all the functionals recover TF for a constant density, they have the exact \( c_0 \) coefficient.

The deviations

\[ \Delta c_i = 1000(c_i^{\text{approx}} - c_i^{\text{exact}}) \quad i = 1, 2, \]

are reported in Table II for each functional. As it might be expected, the best results are found for revAPBEK and APBEK which were constructed from the semiclassical theory of the neutral atom. The Laplacian-level meta-GGA provide a good performance, improving over GE2 for \( c_1 \), which is the leading term of quantum effects, beyond the Thomas-Fermi theory.

B. Uniform-electron-gas scaling

The uniform-electron-gas scaling is obtained for \( \beta = -\frac{1}{3} \) and \( \lambda \rightarrow \infty \). Under these conditions, in fact, the density becomes very slowly varying, being almost constant over the space. Hence, the gradient expansion

\[ T_s^{\text{exact}}[n_\lambda] = T_s^{TF}[n_\lambda] + T_s^{(2)}[n_\lambda] + T_s^{(4)}[n_\lambda] + T_s^{(6)}[n_\lambda] + \ldots , \]

which was derived from small perturbations of the uniform electron gas, is (almost) exact in this limit. As a consequence all the Laplacian-level meta-GGA kinetic functionals, which recover GE4 in the slowly varying density, become very accurate under the uniform-electron-gas scaling.

C. Homogeneous scaling

Setting \( \beta = 0 \) and considering the limit \( \lambda \rightarrow 0 \) the homogeneous scaling is realized. This scaling is a valuable tool in DFT and was used to study the static correlation and the delocalization errors, as well as to construct kinetic energy functionals.

Following Ref. 101 we study the ability of different functionals to satisfy the homogeneous scaling by considering the hydrogen density \( n_H \) and the associated effective scaling order

\[ s_{\text{eff}} = \int_0^1 \frac{\ln \left( \frac{[T_s[n_H]]}{\ln(\lambda)} \right) - \ln \left( [T_s[n_H]] \right) }{\ln(\lambda)} d\lambda . \]

This provides a measure for the scaling behavior of a generic KE functional. The exact result is \( s_{\text{eff}}^{\text{exact}} = 1 \) (see eq. 21).
TABLE I: Error statistics for different density scaling tests. The best Laplacian-level meta-GGA result in each line is highlighted in bold face; the worst Laplacian-level meta-GGA result for each test is underlined.

|                      | TF    | GE2   | APBEK | revAPBEK | GE4   | MGE4  | MGGA | L0.4 | L0.6 |
|----------------------|-------|-------|-------|----------|-------|-------|-------|------|------|
|                      |       |       |       |          |       |       |       |      |      |
| Thomas-Fermi scaling |       |       |       |          |       |       |       |      |      |
| (β = 1/3)            |       |       |       |          |       |       |       |      |      |
| Δc₁                  | -160.87 | -36.25 | -3.77 | -2.20 | -20.39 | -17.28 | -8.90 | -23.40 | -22.39 |
| Δc₂                  | 115.50 | 66.13 | 17.14 | 27.00 | 73.25 | 72.84 | 47.53 | 79.04 | 74.75 |
| Homogeneous scaling  |       |       |       |          |       |       |       |      |      |
| (β = 0)              |       |       |       |          |       |       |       |      |      |
| Δsᵥff               | 0.67  | 0.55  | 0.59  | 0.58  | 0.47  | 0.49  | 0.02  | 0.54  | 0.53  |
| Fractional scaling   |       |       |       |          |       |       |       |      |      |
| (β = 1)              |       |       |       |          |       |       |       |      |      |
| Δsᵥff               | 0.67  | 0.47  | 0.50  | 0.50  | 0.38  | 0.42  | 0.02  | 0.45  | 0.44  |
| ∆                   | 12.26 | 34.71 | 33.84 | 35.89 | 38.05 | 38.89 | 7.77  | 37.15 | 37.15 |

The errors on $s_{vff}$ for different functionals are reported in Table I. All the Laplacian-level meta-GGA functionals perform slightly better than GGA ones, showing that the inclusion of the Laplacian dependence can help to improve the physical behavior of the functional. Notably, the MGGA functional outstands over the other meta-GGAs, scaling almost perfectly under the homogeneous scaling. This finding supports the conclusion that this sophisticated approximation can capture very well the physics of the von Weizsäcker functional.

D. Fractional scaling

The fractional scaling is defined by $β = 1$ and the limit $λ \to 0$. It describes the physics of systems with a fractional number of electrons. In particular, the fractional scaling is related to the disintegration of the hydrogen atom into fragments with fractional nuclear and electronic charge, which is a model for atomization processes in molecules.

In analogy with Ref. 101, we define the kinetic disintegration energy of the hydrogen atom into fragments with partial charge $λ$ and $1 - λ$ as

$$M(λ) = T_s(1) - T_s(λ) - T_s(1 - λ),$$  \hspace{1cm} (27)

where $T_s(1)$ is the kinetic energy of the hydrogen atom and $T_s(λ)$ the KE of the neutral fragment with charge $λ$. The disintegration error is then defined as

$$ΔM = \int_0^1 [M^{DFT}(λ) - M^{exact}(λ)] dλ. \hspace{1cm} (28)$$

In Table I we report, for all the functionals the errors on the effective scaling order (Eq. (27)), with respect to the exact result $s_{vff} = 3$, and the disintegration error of Eq. (28). The results show that the effective scaling order follows a similar trend as for the homogeneous scaling. The inspection of the values of $ΔM$ provides a couple of additional considerations, that show the importance of fractional scaling: (1) The MGGA functional works very well for both $s_{vff}$ and the disintegration problem, showing that this functional can incorporate most of the von Weizsäcker physics without error compensations; (2) The TF functional displays the worst $s_{vff}$ value, but the smallest error for the disintegration problem (except for MGGA), showing an important error cancellation.

