Perspective: Advances and challenges in treating van der Waals dispersion forces in density functional theory

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Electron dispersion forces play a crucial role in determining the structure and properties of biomolecules, molecular crystals and many other systems. However, an accurate description of dispersion is highly challenging, with the most widely used electronic structure technique, density functional theory (DFT), failing to describe them with standard approximations. Therefore, applications of DFT to systems where dispersion is important have traditionally been of questionable accuracy. However, the last decade has seen a surge of enthusiasm in the DFT community to tackle this problem and in so-doing to extend the applicability of DFT-based methods. Here we discuss, classify, and evaluate some of the promising schemes to emerge in recent years. A brief perspective on the outstanding issues that remain to be resolved and some directions for future research are also provided.

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I. INTRODUCTION

The theoretical description of matter as well as of many chemical, physical, and biological processes requires accurate methods for the description of atomic and molecular-scale interactions. Whilst there are many quantum mechanical approaches, in the past few decades Kohn-Sham density functional theory (DFT)\(^1,2\) has established itself as the theoretical method of choice for this task, undergoing a meteoric rise in large parts of physics, chemistry, and materials science. The rise of DFT and its uptake in academia and industry has been widely discussed, and was perhaps illustrated most clearly in Burke’s recent Spotlight article on DFT.\(^3\)

Although DFT is in principle exact, in practice approximations must be made for how electrons interact with each other. These interactions are approximated with so-called exchange-correlation (XC) functionals and much of the success of DFT stems from the fact that XC functionals with very simple forms often yield accurate results. However there are situations where the approximate form of the XC functional leads to problems. One prominent example is the inability of “standard” XC functionals to describe long-range electron correlations, otherwise known as electron dispersion forces; by standard XC functionals we mean the local density approximation (LDA), generalized gradient approximation (GGA) functionals or the hybrid XC functionals. The “lack” of dispersion forces – often colloquially referred to as van der Waals (vdW) forces – is one of the most significant problems with modern DFT and, as such, the quest for DFT-based methods which accurately account for dispersion is becoming one of the hottest topics in computational chemistry, physics, and materials science. Fig. 1 underlines this point, where it can be seen that over 800 dispersion-based DFT studies were reported 2011 compared to fewer than 80 in the whole of the 1990s.

Dispersion can be viewed as an attractive interaction originating from the response of electrons in one region to instantaneous charge density fluctuations in another. The leading term of such an interaction is instantaneous dipole-induced dipole which gives rise to the well known \(-1/r^6\) decay of the interaction energy with interatomic separation \(r\). Standard XC functionals don’t describe dispersion because: (a) instantaneous density fluctuations...
are not considered; and (b) they are “short-sighted” in that they consider only local properties to calculate the XC energy. The consequence for two noble gas atoms, for example, is that these functionals give binding or repulsion only when there is an overlap of the electron densities of the two atoms. Since the overlap decays exponentially with the interatomic separation, so too does any binding. We show this in Fig. 2 for one of the most widely used GGAs — the so-called Perdew-Burke-Ernzerhof (PBE) functional — for a binding curve between two Kr atoms.

![Figure 2](image-url) **FIG. 2.** Binding curves for the Kr dimer obtained with the PBE exchange-correlation functional and an accurate model potential. Dispersion originates from fluctuations in the electron density which polarize different atoms and molecules, as shown schematically in the lower right diagram. The interaction then exhibits the well-known $-1/r^6$ decay. This $-1/r^6$ decay is not reproduced with PBE (or other semi-local functionals) which instead gives an exponential decay for the interaction because PBE relies on the overlap of density to obtain the interaction.

The binding of noble gases is a textbook dispersion bonded system. However, it has become increasingly apparent that dispersion can contribute significantly to the binding of many other types of materials, such as biomolecules, adsorbates, liquids, and solids. Fig. 3 illustrates a more “real world” example, where an accurate description of dispersion is critical. The figure reports binding energies obtained from PBE and an accurate reference method for DNA base pairs in two different configurations. In the first configuration the binding between the base pairs is dominated by hydrogen bonding. Hydrogen bonding is governed mainly by electrostatics and a standard functional such as PBE predicts reasonable hydrogen bond strengths and as a result the stability of the dimer is quite close (within 15%) to the reference value. However, in the other “stacked” configuration the binding is dominated by dispersion forces and for this isomer PBE is hopeless, hardly predicting any binding between the base pairs at all. This huge variability in performance is far from ideal and since the stacked arrangement of base pairs is a common structural motif in DNA, the result suggests that DNA simulated with PBE would not be stable.

As we will see throughout this article there are many other examples of the importance of dispersion to the bonding of materials and in recent years a plethora of schemes has been proposed to treat dispersion within DFT. Here we will discuss some of the main approaches developed and in the process attempt to provide a useful classification of them. We also highlight some of the obstacles that must be overcome before improved DFT-based methods for including dispersion are available. By its nature, this article is a limited personal view that cannot cover every development in this thriving field. For the most part we have tried to keep the overview simple, and have aimed it primarily at newcomers to the field, although we do go into more depth at the later stages. For more detailed discussions of some of the methods shown here the interested reader should consult the reviews of Grimme, Tkatchenko et al., Johnson et al., or others. Some relevant developments are also discussed in the reviews of Sherrill and Riley et al. which are more focused on post Hartree-Fock (HF) methods.

