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Nonlocal van der Waals density functional: The simpler the better

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We devise a nonlocal correlation energy functional that describes the entire range of dispersion interactions in a seamless fashion using only the electron density as input. The new functional is considerably simpler than its predecessors of a similar type. The functional has a tractable and robust analytic form that lends itself to efficient self-consistent implementation. When paired with an appropriate exchange functional, our nonlocal correlation model yields accurate interaction energies of weakly-bound complexes, not only near the energy minima but also far from equilibrium. Our model exhibits an outstanding precision at predicting equilibrium intermonomer separations in van der Waals complexes. It also gives accurate covalent bond lengths and atomization energies. Hence the functional proposed in this work is a computationally inexpensive electronic structure tool of broad applicability.

I. INTRODUCTION

In Kohn-Sham density functional theory (DFT), only one contribution to the ground state energy is not known exactly — the correlation energy. Common approximations to the correlation energy have the form of local or semilocal density functionals. One of the numerous limitations of (semi)local functionals is their inherent inability to describe such long-range correlation effects as dispersion (van der Waals) interactions. Empirical dispersion corrections are quite popular and reasonably successful, but they typically entail a departure from pure DFT into the realm of classical force fields.

The proper physics of long-range van der Waals interactions can be captured with fully nonlocal correlation functionals. However, the rigor usually comes at the cost of an explicit and cumbersome dependence on Kohn-Sham orbitals, both occupied and virtual, as exemplified by the random-phase approximation and related methods (see e.g. Refs. 6 and 7 and references therein). An elegant compromise between rigor and computational tractability is achieved in the recently introduced class of nonlocal correlation functionals that treat the entire range of dispersion interactions in a general and seamless fashion, yet include no explicit orbital dependence and use only the electron density as input. In this article we devise a nonlocal van der Waals functional that is considerably simpler yet more accurate than any of its precursors. We give no derivation for the new functional form, but we show that it recovers all the relevant limits and satisfies many exact constraints. We suggest a suitable combination of exchange and correlation terms that performs remarkably well in all our benchmark tests.

II. FORMALISM

We write the nonlocal correlation energy as

$$E_{\text{c}}^{\text{nl}} = \frac{\hbar}{2} \int \int \text{d}r \text{d}r' n(r) \Phi(r, r') n(r'),$$

(1)

where $n$ is the total electron density. Building upon the insights gained in Refs. 10–13, we write the correlation kernel as

$$\Phi = \frac{3e^4}{2m^2 g' (g + g')}$$

(2)

with

$$g = \omega_0(r) R^2 + \kappa(r)$$

(3)

and similarly

$$g' = \omega_0(r') R^2 + \kappa(r').$$

(4)

In the above equations, $R = |r - r'|$ and

$$\omega_0(r) = \sqrt{\omega_0^2(r) + \frac{\omega_2^2(r)}{3}},$$

(5)

with the local plasma frequency defined via $\omega_p^2 = 4\pi ne^2/m$ and the local band gap given by

$$\omega_2^2(r) = C \frac{\hbar^2}{m^2} \left| \frac{\nabla n(r)}{n(r)} \right|^4,$$

(6)

where $C$ is a parameter adjusted to give accurate asymptotic van der Waals $C_6$ coefficients, as elaborated in Section IV.

In the $R \to \infty$ limit,

$$\Phi \to -\frac{3e^4}{2m^2 \omega_0(r) \omega_0(r') [\omega_0(r) + \omega_0(r')] R^6},$$

(7)

so that $E_{\text{c}}^{\text{nl}}$ of Eq. (1) has precisely the same long-range behavior as the nonlocal energy in VV09 of Ref. 11. A detailed analysis and physical justification of this asymptotic form was given in Ref. 13.

For $R \to 0$, the correlation kernel of Eq. (2) behaves as

$$\Phi = -A + BR^2 + \ldots,$$

(8)
in accordance with the result of Koide for the $R \to 0$ asymptotic behavior of the dispersion interaction energy. In Eqs. (3) and (4), we introduced the quantity

$$\kappa(r) = b \frac{v_F^2(r)}{\omega_p(r)} = 3 b \frac{\omega_p(r)}{k_s^2(r)},$$

(9)

where $v_F = (3\pi^2 n)^{1/3} \hbar/m$ is the local Fermi velocity, $k_s = \sqrt{3} \omega_p/v_F$ is the Thomas–Fermi screening wave vector, and $b$ is an adjustable parameter that controls the short-range damping of the $R^{-6}$ asymptote.