IV. COMPUTATIONAL DETAILS

A. Kinetic energies

To test the different functionals we assessed their ability to compute the KE of different systems:

- **Model one- and two-electron systems.** These include the one-electron Gaussian, hydrogen, and cupless-hydrogen densities, as well as the Hooke’s atom at various values of the harmonic potential. For these systems the reference values were computed via the von Weizsäcker formula. In all calculations exact densities were employed.

- **Jellium systems.** We considered: (i) a series of Na jellium clusters ($r_s = 3.93$) with magic electron number 2, 8, 18, 20, 34, 40, 58, 92, and 106, used also in Refs. 55, 56, 64; (ii) jellium surfaces with bulk parameter $r_s = 2, 4, 6$ into the liquid drop model (LDM) (as in Refs. 55, 56, 64); and (iii) two interacting jellium slabs at different distances. Each jellium slab has $r_s = 3$ and a thickness of $2λ_F$.

All the calculations were performed using the orbitals and densities resulting from numerical Kohn-Sham calculations within the local density approximation for the exchange-correlation functional ((ii) and (iii)), and the exact exchange functional ((i)).
• **Atoms and ions** We tested the benchmark set of atoms and ions used in Refs. 55, 56, 64. All calculations employed analytic Hartree-Fock orbitals and densities\(^{114}\). We also calculated the ionization KE of noble atoms (until Uuo), using accurate exchange-Kohn-Sham densities and orbitals.

• **Molecules.** We considered the set of molecules including H\(_2\), HF, H\(_2\)O, CH\(_4\), NH\(_3\), CO, F\(_2\), HCN, N\(_2\), CN, NO, and O\(_2\). This set was already used in Refs. 55, 56, 113. The noninteracting kinetic energies of test molecules were calculated using the PROAMV code\(^{116}\). The required Kohn-Sham orbitals were obtained by Kohn-Sham calculations performed with the uncontracted 6-311+G(3df,2p) basis set, the Becke 1988 exchange functional\(^{117}\), and Perdew-Wang correlation functionals\(^{118}\).

**B. FDE calculations**

The FDE calculations were performed using the FDE script as implemented in the TURBOMOLE program package\(^{119}\). Details about our KSCED implementation in TURBOMOLE are discussed in Refs. 64, 59. The implementation of Laplacian-level meta-GGA functionals is briefly presented in the appendix. In all calculations, the PBE\(^{120}\) XC functional and def2-TZVPDD monomolecular basis sets\(^{121,122}\) were used. The use of a monomolecular approach was needed to guarantee good convergence for calculations using the MGE4 and MGGA KE functionals. These functionals show in fact marked oscillations in the KE potential in the tail of the density. Therefore, they can give rise to important numerical problems in a supermolecular basis-set calculation. The quality of our results was tested by comparing monomolecular and supermolecular results for the GGA KE functionals as well checking the convergence with increasingly large basis sets. We found that the def2-TZVPPDD basis set, adding diffuse basis functions to the def2-TZVPP\(^{121}\) basis, provides finally a reliable description of all the systems considered in this paper.

The FDE calculations were performed on the following test systems, characterized by different interaction characters: He-Ne, He-Ar, Ne-Ne, Ne-Ar, CH\(_4\)-Ne, CH\(_4\)-CH\(_4\) (weak interaction); H\(_2\)S-H\(_2\)S, HCl-HCl, H\(_2\)S-HCl, CH\(_3\)Cl-HCl, CH\(_3\)SH-HCN, CH\(_3\)SH-HCl (dipole-dipole interaction); NH\(_3\)-NH\(_3\), HCl-HCl, H\(_2\)O-H\(_2\)O, NH\(_3\)-H\(_2\)O, HF-NCH\(_3\), (HCONH\(_2\))\(_2\), (HCOOH)\(_2\) (hydrogen bond). The geometry of the complexes was taken from Refs. 55, 123, 124. The subsystems A and B are the monomer units.

The error on the total embedding energy was computed as\(^{99,93}\)

\[
\Delta E = E^{FDE}[n_A; n_B] - E^{KS}[n^{KS}], \tag{29}
\]

where \(E^{FDE}[n_A; n_B]\) is the FDE total energy obtained from the embedded subsystem densities \(n_A\) and \(n_B\), whereas \(E^{KS}\) is the conventional Kohn-Sham total energy corresponding to the ground state density \(n^{KS}\). The performance of the different approaches was evaluated, within each group of molecules, by computing the mean absolute error (MAE).

The errors on the embedding densities were studied by considering the deformation density

\[
\Delta n(r) = n_A(r) + n_B(r) - n^{KS}(r). \tag{30}
\]

Some plots for different systems were realized by representing the plane-averaged deformation density

\[
\langle \Delta n \rangle_{xy}(z) = \int \int |\Delta n(x, y, z)| \, dx \, dy, \tag{31}
\]

where we used Cartesian coordinates explicitly and the \(z\) direction is along the intermolecular axis. Finally, a quantitative measurement of the absolute error associated with a given embedding density was obtained by computing the embedding density error

\[
\xi = \frac{1000}{N} \int |\Delta n(r)| \, dr, \tag{32}
\]

with \(N\) the number of electrons. In the evaluation of \(\xi\), only valence electron densities were considered; Core densities are in fact much higher than valence ones and would largely dominate the calculation of \(\xi\). On the other hand, core densities are not very important for the determination of chemical and physical properties of the interaction between the subsystems, which are of interest here.

**V. KINETIC ENERGIES OF MODEL SYSTEMS**

In this section we present the KE results of one-electron densities, the Hooke’s atom, jellium surfaces, jellium clusters, interacting jellium slabs, atomic systems, and molecules. All results are summarized in Table II where we report mean absolute relative errors (MARE) for each test. In addition, the last line shows the average performance relative to the GE2 method, defined as

\[
\text{MARE(GE2)} = \frac{1}{N} \sum_{i=1}^{N} \frac{\text{MARE}_i}{\text{MARE}_{\text{GE2}}}, \tag{33}
\]

where the sum runs over all the \(N\) tests and \(\text{MARE}_i\) is the MARE of the \(i\)-th test.