In the uniform density limit, Eq. (2) reduces to

$$\Phi^{uni} = -\frac{3e^4}{4m^2} \left[ \frac{\omega_p}{\sqrt{3}} R^2 + b \frac{v_F^2}{\omega_p} \right]^{-3},$$

(10)

and Eq. (11) gives the following energy density per electron:

$$\varepsilon_r^{uni} = 2\pi \hbar n \int_0^\infty R^2 \Phi^{uni} dR = -3\pi^2 \hbar e^4 n \left[ \frac{3}{32 m^2 v_F^3} \right]^{3/4}$$

$$= -\frac{e^2}{32 a_0} \left[ \frac{3}{b^2} \right]^{3/4} = -\beta,$$

(11)

where $a_0 = \hbar^2/m e^2$ is the Bohr radius and $\beta$ is a density-independent constant.

It is instructive to rewrite Eq. (10) in a different form:

$$\Phi^{uni} = -\frac{9e^4 \sqrt{3}}{4m^2 \omega_p R^6} \left[ 1 + \frac{3\sqrt{3} b}{(k_s R)^2} \right]^{-3}.$$  

(12)

The above equation shows that the $R^{-6}$ asymptote is damped at short range on the length scale given by $k_s R$.

Finally, we define our van der Waals density functional as

$$E_c^{VV10} = E_c^{eq} + \beta N$$

$$= \int \text{d}r \ n(r) \left[ \beta + \frac{\hbar}{2} \int \text{d}r' n(r') \Phi(r, r') \right],$$

(13)

where $N = \int \text{d}r n(r)$ is the number of electrons and $\beta$ is determined by Eq. (11). By construction, $E_c^{VV10}$ of Eq. (13) vanishes in the uniform density limit. This is a useful property: we can pair $E_c^{VV10}$ with an existing exchange-correlation (XC) functional without affecting the description of the uniform electron gas.

It is a nontrivial task to determine how Eq. (13) behaves in the slowly varying density limit. At any rate, it has been argued that recovery of the density gradient expansion for correlation energy is of little relevance for real systems.

The van der Waals functional of Eq. (13) is very simple and easily implementable. The double integral in Eq. (11) is in practice evaluated as a double sum over a numerical grid. For $R \to 0$, the kernel of Eq. (2) goes to a constant, avoiding any complications with the numerical integration. In the inner integration loop, only simple arithmetic operations need to be performed. Hence this new functional is computationally less expensive than the nonlocal functionals of Refs. [8][11].

The previous version of the nonlocal van der Waals functional, VV09 proposed by us in Ref. [11], was more computationally demanding because it used a rather elaborate damping function. Evaluation of the nonlocal correlation energy in VV09 required computing a square root, an exponent, and an error function for each pair of grid points in the double sum. The energy derivatives, needed for the self-consistent implementation of VV09, were also quite complicated. With the new VV10 model of Eq. (13), these energy derivatives are much simplified. The self-consistent implementation of VV10 within a Gaussian basis set code is described in detail in the next section.

### III. IMPLEMENTATION

For the sake of brevity, in this section we switch to atomic units ($\hbar = m = e = 1$). In these units, $\kappa$ of Eq. (9) is expressed rather simply as

$$\kappa(r) = b \frac{3 \pi^2 n(r)}{2 \sqrt{9\pi}}$$

(14)

Our implementation of VV10 is very similar to the implementation of its predecessor VV09, described in Ref. [16]. We express the electron density via a Gaussian basis set:

$$n(r) = \sum_{\mu \nu} P_{\mu \nu} \chi_\mu(r) \chi_\nu(r),$$

(15)

where $\chi_\mu$ and $\chi_\nu$ are basis functions, and $P_{\mu \nu}$ are the density matrix elements. For the self-consistent treatment, we need to find the derivatives of $E_c^{VV10}$ with respect to $P_{\mu \nu}$:

$$\frac{dE_c^{VV10}}{dP_{\mu \nu}} = \int \text{d}r \chi_\mu(r) \frac{\delta E_c^{VV10}}{\delta n(r)} \chi_\nu(r).$$

(16)

To that end, we employ the standard formalism developed for semilocal XC functionals:

$$\frac{dE_c^{VV10}}{dP_{\mu \nu}} = \int \text{d}r \left[ F_n \chi_\mu \chi_\nu + 2 F_\gamma \nabla n \cdot \nabla (\chi_\mu \chi_\nu) \right].$$

(17)

We denote $\gamma = |\nabla n|^2$ for convenience. $F_n$ and $F_\gamma$ in Eq. (17) are computed as

$$F_n(r) = \beta + \int \text{d}r' n(r') \Phi$$

$$+ n(r) \left[ \frac{\partial \kappa}{\partial n}(r) U(r) + \frac{\partial \omega_0}{\partial n}(r) W(r) \right]$$

(18)

and

$$F_\gamma(r) = n(r) \frac{\partial \omega_0}{\partial \gamma}(r) W(r),$$

(19)
where

\[ U(r) = -\int dr' n(r') \Phi \left[ \frac{1}{g} + \frac{1}{g + g'} \right] \]  

and

\[ W(r) = -\int dr' n(r') |r - r'|^2 \Phi \left[ \frac{1}{g} + \frac{1}{g + g'} \right]. \]  

In the above equations, \( \Phi, g, \) and \( g' \) are assumed to be functions of both \( r \) and \( r' \).

Within an atom-centered basis set implementation, the gradient of \( E^\mathrm{VV10}_c \) with respect to the displacement of nucleus \( A \) has three contributions:

\[ \nabla A E^\mathrm{VV10}_c = g^A_{\text{GBF}} + g^A_{\text{weights}} + g^A_{\text{grid}}. \]  

\( g^A_{\text{GBF}} \) denotes the contribution of the Gaussian basis functions. This term can be evaluated by plugging \( F_n \) and \( F_\gamma \) into Eq. (9) of Ref. \[17\] instead of \( \partial f/\partial n \) and \( \partial f/\partial \gamma \).

The last two terms in Eq. (22) are due to the use of atom-centered numerical integration quadratures. We employ the atomic partitioning scheme of Becke,\[18\] which separates the molecular integral into atomic contributions:

\[ E^\mathrm{VV10}_c = \sum_A \sum_{i \in A} w_{Ai} n(r_{Ai}) \left[ \beta \right. \]

\[ + \frac{1}{2} \sum_B \sum_{j \in B} w_{Bj} n(r_{Bj}) \Phi(r_{Ai}, r_{Bj}) \left. \right]. \]  

where \( w_{Ai} \) and \( w_{Bj} \) are the quadrature weights, and the grid points \( r_{Ai} \) are given by \( r_{Ai} = R_A + r_i \), where \( R_A \) is the position of nucleus \( A \), with the \( r_i \) defining a one-center integration grid. The quadrature weights depend on the nuclear configuration and hence have a nonzero gradient with respect to nuclear displacements:

\[ g^A_{\text{weights}} = \sum_B \sum_{i \in B} (\nabla_A w_{Bi}) n(r_{Bi}) \left[ \beta \right. \]

\[ + \sum_C \sum_{j \in C} w_{Cj} n(r_{Cj}) \Phi(r_{Bi}, r_{Cj}) \left. \right]. \]  

The weight derivatives \( \nabla_A w_{Bi} \) can be found in Ref. \[17\].

The last term in Eq. (22) arises because each one-center quadrature grid moves together with its parent nucleus and the nonlocal correlation kernel \( \Phi \) depends explicitly on the distance between the grid points. The \( g^A_{\text{grid}} \) term can be computed as:

\[ g^A_{\text{grid}} = \sum_{i \in A} w_{Ai} n(r_{Ai}) \]

\[ \times \sum_{B \neq A} \sum_{j \in B} w_{Bj} n(r_{Bj}) Q(r_{Ai}, r_{Bj}) |r_{Ai} - r_{Bj}|, \]  

where

\[ Q(r, r') = -2\Phi \left[ \frac{\omega_0(r)}{g} + \frac{\omega_0(r')}{g'} + \frac{\omega_0(r) + \omega_0(r')}{g + g'} \right]. \]  

Analytic gradients, computed via Eq. (22), enable us to perform structural optimizations routinely and efficiently.

IV. ADJUSTMENTS

\( E^\mathrm{VV10}_c \) of Eq. (13) can in principle be paired with nearly any existing XC functional. However, to avoid double-counting as much as possible, it is preferable to combine \( E^\mathrm{VV10}_c \) with a functional that gives no significant binding in van der Waals complexes. Since predictions of XC functionals for weakly interacting systems can differ drastically, the parameter \( b \), controlling the short-range behavior of \( E^\mathrm{VV10}_c \), has to be adjusted for a particular parent XC approximation.

The Perdew–Wang 86 (PW86) exchange functional\[19\] has been shown to describe the repulsive parts of van der Waals potentials rather well,\[20,21\] and a refitted version of PW86 was recently proposed.\[22\] We denote this ‘refitted PW86’ as RPW86 here. In the rest of this article we will consider one particularly apt combination of the exchange and correlation terms:

\[ E^{\mathrm{VV10}_c} = E^\mathrm{epW86}_c + E^\mathrm{pBE}_c + E^\mathrm{VV10}_c, \]  

where \( E^\mathrm{pBE}_c \) is the semi-local correlation energy functional in the generalized gradient approximation of Perdew, Burke, and Ernzerhof (PBE)\[23\]. Hereafter, we will often refer to the XC functional of Eq. (27) simply as VV10.