**A. One-electron densities**

We tested the different KE functionals on three model one-electron densities, namely, the hydrogen (H), the Gaussian (G), and the cuspidal densities. These are defined as

\[
n_H(r) = \frac{e^{-2r}}{\pi}, \quad n_G(r) = \frac{e^{-r^2}}{\pi^{3/2}}, \quad n_C(r) = \frac{(1 + r)e^{-r}}{32\pi}. \tag{34}
\]
TABLE II: Mean absolute relative errors (MARE) in %, for different tests, and absolute relative errors for one-electron densities (G, H, and C). The best Laplacian-level meta-GGA result in each line is highlighted in bold face; the worst Laplacian-level meta-GGA result for each test is underlined. The last line reports the MARE relative to the GE2 performance (see text for details). The star symbol stands for a Laplacian-level meta-GGA KE functional that is better than the best GGA.

|          | GGAs     | meta-GGAs |
|----------|----------|-----------|
|          | TF       | GE2       | APBEK | revAPBEK | GE4 | MGE4 | MGGA | L0.4 | L0.6 |
| Total kinetic energies |
| H        | 8.2      | 2.9       | 2.2   | 3.1      | 5.8 | 5.5   | 2.5  | 4.4  | 4.5  |
| G        | 10.1     | 1.0       | 1.8   | 0.8      | 15.3| 4.2   | 3.7  | *0.7 | 1.3  |
| C        | 5.3      | 5.8       | 4.6   | 5.6      | 8.9 | 7.9   | *2.1 | 6.8  | 9.6  |
| Hooke’s atom KE | 25.7     | 14.6      | 18.8  | 17.8     | 14.6 | *10.3 | *14.3 | 15.5 | *14.5 |
| Jell. clust. KE | 4.4      | 1.0       | 1.0   | 0.8      | 1.9 | 1.2   | *2.5 | 0.9  | 1.0  |
| Jell. slabs KE | 1.89     | 0.57      | 0.55  | 0.46     | *0.21| *0.36 | *0.36 | *0.42 | *0.39 |
| Atoms’ KE | 8.4      | 1.1       | 0.8   | 1.2      | 2.5 | 2.4   | 1.4  | 1.9  | 2.0  |
| Molecules’ KE | 9.7      | 0.9       | 0.5   | 0.4      | 1.0 | 1.2   | 1.4  | 0.8  | 0.8  |
| MARE(GE2) | 5.42     | 1         | 0.94  | 0.88     | 2.16| 1.42  | 1.43 | 1.07 | 1.15 |
| Kinetic energy differences |
| Jell. clust. DKE | 17.7     | 27.2      | 18.9  | 23.1     | 50.2| 39.0  | *17.9 | 29.3 | 31.5 |
| Jell. surf. LDM | 8.1      | 3.3       | 3.9   | 3.6      | *1.7 | *1.9  | *2.5 | *2.8 | *2.8 |
| Jell. slabs DKE | 17.28    | 5.02      | 4.17  | 3.45     | *1.31| 3.53  | 3.53 | *2.85 | *2.65 |
| Atoms’ IKE | 49.5     | 42.2      | 45.8  | 44.4     | *39.2| *40.7 | 51.8 | 41.3 | 41.4 |
| Molecules’ AKE | 106.0    | 184.0     | 142.0 | 155.0    | 222.0| 207.0 | *108.0 | 216.0 | 217.0 |
| MARE(AKE) | 1.66     | 1.00      | 0.91  | 0.90     | 0.95 | 0.96  | *0.79 | 0.93 | 0.94 |

They were used in the construction of several XC functionals, being simple models for simple iso-orbital regions.

For these model densities, the von Weizsäcker functional is exact and behaves as $\tau^W \sim n$. A similar behavior is found for GE2 and all the GGAs, which therefore perform rather well for this problem, with errors below 6%. On the other hand, for GE4, in iso-orbital regions, we have $\tau^{GE4} \sim n^{1/3}$. For this reason GE4 performs significantly worse than GE2 in all cases (errors up to 15%) (note that it is in general also worse than TF, that has $\tau^{TF} \sim n^{5/3}$). Finally, the correct behavior is restored for the other Laplacian-level meta-GGAs, which thus describe these one-electron densities reasonably well. In particular, in two cases, Laplacian-level meta-GGA functionals are more accurate than the best GGA functional: MGGA is the most accurate for the delocalized C density, and the L0.4 functional is the most accurate for the Gaussian density.

B. Hooke’s atom

The Hooke’s atom consists of two interacting electrons in an isotropic harmonic potential of frequency $\omega$. At small values of $\omega$, the electrons are strongly correlated. At large values of $\omega$, they are tightly bound. The exact ground state solutions for the Hooke’s atom are known for special values of $\omega$. We consider here the first nine values of $\omega$ for which an analytical solution is available: from $\omega = 0.25$ (strongly bound electrons), to $\omega = 3.597 \times 10^{-6}$ (strongly-correlated electrons).

The MARE with respect to the exact von Weizsäcker values ($\left(\frac{T_s^{approx} - T_s^W}{T_s^W}\right) \times 100$), are reported in Table III. The best Laplacian-level meta-GGA results are obtained with MGE4, MGGA and L0.6, which are also

![FIG. 3: Kinetic energy deviations from the exact values of several KE functionals, versus the classical electron distance in the Hooke atom. The von Weizsäcker KE functional is the exact reference for a two-electron closed-shell system.](image)
superior to the best GGAs.

In Fig. 3 we report the individual deviations for the smallest values of the classical electron distance $r_0 = (\omega^2/2)^{-1/3}$. The L0.4 and L0.6 functionals provide the best description over the whole range of frequencies, whereas the good performance of MGE4 originates from a high accuracy in the strongly correlated regime, while in the tightly bound regime much larger errors are found.

C. Jellium clusters

The KE MAREs for jellium clusters are reported in Table III. The best Laplacian-level meta-GGA functional is L0.4, the worst is MGGA.