There are two adjustable parameters in our method, but only \( C \) in Eq. (6) affects the asymptotic dispersion \( C_0 \) coefficients. We fit \( C \) to minimize the average error for the benchmark set of 54 \( C_6 \) coefficients compiled in Tables II and III of Ref. \[13\]. Using self-consistent electron densities from RPW86-PBE (i.e. using \( E_{xc} = E^\mathrm{pW86} + E^\mathrm{pBE}_c \)), we find the optimal value of \( C = 0.0093 \), which gives the mean absolute percentage error (MAPE) of 14% for the whole test set of 54 species. The largest errors in \( C_6 \) occur for metal atoms. When all the metal atoms (Li, Na, Mg, Al, Zn, Ga) are removed from the benchmark set and only the remaining 48 species are considered, the MAPE in \( C_6 \) drops to 9%. Note that a slightly smaller value of the parameter \( C \) was used in Refs. \[11, 13\], since the source of electron densities was different.\[24\]

Another adjustable parameter \( b \), introduced in Eq. (10), controls the short range damping of \( \Phi \). Using \( E^{\mathrm{VV10}_c} \) of Eq. (27), we fitted \( b \) on the training set of 22 binding energies (the S22 set of Ref. \[23\]), with the computational details described in Section IV and the results given in Section VI A. We obtained the best fit with \( b = 5.9 \). Correspondingly, in Eq. (19) we use

\[ \beta = \frac{1}{32} \left[ \frac{3}{5.9^2} \right]^{3/4} = 0.00497 \]
where $E_{\text{c}}^{\text{LDA}}$ is the correlation energy in the local density approximation, for which we use the parameterization of Perdew and Wang.\textsuperscript{28}

All reported calculations were performed self-consistently. For weakly bound systems, the interaction energies were counterpoise corrected. The unpruned Euler–Maclaurin–Lebedev (99,590) quadrature grid was used to evaluate all local and semilocal contributions ($E_x^\text{PW86}$, $E_c^{\text{PBE}}$, and $E_{\text{c}}^{\text{LDA}}$) and the (75,302) grid was used for the nonlocal components. Deviations of computed binding energies from the reference values are analyzed with the help of mean errors (ME), mean absolute errors (MAE), and mean absolute percentage errors (MAPE). In all tables, binding energies are reported as positive values. Hence a negative ME indicates underbinding while a positive ME means overbinding.

In our recent article\textsuperscript{16} assessing the performance of VV09, we used the same benchmark sets of weakly bound systems and the same computational details. Therefore, the results reported in Ref.\textsuperscript{16} can be directly compared to the results obtained with the new functionals, presented in Section \textbf{VI} below.

\section{RESULTS FOR WEAKLY INTERACTING SYSTEMS}

\subsection{The S22 benchmark set}

Binding energies for the S22 test set, computed with vdW-DF2 and VV10 at the geometries of Ref.\textsuperscript{25} are given in Table \textbf{I}. The error statistics are summarized in Table \textbf{II}. Updated reference values of binding energies for the S22 set were recently obtained in Refs.\textsuperscript{29} and \textsuperscript{30}. For our benchmark set, we picked the values that we deemed the most converged with respect to the basis set size: we took the reference values for most of the systems with aug-cc-pVTZ basis set and counterpoise correction. Molecular structures are from Ref.\textsuperscript{25} and reference binding energies are from Ref.\textsuperscript{28} for both of them. The full XC energy is presented in Section \textbf{VI} below.

\begin{table}[h]
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\begin{tabular}{lccc}
\hline
Complex (symmetry) & Ref. & VV10 & vdW-DF2 \\
\hline
\textit{Dispersion-bound complexes} (8) & & & \\
CH$_4$ dimer ($D_{2d}$) & 0.53 & 0.50 & 0.68 \\
C$_2$H$_4$ dimer ($D_{2d}$) & 1.48 & 1.42 & 1.32 \\
Benzenes–CH$_4$ ($C_2v$) & 1.45 & 1.45 & 1.29 \\
Benzenes dimer ($C_{2h}$) & 2.66 & 2.71 & 2.15 \\
Pyrazine dimer ($C_{2h}$) & 4.26 & 4.02 & 3.30 \\
Uracil dimer ($C_{2h}$) & 9.78 & 9.70 & 8.76 \\
Indole–Benzene ($C_{2h}$) & 4.52 & 4.54 & 3.44 \\
Adenine–Thymine ($C_{2h}$) & 11.66 & 11.42 & 9.58 \\
\textit{Mixed complexes} (7) & & & \\
C$_2$H$_4$–C$_2$H$_2$ ($C_{2h}$) & 1.50 & 1.68 & 1.53 \\
Benzenes–H$_2$O ($C_2v$) & 3.28 & 3.31 & 2.80 \\
Benzenes–NH$_3$ ($C_2v$) & 2.32 & 2.28 & 1.99 \\
Benzenes–HCN ($C_2v$) & 4.54 & 4.30 & 3.55 \\
Benzenes dimer ($C_{2v}$) & 2.72 & 2.54 & 2.06 \\
Indole–Benzene ($C_{2v}$) & 5.63 & 5.27 & 4.20 \\
Phenol dimer ($C_1$) & 7.10 & 6.99 & 5.97 \\
\textit{Hydrogen-bonded complexes} (7) & & & \\
NH$_3$ dimer ($C_{2h}$) & 3.15 & 3.43 & 2.97 \\
H$_2$O dimer ($C_2v$) & 5.00 & 5.50 & 4.78 \\
Formic acid dimer ($C_{2h}$) & 18.75 & 19.96 & 16.77 \\
Formamidine dimer ($C_{2h}$) & 16.06 & 16.71 & 14.43 \\
Uracil dimer ($C_{2v}$) & 20.64 & 21.10 & 18.69 \\
2-pyridone–2-aminoypyridine ($C_1$) & 16.94 & 18.05 & 15.37 \\
Adenine–Thymine WC ($C_1$) & 16.74 & 17.42 & 14.74 \\
\hline
\end{tabular}
\caption{Binding energies (in kcal/mol) for the S22 test set. All calculations were performed self-consistently with the aug-cc-pVTZ basis set and counterpoise corrected. Molecular structures are from Ref.\textsuperscript{25} and reference binding energies are from Ref.\textsuperscript{28} for all systems except Adenine–Thymine complexes, for which the numbers from Ref.\textsuperscript{30} were used.}
\end{table}