The errors on the kinetic energy per electron ($\Delta T_s/N = (T_s^{approx}(N) - T_s^{exact}(N))/N$) for the different clusters are reported in Fig. 4 for some selected KE functionals (TF, GE2, GE4, MGE4, MGGA, L0.4). The data in Fig. 4 show that almost of all functionals (but MGGA and TF) work very well for medium and large clusters (i.e. for N ≥ 18 the is error below 1 mHa) whereas for smaller clusters definitely larger errors are found. On the other hand, the TF functional yields always an underestimation of the kinetic energy, less dependent on the cluster’s size, whereas the MGGA functional shows strong oscillations. In Fig. 4 it is worth to note the very close performances of GE2 and L0.4 for all N, as these two functionals are quite different from each other (see Fig. 4). This small error originates from an error cancellation between the region inside the cluster ($s < 1$ and $|q| < 1$) where L0.4 is (by construction) almost the same of GE4 and thus larger than GE2, and the region outside the cluster ($s$ and $q$ are large) where GE2 is larger than L0.4, as the former diverges with $s$ while the latter approaches a constant (see Eq. (??)).

The dependence of the KE errors on the clusters’ size suggests that interesting results may be obtained for the cluster disintegration problem. We consider the disintegration of the cluster with $N = 106$ into smaller magic clusters. The energy associated with this process is called disintegration kinetic energy (DKE) and is defined as:

$$DKE = \sum_i m_i T_s(N_i) - T_s(106)$$

where $m_i$ are the masses of the smaller clusters.

In Fig. 4 we report the individual deviations for KE, with MARE 4.4.

D. Jellium surfaces and slabs

The LDM MARE of several jellium surfaces are reported in Table III. The best results are found for GE4, and in general all Laplacian-level meta-GGA functionals (recovering GE4) overcomes the best GGAs. In fact the uniform-electron-gas scaling becomes important in these systems.

We also calculated the KE energy of two interacting jellium slabs, for various distances $0 ≤ z ≤ \lambda_F$ between the slabs. The MARE represents $(1/\lambda_F) \int_0^{\lambda_F} dz |T_s(z) - T_s^{exact}(z)|/T_s^{exact}(z)$, and is reported in Table III for each functional. Table III shows that GE4 and all the Laplacian-level meta-GGA functionals are significantly better than GGAs.

Finally we also consider the following KE difference

$$DKE = (1/\lambda_F) \int_0^{\lambda_F} dz |[T_s(z) - T_s(0)] - [T_s^{exact}(z) - T_s^{exact}(0)]|/T_s^{exact}(z) - T_s^{exact}(0),$$

which represents the dissociation KE of a jellium slab into two pieces and can be used as an indicator of the quality of the functionals in describing bonding regions. Also in this case Table III shows that Laplacian-level meta-GGA functionals (GE4, L0.4 and L0.6) are significantly better than GGAs.

E. Atoms, ions and molecules

The KE MAREs of a benchmark set of atoms and ions are reported in Table III. The GGAs yield a MARE of about 1%, whereas all the Laplacian-level meta-GGAs are about twice worse, with a MARE in the range 1.4-2.5%.

As an additional test, we considered the ionization kinetic energies (IKE=$T_s^{atom} - T_s^{ion}$) of the noble gases (up to Uuo). Note that because of the virial theorem, IKEs are just equal to the regular ionization potentials. The
MARE IKEs are reported in Table III. All the Laplacian-level meta-GGA functionals (but MGGA) are better than the GGAs. This result traces back to the fact that the GE4-based functionals behave better than GGAs for the homogeneous and fractional scalings, and thus for systems with fluctuating number of electrons.

Then we considered the total and atomization KEs (AKE) of a set of molecules. We recall that the latter is a hard test for any kinetic energy functional and that most of the functionals even fail to yield a qualitative description of AKEs.35 Laplacian-level meta-GGA functionals show a MARE for the total KEs below or close to 1%56, which is close to GE2 one but much worse than the best GGA. For the AKE the trend is similar to the disintegration of the 106e− jellium cluster.

F. Summary and overall assessment

Table I reports the global MARE (relative to GE2) for the total KE and the KE differences. Concerning the total KE, the best Laplacian-level meta-GGA functional is L0.4, followed by L0.6. Both functionals are largely better than GE4 (MARE reduced to one half) but are a little worse than the GGA functionals. However, when KE differences are considered, the best performance is yield by the MGGA functional, which definitely overcomes the best GGA. The other Laplacian-level meta-GGA functionals also show very good performances for KE difference, with MARE(GE2) in the range 0.93-0.96, close to the best GGA.

These results show that in general the inclusion of the Laplacian can improve the description of the noninteracting kinetic energy. However, the proper dependence on this parameter is not captured in a systematic way by any of the functionals that we examined here. As a result the Laplacian-level meta-GGA functionals perform in a rather erratic way overcoming the GGAs for some properties and systems, but also showing sudden failures for other cases. Anyway, the results summarized in Table I indicate that the L0.4 (and L0.6) functional have a more regular behavior than other Laplacian-level meta-GGAs and can be competitive in numerous applications.

VI. FDE CALCULATIONS

In this section we present the results of FDE calculations on different test systems using the Laplacian-level meta-GGA functionals considered in this paper, but GE4 which gives very poor results, even failing to converge in some cases, possibly due to its wrong tail behavior.

A. Embedding densities

The errors on embedding densities (see Eq. (??)) are reported in Table III. We recall that this is an important test for embedding approaches because it provides direct insight into the quality of the embedding potential.64,89,90,129-133

Inspection of the data shows that the L0.4 and L0.6 functionals perform very similarly and are in line with the GGA functionals. On the other hand, MGE4 and MGGA provide significantly larger errors. For weakly interacting systems both functionals display a very poor performance and interestingly almost the same results. This finding is rationalized considering that, for dispersion-dominated, systems the bond region is characterized by small and medium values of s and quite large values of q (q ≥ 5). Thus, in this region the two functionals, and the corresponding kinetic potentials, are the same by construction (in this region $F^{s}_{MGE4} > F^{s}_{MGE4}$, hence $F^{s}_{MGGA} = F^{s}_{MGE4}$; see Eqs. (??) and (??)). On the other hand, for the other kinds of interactions only MGGA shows a poor behavior whereas MGE4 performs rather well, being even the best Laplacian-level meta-GGA for hydrogen bonds. In this cases in fact the bonding region is characterized by small and medium values of s but small values of q (q ∼ 0.5) so that MGE4 recovers GE4 which is a reasonable limit. However, this is exactly the range of values where the sharp interpolating function of MGGA assumes its intermediate values. Therefore, the MGGA potential is strongly oscillating in this region (see Fig. 1).