\begin{table}[h]
\centering
\begin{tabular}{lccc}
\hline
& VV10 & vdW-DF2 \\
\hline
\textit{Dispersion-bound complexes} (8) & & & \\
ME & −0.07 & −0.73 \\
MAE & 0.09 & 0.76 \\
MAPE (%) & 2.6 & 18.0 \\
\textit{Mixed complexes} (7) & & & \\
ME & −0.10 & −0.71 \\
MAE & 0.16 & 0.72 \\
MAPE (%) & 4.8 & 16.8 \\
\textit{Hydrogen-bonded complexes} (7) & & & \\
ME & 0.70 & −1.36 \\
MAE & 0.70 & 1.36 \\
MAPE (%) & 6.1 & 8.8 \\
\textit{Total} (22) & & & \\
ME & 0.16 & −0.92 \\
MAE & 0.31 & 0.94 \\
MAPE (%) & 4.4 & 14.7 \\
\hline
\end{tabular}
\caption{Summary of deviations from the reference values of the binding energies reported in Table \textbf{I}. ME and MAE are in kcal/mol, MAPE is in percents.}
\end{table}
from Ref. [30] except for the Adenine–Thymine complexes (in both stacked and planar configurations), for which we used the numbers from Ref. [31].

As Tables I and II show, VV10 performs remarkably well for all dispersion-bound and mixed complexes, but yields somewhat larger errors for hydrogen-bonded systems. The parameter $b$ inside $E_{VV10}^c$ was fitted to the binding energies of the S22 set, but it is by no means trivial to achieve such accuracy by adjusting just one global parameter.

vdW-DF2 overbinds some of the smaller systems, specifically $\text{CH}_4$ and $\text{C}_2\text{H}_4$–$\text{C}_2\text{H}_2$, but underbinds all other complexes in the S22 set, when the geometries from Ref. [29] are used. This underbinding tendency intensifies as the monomer sizes increase. A negative ME for the binding energies of the S22 set was also found in Ref. [9]. However, intermonomer separations were optimized in Ref. [9], yielding larger binding energies, as compared to our results for the fixed geometries.

Binding energies calculated for a standard set of near-equilibrium geometries certainly do not tell the full story about the performance of a functional. For a more comprehensive assessment, it is necessary to determine whether equilibrium intermonomer separations are accurately located and whether reasonable interaction energies are predicted far from equilibrium. To this end, we analyze binding energy curves for several complexes.

### B. Interaction energy curves

We computed interaction energy curves for six weakly-bound systems using VV10 and vdW-DF2. Figure 1 shows the results for the argon and krypton dimers, Figure 2 for the methane dimer ($D_{3d}$ symmetry) and the benzene–methane complex ($C_{3v}$), and Figure 3 for benzene dimers in two configurations: T-shaped ($C_{2v}$) and stacked sandwich-shaped ($D_{hb}$). We applied counterpoise corrections at all intermonomer distances. For Ar$_2$ and Kr$_2$, the reference values were taken from Ref. [31], and for the other complexes from Ref. [32]. For molecular complexes, we used the same fixed monomer geometries as in Ref. [32].