To understand better these results we consider in Fig. 5 the plot of the plane-averaged deformation density in two typical cases: for the hydrogen-bond complex HF-NCH and for the weakly-interacting Ne-Ar system. The figure confirms the findings of Table I and additionally indicates that the GGA and L0.4/L0.6 densities are in fact similar over the whole space (not only after integration). On the contrary, MGE4 is very similar to L0.4 for hydrogen-bond complexes, but almost identical to MGGA for weakly interacting complexes. Note that the HF-NCH complex has larger density in the bond than Ne-Ar, thus the relative errors in the bond are larger in the latter case that in the former.

B. Embedding energies

The errors on the embedding energies obtained from FDE calculations using different kinetic functionals are reported in Table IV. The data show that the L0.4 functional has MAE = 1.06 mHa, which is comparable to that of the state-of-the-art GGA KE functionals and lower than the errors originating from the XC approximation.124 In particular, L0.4 yields the best results (MAE = 0.64 mHa) for the dipole-dipole interaction systems, also outperforming revAPBEK. On the other hand, lower accuracy is obtained for the weakly interacting systems: this drawback can be related to the inaccuracy of the gradient expansion for this class of systems, as it will be explained below (see next section). In general, we can state the important result that the L0.4 Laplacian-level functional can be effectively used to ap-
TABLE III: Global absolute errors on embedding densities (see Eq. (??)), resulting from FDE calculations with different KE functionals on several classes of systems (weak, dipole, and hydrogen-bonded systems). The mean absolute error (MAE) for each set of molecules, and the total MAE are also reported. The best (worst) Laplacian-level meta-GGA KE functional on each line is highlighted in bold (underline) style. The star symbol stands for a Laplacian-level meta-GGA KE functional that is better than the best GGA.

| system     | GGAs | meta-GGAs |
|------------|------|-----------|
|            | TF   | GE2       | APBEK | revAPBEK | MGE4 | MGGA | L0.4 | L0.6 |
| Weak interaction |      |           |       |          |      |      |      |
| He-Ne      | 0.71 | 0.71      | 0.70  | 1.46     | 1.53 | 0.76 | 0.76 |
| He-Ar      | 0.78 | 0.78      | 0.78  | 1.88     | 2.01 | 0.85 | 0.83 |
| Ne-Ne      | 0.17 | 0.12      | 0.08  | 1.58     | 1.58 | 0.26 | 0.21 |
| Ne-Ar      | 0.22 | 0.14      | 0.10  | 2.02     | 2.02 | 0.31 | 0.19 |
| CH₄-Ne     | 0.04 | 0.06      | 0.18  | 2.32     | 2.32 | 0.38 | 0.21 |
| CH₄-CH₄    | 0.59 | 0.22      | 0.60  | 5.89     | 5.64 | 0.36 | 0.77 |
| MAE        | 0.45 | 0.86      | 0.34  | 0.37     | 2.53 | 2.52 | 0.47 | 0.47 |
| Dipole-dipole interaction |      |           |       |          |      |      |      |
| H₂S-H₂S    | 2.19 | 2.01      | 2.02  | 4.17     | 4.67 | 2.73 | 2.86 |
| HCl-HCl    | 2.70 | 2.47      | 2.50  | 4.56     | 4.76 | 2.76 | 2.86 |
| H₂S-HCl    | 5.07 | 4.79      | 4.78  | 6.59     | 4.73 | 4.77 |
| CH₃Cl-HCl  | 3.35 | 3.04      | 3.08  | 5.35     | 3.15 | 3.28 |
| CH₃SH-HCN  | 1.86 | 1.91      | 2.05  | 5.08     | 2.56 | 2.68 |
| CH₃SH-HCl  | 5.47 | 5.22      | 5.24  | 6.60     | 5.13 | 5.15 |
| MAE        | 3.44 | 3.24      | 3.29  | 3.65     | 5.47 | 3.51 | 3.60 |
| Hydrogen-bond interaction |      |           |       |          |      |      |      |
| NH₃-NH₃    | 2.32 | 2.04      | 2.12  | 2.50     | 5.44 | 2.92 | 2.91 |
| HCl-HCl    | 2.60 | 2.22      | 2.19  | 2.28     | 5.36 | 2.35 | 2.36 |
| H₂O-H₂O    | 2.94 | 2.55      | 2.58  | 2.67     | 6.02 | 3.02 | 2.96 |
| NH₃-H₂O    | 4.57 | 4.26      | 4.26  | 4.04     | 7.38 | 4.52 | 4.47 |
| HF-NCH     | 4.49 | 4.22      | 4.22  | 4.09     | 6.80 | 4.29 | 4.25 |
| (HCONH₂)₂  | 3.17 | 3.17      | 3.30  | *3.13    | 6.78 | 3.92 | 3.78 |
| (HCOOH)₂   | 4.84 | 4.65      | 4.69  | *4.03    | 6.85 | 4.82 | 4.69 |
| MAE        | 3.57 | 3.38      | 3.34  | *3.25    | 6.38 | 3.69 | 3.63 |
| Overall assessment |      |           |       |          |      |      |      |
| MAE        | 1.81 | 1.66      | 1.69  | 2.31     | 3.52 | 1.86 | 1.86 |

approximate the non-additive kinetic energy functional in embedding calculations, yielding accurate total embedding energies for non-covalently interacting systems.