For all six complexes, VV10 reproduces the equilibrium intermonomer separations with remarkable precision. Interaction energies yielded by VV10 agree quite well with the reference values at all separations. The well depths are predicted with reasonable accuracy: VV10 slightly underbinds $\text{CH}_4$ and the T-shaped ($C_2h$) benzene–methane complex ($D_{3d}$), but it slightly overbinds Ar$_2$ and Kr$_2$ and more strongly overbinds the sandwich-shaped ($C_6h$). For the benzene–methane complex, the deviations of VV10 from the reference curve are within the uncertainties of the reference interaction energies at nearly all separations. The long-range falloff of the potential energy curves is reproduced very well by VV10 in nearly all cases.

vdW-DF2 tends to predict the energy minima at somewhat too large separations. It significantly overbinds the complexes of small monomers: Ar$_2$, Kr$_2$, and $\text{CH}_4$. For all systems in Figures 1–3 and 5 except Ar$_2$, the repulsive walls given by vdW-DF2 are too steep and shifted towards larger distances as compared to the reference curves. As was shown in Ref. [13], vdW-DF2 strongly underestimates asymptotic $C_6$ coefficients. As a result, for systems where the asymptotic interactions are dominated by dispersion, vdW-DF2 will yield too shallow potential energy curves at long range. This underestimation is vis-

![Figure 1](image1.png)

**FIG. 1.** Interaction energy curves for (a) the argon dimer and (b) the krypton dimer. Reference curves are from Ref. [31]. VV10 and vdW-DF2 results were obtained self-consistently with the aug-cc-pVQZ basis set.
FIG. 2. Interaction energy curves for (a) the methane dimer and (b) the benzene–methane complex. $R$ is the distance between the centers of mass of the monomers. Reference values are from Ref. 32. VV10 and vdW-DF2 results were obtained self-consistently with the aug-cc-pVTZ basis set.

FIG. 3. Interaction energy curves for (a) the T-shaped and (b) the stacked sandwich-shaped benzene dimers. $R$ is the distance between the centers of mass of the monomers. Reference values are from Ref. 32. VV10 and vdW-DF2 results were obtained self-consistently with the aug-cc-pVTZ basis set.

ible at large separations for Ar$_2$, Kr$_2$, and the T-shaped (C$_6$H$_6$)$_2$. The effects of the poor vdW-DF2 asymptotics are expected to be more noticeable at very large intermonomer distances or for very large monomers. Such cases are not significantly represented in this study.

As Figure 3(b) shows, both VV10 and vdW-DF2 significantly overestimate the equilibrium binding energy in the sandwich-shaped benzene dimer. This may be partially caused by the lack of Axilrod-Teller-type three-body interactions in the functionals of the form of Eq. (1). Another possible source of errors is the rPW86 exchange functional. It was found in Ref. 22 that PW86 exchange is more repulsive than Hartree–Fock for the T-shaped benzene dimer, but less repulsive for the sandwich-shaped configuration.
VII. RESULTS FOR ATOMS AND COVALENT BONDS

We have shown that VV10 describes weakly-bound systems quite well. In this section, we show that VV10 is a broadly applicable method that also treats covalent and ionic bonds well.

A. Total energies of atoms

In Table III we report the total energies of atoms up to Ar, computed with rPW86-PBE (that is rPW86 exchange with PBE correlation) as well as VV10, which is rPW86-PBE with the addition of $E^{\text{VV10}}_c$ as shown in Eq. (27). We also include the popular semilocal XC functional PBE. We used the aug-cc-pVQZ basis set with all beyond-$f$ polarization functions removed. This basis set is expected to yield energies close to the basis set limit. As the benchmark, we use the accurate non-relativistic values of atomic energies from Refs. [33] and [34].

Among the three considered functionals, PBE gives the most accurate atomic energies. PBE total energies are systematically above the exact values, except for the H atom, for which PBE gives a nearly exact energy. On the contrary, rPW86-PBE and VV10 yield total energies that are systematically too low.

We find $E^{\text{VV10}}_c$ of Eq. (22) to be positive not only for atoms, but for all systems considered in this study. We have no proof that $E^{\text{VV10}}_c$ is always strictly nonnegative, i.e. that $\beta N \geq -E^{\text{nl}}_c$, but we have found no exceptions so far. The positive contribution of $E^{\text{VV10}}_c$ brings the rPW86-PBE total energies in better agreement with the exact values, as Table III shows.

We did not include vdW-DF2 in Table III because this functional is not defined for open-shell systems. For the five closed-shell atoms in the set of Table III, vdW-DF2 yields the ME of $-0.048$ hartree per electron. vdW-DF2 total energies are much too low primarily because this functional includes $E^{\text{vdW}}_c$ as a constituent and LDA is known to severely overestimate the magnitudes of correlation energies in atoms.

B. Atomization energies

We computed atomization energies for the AE6 set developed by Lynch and Truhlar. This set includes only six molecules, but it is quite diverse and was constructed to be representative, that is to reproduce mean errors in much larger sets. We used the molecular geometries suggested by the authors of the set. The results and the mean errors are given in Table IV.

rPW86-PBE, which is the backbone of VV10, yields considerably more accurate atomization energies as compared to the popular PBE functional. The addition of the nonlocal correlation term $E^{\text{VV10}}_c$ to rPW86-PBE increases the atomization energies in all cases, as the VV10 results in Table IV show. In other words, the addition of $E^{\text{VV10}}_c$ makes covalent bonds somewhat stronger. As a result, VV10 yields smaller errors than rPW86-PBE for three molecules in the AE6 set, but larger errors for the other three molecules. On average, the performance of VV10 for the AE6 set is quite similar to its parent functional rPW86-PBE.