Concerning the other Laplacian-level meta-GGA functionals, slightly worse results are found with L0.6, which gives in any case rather good results, in line with APBEK. For the MGE4 and MGGA functionals, similar considerations as for the case of the embedding densities apply. In fact, as shown in more details in next section, for weakly-interacting systems the embedding energy is mainly determined by the region having moderately large $s$-values and large values of the reduced Laplacian ($q ≳ 5$). Hence, the two functionals perform very similarly and yield strongly underestimated embedding energies (see next section). For dipole-dipole and hydrogen bond interactions, instead, the relevant region for the embedding energy is defined (see next section) by relatively small values of the reduced parameters ($s ≲ 1$ and $|q| ≲ 1$). Thus, MGE4 correctly tends to GE4, which is a rather good approximation for this limit, whereas MGGA is dominated by the interpolating function.
TABLE IV: Embedding energy errors (see Eq. (??)) in mHa, resulting from FDE calculations with different KE functionals on several classes of systems (weak, dipole, and hydrogen-bonded systems). The second column reports the benchmark binding energy $E_b$ from Ref. 124. The mean error (ME), MAE, and MARE are indicated for each set of molecules. At the bottom of the table, the total MAEs are also reported. The best (worst) Laplacian-level meta-GGA value on each line is highlighted in bold (underline) style. The star symbol stands for a Laplacian-level meta-GGA KE functional that is better than the best GGA.

| system    | $E_b$ | TF | GE2 | APBEK | revAPBEK | MGE4 | MGGA | L0.4 | L0.6 |
|-----------|-------|----|-----|-------|----------|------|------|------|------|
| **Weak interaction** |       |    |     |       |          |      |      |      |      |
| He-Ne     | 0.06  | 0.38 | -0.81 | 0.39  | 0.35     | -3.28 | -3.24 | 0.48 | 0.50 |
| He-Ar     | 0.10  | 0.43 | -0.81 | 0.44  | 0.40     | -3.20 | -3.13 | 0.55 | 0.57 |
| Ne-Ne     | 0.13  | 0.29 | -1.67 | 0.27  | 0.18     | -4.45 | -4.45 | 0.50 | 0.52 |
| Ne-Ar     | 0.21  | 0.34 | -1.84 | 0.29  | 0.16     | -4.44 | -4.44 | 0.57 | 0.57 |
| CH$_4$-Ne | 0.35  | 0.37 | -2.05 | 0.32  | 0.18     | -7.11 | -7.11 | 0.63 | 0.64 |
| C$_2$H$_6$-Ne | 0.75 | 1.05 | -4.49 | 0.61  | 0.14     | -13.39 | -13.39 | 1.59 | 1.45 |
| CH$_4$-CH$_4$ | 0.81 | 0.78 | -4.39 | 0.21  | -0.29   | -13.29 | -13.13 | 1.05 | 0.67 |
| **MAE** | 0.52  | 2.29 | 0.36 | 0.24  | 7.02     | 6.98  | 0.77  | 0.70 |
| **Dipole-dipole interaction** |       |    |     |       |          |      |      |      |      |
| H$_2$S-H$_2$S | 2.63 | 2.15 | -4.68 | 0.67  | -0.23    | -3.83 | -7.80 | *-0.04 | -1.19 |
| HCl-HCl   | 3.20  | 3.26 | -4.76 | 1.49  | 0.44     | -3.67 | -8.76 | 0.73  | -0.60 |
| H$_2$S-HCl | 5.34  | 5.28 | -4.83 | 2.46  | 1.05     | -3.91 | -12.55 | *0.44  | *-1.03 |
| CH$_3$CH$_2$HCl | 5.66 | 5.92 | -7.08 | 2.42  | 0.63     | -7.37 | -15.40 | 1.12  | -0.82 |
| CH$_3$SH-NCN | 5.72 | 2.79 | -7.34 | 0.53  | -0.79    | -9.84 | -16.55 | *-0.24 | -1.78 |
| CH$_3$SH-HCl | 6.63 | 8.46 | -6.30 | 3.80  | 1.72     | -7.23 | -19.07 | 1.12  | -0.82 |
| **MAE** | 4.64  | 5.83 | 1.90 | 0.81  | 13.36    | *0.64 | 1.01  |
| **Hydrogen-bond interaction** |       |    |     |       |          |      |      |      |      |
| NH$_3$-NH$_3$ | 5.02 | 2.63 | -6.20 | 0.63  | -0.55    | -5.21 | -11.44 | *-0.50 | -1.94 |
| HCl-HCl   | 7.28  | 5.23 | -5.87 | 2.77  | 1.29     | -4.21 | -15.32 | 1.42  | -0.49 |
| H$_2$O-H$_2$O | 7.92 | 4.77 | -6.82 | 1.84  | 0.24     | -5.42 | -16.22 | *-0.17 | -1.93 |
| NH$_3$-H$_2$O | 10.21 | 5.97 | -6.98 | 2.17  | 0.34     | -6.28 | -19.97 | *-0.86 | -2.65 |
| HF-NCH   | 11.33 | 8.26 | -6.62 | 3.78  | 1.67     | -5.85 | -23.46 | *0.17  | -1.89 |
| (HCONH$_2$)$_2$ | 23.81 | 10.59 | -17.72 | 1.43  | -2.76    | -17.85 | -52.06 | *-5.42 | -9.11 |
| (HCOOH)$_2$ | 25.74 | 18.81 | -15.77 | 6.13  | 0.99     | -15.49 | -60.77 | *-3.49 | -7.26 |
| **MAE** | 8.04  | 9.43 | 2.68 | 1.12  | 28.46    | 1.72  | 3.61  |
| **Overall assessment** |       |    |     |       |          |      |      |      |      |
| **MAE** | 4.39  | 5.85 | 1.63 | 0.72  | 7.27     | 16.41 | 1.06  | 1.81  |