C. Bond lengths

In Table V we have compiled a test set of 20 small closed-shell molecules for which experimental equilibrium bond lengths ($r_e$) are known. The test set consists mostly of diatomics, but includes several polyatomic molecules of high symmetry whose geometry is completely determined by a single bond length. Most of the molecules in the set are covalently bound, except for the...
**TABLE V.** Equilibrium bond lengths \((r_e)\) computed using the aug-cc-pVTZ basis set. All values are in Å. Experimental data are from Ref. 37.

<table>
<thead>
<tr>
<th></th>
<th>Expt.</th>
<th>PBE</th>
<th>rPW86-LDA</th>
<th>vdW-DF2</th>
<th>rPW86-PBE</th>
<th>VV10</th>
</tr>
</thead>
<tbody>
<tr>
<td>(H_2)</td>
<td>0.741</td>
<td>0.751</td>
<td>0.734</td>
<td>0.736</td>
<td>0.745</td>
<td>0.745</td>
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<tr>
<td>(LiF)</td>
<td>1.564</td>
<td>1.583</td>
<td>1.592</td>
<td>1.592</td>
<td>1.586</td>
<td>1.585</td>
</tr>
<tr>
<td>(LiCl)</td>
<td>2.021</td>
<td>2.028</td>
<td>2.041</td>
<td>2.041</td>
<td>2.031</td>
<td>2.031</td>
</tr>
<tr>
<td>(Li_2)</td>
<td>2.673</td>
<td>2.726</td>
<td>2.663</td>
<td>2.670</td>
<td>2.699</td>
<td>2.700</td>
</tr>
<tr>
<td>(CH_4)</td>
<td>1.087</td>
<td>1.096</td>
<td>1.088</td>
<td>1.090</td>
<td>1.092</td>
<td>1.093</td>
</tr>
<tr>
<td>(CO)</td>
<td>1.128</td>
<td>1.137</td>
<td>1.136</td>
<td>1.136</td>
<td>1.137</td>
<td>1.137</td>
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<tr>
<td>(CO_2)</td>
<td>1.160</td>
<td>1.172</td>
<td>1.174</td>
<td>1.173</td>
<td>1.172</td>
<td>1.172</td>
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<tr>
<td>(CS_2)</td>
<td>1.553</td>
<td>1.566</td>
<td>1.573</td>
<td>1.571</td>
<td>1.568</td>
<td>1.567</td>
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<tr>
<td>(N_2)</td>
<td>1.098</td>
<td>1.103</td>
<td>1.101</td>
<td>1.101</td>
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<td>1.102</td>
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<tr>
<td>(HF)</td>
<td>0.917</td>
<td>0.932</td>
<td>0.934</td>
<td>0.935</td>
<td>0.934</td>
<td>0.934</td>
</tr>
<tr>
<td>(F_2)</td>
<td>1.412</td>
<td>1.413</td>
<td>1.459</td>
<td>1.453</td>
<td>1.433</td>
<td>1.432</td>
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<tr>
<td>(NaCl)</td>
<td>2.361</td>
<td>2.385</td>
<td>2.410</td>
<td>2.401</td>
<td>2.390</td>
<td>2.385</td>
</tr>
<tr>
<td>(NaBr)</td>
<td>2.502</td>
<td>2.528</td>
<td>2.560</td>
<td>2.547</td>
<td>2.538</td>
<td>2.532</td>
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<tr>
<td>(Na_2)</td>
<td>3.079</td>
<td>3.079</td>
<td>3.006</td>
<td>2.987</td>
<td>3.035</td>
<td>3.024</td>
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<tr>
<td>(SiO)</td>
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<td>1.536</td>
<td>1.543</td>
<td>1.540</td>
<td>1.539</td>
<td>1.538</td>
</tr>
<tr>
<td>(P_2)</td>
<td>1.893</td>
<td>1.911</td>
<td>1.922</td>
<td>1.918</td>
<td>1.914</td>
<td>1.913</td>
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<tr>
<td>(Cl_2)</td>
<td>1.275</td>
<td>1.291</td>
<td>1.289</td>
<td>1.290</td>
<td>1.290</td>
<td>1.290</td>
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<tr>
<td>(HCl)</td>
<td>1.988</td>
<td>2.019</td>
<td>2.085</td>
<td>2.073</td>
<td>2.040</td>
<td>2.037</td>
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<tr>
<td>(HBr)</td>
<td>1.415</td>
<td>1.433</td>
<td>1.436</td>
<td>1.437</td>
<td>1.435</td>
<td>1.435</td>
</tr>
<tr>
<td>(Br_2)</td>
<td>2.281</td>
<td>2.311</td>
<td>2.396</td>
<td>2.380</td>
<td>2.343</td>
<td>2.336</td>
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<tr>
<td>ME</td>
<td>0.017</td>
<td>0.024</td>
<td>0.021</td>
<td>0.021</td>
<td>0.018</td>
<td>0.017</td>
</tr>
<tr>
<td>MAE</td>
<td>0.017</td>
<td>0.033</td>
<td>0.031</td>
<td>0.031</td>
<td>0.023</td>
<td>0.022</td>
</tr>
</tbody>
</table>

alkali metal halides where the bonding is ionic.