C. Energy decomposition analysis

To analyze in more details Laplacian-level kinetic energy functionals, we extend the idea proposed in Ref. 64. Thus, we perform a decomposition of the non-additive KE in terms of the reduced gradient and Laplacian contributions. Similarly with the GGA case, we define the following transformation of the Thomas-Fermi kinetic energy density

$$t[n](s,q) = \int \tau_{TF}[n](r)\delta(s-s(r))\delta(q-q(r))\,dr, \quad (35)$$

so that the Thomas-Fermi kinetic energy is

$$T_s^{TF}[n] = \int \int t[n](s,q)\,ds\,dq, \quad (36)$$

where $t[n](s,q)$ is the $(s,q)$-decomposed homogeneous electron gas (HEG) KE distribution. For a Laplacian-level meta-GGA KE functional we have

$$T_s[F_s,n] = \int \int t[n](s,q)F_s(s,q)\,ds\,dq. \quad (37)$$

Equation (??) states that the total kinetic energy is the scalar product (in the $(s,q)$-space) of $t(s,q)$ and the ki-
with energy error, for given approximated embedded density, cannot be resolved in the total KEs (see Eq. \((\textit{???})\) of the non-additive kinetic energy is linear in the compositional enhancement factor \(s, q\)). Thus, in this formalism \(T_n\) is also a functional of \(F_s\). Then, as the definition of the non-additive kinetic energy is linear in the composing total KEs (see Eq. \((\textit{???})\)), we also obtain

\[
T_n^{\text{add}}[F_s; n_A; n_B] = \int \int t^{\text{add}}[n_A; n_B](s, q)F_s(s, q)\, ds\, dq,
\]

with

\[
t^{\text{add}}[n_A; n_B](s, q) = t[n_A + n_B](s, q) + t[n_A](s, q) - t[n_B](s, q).
\]

Equation \((\textit{???})\) is a generalization of Eq. \((19)\) of Ref. \[64\]. Note that the \((s, q)\)-decomposition is a more powerful tool as compared to the \(s\)-decomposition, because the former can distinguish important density regions: \(s \approx 0, q > 0\) in the bond, \(s \leq 0.4, q < 0\) near the nucleus, which cannot be resolved in the \(s\)-only decomposition.

Following Ref. \[64\], we thus have that the embedding energy error, for given approximated embedded density, can be written as:

\[
\Delta E[F_s] \approx \int \int t^{\text{add}}[n_A; n_B](s, q)F_s(s, q)\, ds\, dq + \Delta W
\]

where \(\Delta W\) is a constant, i.e. a known bifunctional of the embedded densities.

We note that, if a GGA enhancement factor is considered, using \(\int_{-\infty}^{\infty} dx \delta(x) = 1\), Eq. \((\textit{???})\) correctly turns into the GGA expression

\[
\Delta E[F_s] \approx \int F_s(s) \left[ \int t^{\text{add}}[n_A; n_B](s, q) dq \right] ds + \Delta W
\]

\[
= \int F_s(s)t^{\text{add}}[n_A; n_B]_{q}(s)ds + \Delta W,
\]

i.e. the \(q\)-averaged integral non-additive \(s, q\)-decomposed HEG kinetic energy distribution \(t^{\text{add}}[n_A; n_B]_{q}(s)\) corresponds to the non-additive HEG kinetic energy distribution obtained in Ref. \[64\] for the \(s\)-decomposition of a GGA functional (see Eqs. \((14)\) and \((19)\) of Ref. \[64\]).

Equation \((\textit{???})\) can be used to understand the role of \(F_s(s, q)\) and the performance of different Laplacian-meta-GGAs in terms of the shape of their enhancement factor in the \((s, q)\)-space.

As an application of the present \((s, q)\)-decomposition we consider the HF-NCH and Ne₂ complexes. The L0.4 functional gives almost the exact energy for the former (embedding error of only 0.17mHa) whereas it is quite inaccurate for the latter (embedding error larger than the binding energy). Figure \[5\] reports, for the two systems, the non-additive \((s, q)\)-decomposed HEG kinetic energy distribution \(t^{\text{add}}(s, q)\) (upper-panel), as well as the plot of \(t^{\text{add}}_{q}(s)\) (lower panel), both calculated by fixing the embedded densities to the APBEK ones. Integration in Eq. \((\textit{???})\) was performed numerically, representing the delta function with a Gaussian model with broadening \(\sigma=0.07\).

The plots on the bottom panels show that, in analogy with the GGA case studied in Ref. \[64\], the correct embedding energy comes from a delicate balancing of positive contributions at small \(s\) values and negative contributions at larger values of \(s\) in the scalar product (Eq. \((\textit{???})\)). On the other hand, richer structures are present considering \(t^{\text{add}}(s, q)\) and several considerations can be done:

i) Contributions from negative \(q\) are negligible, meaning that the core regions are not important for \(t^{\text{add}}(s, q)\). This information cannot be extracted from the \(s\)-only decomposition because both the core region as well as the valence region have \(s \leq 0.4\).

ii) for HF-NCH the \(q\)-dependence of \(t^{\text{add}}(s, q)\) is quite weak, explaining the success of GGA approximations, which rely on the \(q\)-averaged integral of \(t^{\text{add}}(s, q)\). Moreover, the non-zero values of \(t^{\text{add}}(s, q)\) are mostly confined in the range \(0 \leq q \leq 2\) and \(0 \leq s \leq 2\), which is related to the bonding region (the value of \(q\) at the center of the bond is \(q \approx 0.5\)). In this region of the \(s, q\)-space (small \(s\) and small \(q\)) the fourth-order gradient expansion can be correct, explaining the very low embedding
errors of L0.4 (which recovers GE4). Similar plots have been obtained for other hydrogen-bond or dipole-dipole interaction systems.

iii) For Ne₂, the $t^{\text{nadd}}(s, q)$ looks very different. The most important structures are now at $s > 1.5$ and $4 \leq q \leq 8$. In fact, due to the weakly interacting character of the system, the value of the reduced Laplacian at the center of the bond is $q \approx 6.8$. The plot thus shows how difficult can be the construction of an accurate Laplacian-level kinetic functional for FDE. In fact, the embedding energy error depends on the product over the whole $(s, q)$-space of $t^{\text{nadd}}(s, q)$ and the kinetic enhancement factor $F_s(s, q)$. Hence, an accurate enhancement factor should properly take into account the complexity of the structures at $4 \leq q \leq 8$ and $s > 1.5$. However, the L0.4/L0.6 enhancement factor is only weakly dependent on $q$ for $s > 3$. On the other hand, GGA functionals average over $q$ and loose all $q$-dependent structures: nevertheless, they can still be very accurate for weakly-interaction systems (see e.g. revAPBEK) thanks to an error cancellation in different $q$-regions. This error cancellation is however less likely (and also undesired) for Laplacian-level kinetic functionals.

Finally, the plot of Fig. 6 also explains the fact that MGGA and MGE4 always yield strongly underestimated embedding energies. This fact traces back to the diverging enhancement factor as $s$ increases (see Fig. 1). This exact feature brings an overestimation of the negative contributions of $t^{\text{nadd}}(s, q)$ (always located at large $s$ values) which is not well balanced by the regions responsible for the positive contributions. Thus, too negative embedding energies (even more negative than for GE2) are obtained.

VII. CONCLUSIONS

In this work we investigated the significance of the fourth-order gradient expansion of the kinetic energy. To this end we performed a throughout assessment of several Laplacian-level meta-GGA kinetic energy functionals, with special attention to subsystem DFT applications.

Our study indicated that the inclusion of informations coming from the Laplacian of the density into the functional may play an important role towards a higher accuracy and broader applicability. In fact, GE4 significantly improves over GE2 for solid-state related models (as jel-
lum surfaces, interacting jellium slabs, and large jellium clusters; see Table III as well as for the monovacancy formation in jellium\textsuperscript{11}. Nevertheless, GE4 shows serious drawbacks for small finite systems (e.g., light atoms), due to a non-physical behavior near the nucleus and in the tail of the density, which make the construction of GE4-based functionals a real challenge. In fact, the different Laplacian-level meta-GGAs tested in this paper displayed a quite disappointing unsystematic accuracy, being quite good for some properties and systems and rather poor (at least worse than GGA methods) for others. These problems become especially evident in FDE applications, where the quality of the nonadditive kinetic potential plays a fundamental role, being applied to different densities at the same time. In fact, most of the Laplacian-level meta-GGA functionals considered in the present study perform poorly and several ones also yield severe convergence problems.

The notable exception to this behavior is given by the L0.4 and L0.6 functionals, which perform overall rather close to the best GGAs, especially in embedding calculations of small molecular complexes, while being better than the best GGAs for solid-state related jellium models. The reason for this traces back to the fact that these functionals were constructed not only to recover GE4 in the slowly-varying density limit, but also to achieve a reasonably good behavior in the rapidly-varying regime. This later goal was obtained by mimicking the successful behavior of the (rev)APBEK functional in the rapidly-varying limit. Thus, the L0.4 and L0.6 functionals, appear as promising tools for the application of Laplacian-level meta-GGA kinetic energy functionals in the context of FDE or the density-decomposed orbital-free DFT\textsuperscript{12,13,14}.

We remark, however, that the main message emerging from the present work is that there is still a huge amount of work to be done in the development of Laplacian-level meta-GGA kinetic energy functionals before they can significantly overcome the more simple GGAs for FDE applications to weakly interacting systems. This future work should be focused on studying in deeper details the role played by the Laplacian in different systems and density regimes, so that more complex dependences on the $q$ variable can be developed, especially in the rapidly-varying density regime. In fact, with our assessment work, and in particular through the $(s, q)$-decomposition technique, we showed that the actual Laplacian-level meta-GGA functionals display a reliable $q$-dependence only in the slowly-varying limit, whereas they show limitations in the rapidly-varying regions.

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**Appendix A: Implementation of Laplacian-level meta-GGAs**

For any Laplacian level DFT functional of the form

$$ T_s[n] = \int \tau_s[n, \nabla n, \nabla^2 n](\mathbf{r}) d\mathbf{r} \quad (A1) $$

the functional derivative with respect to the electron density is:

$$ \frac{\delta T_s[n]}{\delta n(\mathbf{r})} = \frac{\partial \tau_s[n](\mathbf{r})}{\partial n(\mathbf{r})} - \nabla \cdot \frac{\partial \tau_s[n](\mathbf{r})}{\partial \nabla n(\mathbf{r})} + \nabla^2 \frac{\partial \tau_s[n](\mathbf{r})}{\partial \nabla^2 n(\mathbf{r})} \quad (A2) $$

The matrix elements between the basis set functions $\{\chi_i\}$ required in FDE calculations are therefore

$$ (v_s)_{\mu\nu} = \int \chi_\mu(\mathbf{r}) \frac{\delta T_s[n]}{\delta n(\mathbf{r})} \chi_\nu(\mathbf{r}) d\mathbf{r} = \int \chi_\mu(\mathbf{r}) \frac{\partial \tau_s[n](\mathbf{r})}{\partial n(\mathbf{r})} \chi_\nu(\mathbf{r}) d\mathbf{r} + \int \frac{\partial \tau_s[n](\mathbf{r})}{\partial \nabla n(\mathbf{r})} \chi_\mu(\mathbf{r}) \chi_\nu(\mathbf{r}) + \int \nabla \chi_\mu(\mathbf{r}) \chi_\nu(\mathbf{r}) + 2\nabla^2 \chi_\mu(\mathbf{r}) \cdot \nabla \chi_\nu(\mathbf{r}) \quad (A3) $$

where we used the first $\left( \int f(\mathbf{r}) \nabla \cdot \mathbf{v}(\mathbf{r}) d\mathbf{r} = -\int \nabla f(\mathbf{r}) \cdot \mathbf{v}(\mathbf{r}) d\mathbf{r} \right)$ and the second Green’s identity $\left( \int f(\mathbf{r}) \nabla^2 g(\mathbf{r}) d\mathbf{r} = \int \nabla^2 f(\mathbf{r}) g(\mathbf{r}) d\mathbf{r} \right)$.
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