Table V reports the results of geometry optimizations carried out using analytic gradients and the aug-cc-pVTZ basis set. We tested vdW-DF2 and VV10 alongside their semilocal parent functionals, rPW86-LDA and rPW86-PBE respectively. We also included PBE for the sake of comparison. For most molecules in Table V, vdW-DF2 yields nearly the same bond lengths as rPW86-LDA, and VV10 nearly the same as rPW86-PBE. Hence the addition of a nonlocal correlation term has an almost negligible effect on covalent and ionic bond lengths. The largest change occurs for Na\(_2\): the Na–Na bond length given by VV10 is 0.011 Å shorter than with rPW86-PBE. This is not surprising since the Na–Na bond is weaker than most covalent bonds. On average for the test set of Table V, the performance of VV10 is on par with PBE, while vdW-DF2 is somewhat less accurate.

**VIII. CONCLUSIONS**

The VV10 correlation model proposed in this work belongs to the family\(^8–11\) of nonlocal van der Waals density functionals defined in terms of the electron density alone and using no orbital input. These functionals\(^8–11\) are general and seamless: they require neither splitting the system into interacting fragments nor any kind of atomic partitioning. They treat inter- and intra-molecular dispersion interactions on equal footing.

VV10 has the same long-range behavior as its predecessor VV09, but the damping mechanism of dispersion interactions at short range is greatly simplified in VV10. This simplification not only makes the functional more efficient and computationally tractable, but it also leads to improved overall accuracy.

An essential aspect of the VV10 formalism is the additional flexibility introduced with the help of an adjustable parameter \(b\) which controls the short range behavior of the nonlocal correlation energy. When \(E_{\text{VV10}}\) of Eq. (13) is added as a correction to an existing XC functional, \(b\) is adjusted to attain a balanced merging of interaction energy contributions at short and intermediate ranges. A particularly successful functional is constructed by adding \(E_{\text{VV10}}\) with \(b = 5.9\) to rPW86-PBE, as shown in Eq. (27). Another parameter \(C = 0.0093\) inside \(E_{\text{VV10}}\) ensures that accurate asymptotic \(C_6\) coefficients are obtained.\(^{24}\)

As our benchmark tests clearly demonstrate, the functional of Eq. (27) is a broadly accurate electronic structure tool: its outstanding predictive power for weakly-bound systems is complemented by its good description of covalent bonds.

**ACKNOWLEDGMENTS**

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**Appendix: VV10 with long-range corrected exchange**

The XC functional of Eq. (27) includes the semilocal rPW86 exchange term as a constituent. Semilo-
The two parameters inside $E_{\text{xc}}$ are adjusted using the same procedure as described in Section IV. As we previously found, $C = 0.0089$ gives accurate $C_6$ coefficients at LC-ωPBE electron densities. We fit the parameter $b$ to the binding energies of the S22 set using the functional of Eq. (A.1) and obtain the optimal value of $b = 6.3$.

In Table VI we summarize the errors of LC-VV10 for the test sets used previously in this article. As compared to the VV10 model with the rPW86 exchange, LC-VV10 performs somewhat better for binding energies of hydrogen bonded complexes, for total energies of atoms, and for bond lengths. However, no improvement is observed for atomization energies, while binding energies of dispersion-bound complexes are slightly worsened on average. Therefore, we suggest using LC-VV10 only when SIE is an issue. In most other cases the VV10 model of Eq. (27) is preferable since the semilocal rPW86 exchange is computationally cheaper and much more easy to implement than an LC hybrid.

It is noteworthy that our model can achieve good performance using a pre-existing and unmodified exchange functional (rPW86 or LC-ωPBE) and adjusting only the parameters in the nonlocal correlation. It might be possible to further improve the performance by tailoring an empirical exchange functional for our specific purpose, in the vein of the recent works of Refs. 12–14. However, we prefer to keep empirical fitting to a minimum.

24 The value of $C = 0.0089$ was obtained in Ref. 11 by fitting to a $C_6$ test set using electron densities produced by the long-range corrected (LC) hybrid functional LC-ωPBE. As compared to LC hybrids, the description of density tails by semilocal XC functionals is systematically less accurate, which is reflected in $\omega$. The readjusted value of $C = 0.0093$ is tailored for use with semilocal XC functionals, such as rPW86-PBE.