**II. A CLASSIFICATION OF THE COMMON DFT-BASED DISPERSION METHODS**

Many DFT-based dispersion techniques have been developed and rather than simply listing them all, it is useful to try to classify them. A natural way to do this is to consider the level of approximation each method makes in obtaining the long range dispersion interactions, that is, the interactions between well separated fragments where dispersion is clearly defined. In doing this it turns out that groups of methods which exploit similar approximations emerge. Here, we simply identify these groupings and rank them from the most approximate to the more sophisticated approaches in a manner that loosely...
resembles the well-known “Jacob’s ladder” of functionals introduced by Perdew. Therefore, by analogy, we introduce a “stairway to heaven” for long range dispersion interactions and place each group of dispersion correction schemes on a different step of the stairway. In complete analogy to the ladder, when climbing the stairway progressively higher overall accuracy can be expected until exact results, and thus heaven, are reached. We stress the point “overall accuracy” because, as with the ladder, climbing the stairway does not necessarily mean higher accuracy for every particular problem but rather a smaller probability to fail, i.e., a statistical improvement in performance. A schematic illustration of the stairway is shown in Fig. 4 and in the following sections each step on the stairway is discussed.

FIG. 4. In analogy with the Jacob’s ladder classification of functionals the “stairway to heaven” is used here to classify and group DFT-based dispersion correction schemes. At ground level are methods which don’t describe the long range asymptotics. Simple $C_6$ correction schemes sit on the first step, on the second step are approaches that utilise environment dependent $C_6$ corrections. The long range density functionals sit on step 3 and on step 4 and above are approaches which go beyond pairwise additive determinations of dispersion. Upon climbing each step of the stairway the level of approximation is reduced and the overall accuracy is expected to increase.

A. Ground – Binding with incorrect asymptotics

First, the ground is occupied by methods that simply don’t describe the long range asymptotics. These approaches give incorrect shapes of binding curves and underestimate the binding of well separated molecules. Some of these methods are, nevertheless, being used for weakly bonded systems. Although this might seem odd, this somewhat misguided approach is used because some standard DFT functionals bind dispersion bonded systems at short separations. A prominent example is the LDA which has been used to study systems where dispersion plays a major role such as graphite or noble gases on metals. However, the results with LDA for dispersion bonded systems have limited and inconsistent accuracy and the asymptotic form of the interaction is incorrect. More promising approaches on the ground level of our stairway are density functionals specifically fitted to reproduce weak interactions around minima as well as specially adapted pseudopotentials.

The “Minnesota functionals” are an example of a new breed of functionals that are fitted to a dataset that includes binding energies of dispersion bonded dimers, amongst other properties. Although these functionals can describe binding accurately at separations around minima, they suffer from the same incorrect asymptotics as the LDA does. On the plus side, the reference data used in their construction also contains properties other than weak interactions (e.g., reaction barriers), so that they can be rather accurate for general chemistry problems. This is a clear advantage over “proper” dispersion correction schemes which often utilise GGA functionals that can be less accurate for such problems.

For electronic structure codes that make use of pseudopotentials, dispersion can also be modelled by adding a specially constructed pseudopotential projector. Within this class are the dispersion corrected atom-centered potentials (DCACP) and the local atomic potentials (LAP) methods. These approaches have shown a lot of promise, however, effort is required to carefully fit the potentials for each element and they are not easily transferable to all-electron methods.

B. Step one – Simple $C_6$ corrections

The basic requirement for any DFT-based dispersion scheme should be that it yields reasonable $-1/r^6$ asymptotic behavior for the interaction of particles in the gas phase, where $r$ is the distance between the particles. A simple approach for achieving this is to add an additional energy term which accounts for the missing long range attraction. The total energy then reads

$$E_{\text{tot}} = E_{\text{DFT}} + E_{\text{disp}},$$

where $E_{\text{DFT}}$ is the DFT total energy computed with a given XC functional. The dispersion interaction is given by

$$E_{\text{disp}} = - \sum_{A,B} C_{6}^{A B} / r_6^{A B},$$

where the dispersion coefficients $C_{6}^{A B}$ depend on the elemental pairs A and B. Within this approach dispersion is assumed to be pairwise additive and can therefore be calculated as a sum over all pairs of atoms A and B. Methods on step 1 of the stairway use coefficients that are tabulated, isotropic (i.e., direction independent) and constant, and these methods are generally termed “DFT-
Mostly because of the simplicity and low computational cost this pairwise $C_6/r^6$ correction scheme is widely used. Nonetheless, it has at least four clear shortcomings which limit the accuracy one can achieve with it. First, the $C_6/r^6$ dependence represents only the leading term of the correction and neglects both many-body dispersion effects\textsuperscript{37} and faster decaying terms such as the $C_8/r^8$ or $C_{10}/r^{10}$ interactions. Second, it is not clear where one should obtain the $C_6$ coefficients. Various formulae, often involving experimental input (ionization potentials and static polarizabilities), have been proposed for this.\textsuperscript{7–9,12} However, this reliance on experimental data limited the set of elements that could be treated to those present in typical organic molecules. The third issue is that the $C_6$ coefficients are kept constant during the calculation, and so effects of different chemical states of the atom or the influence of its environment are neglected. The fourth drawback, which we discuss later, is that the $C_6/r^6$ function diverges for small separations (small $r$) and this divergence must be removed.

For a more widely applicable method a more consistent means of deriving the dispersion coefficients is required. In 2006 Grimme published one such scheme, referred to as DFT-D2.\textsuperscript{17} In this approach the dispersion coefficients are calculated from a formula which couples ionization potentials and static polarizabilities of isolated atoms. Data for all elements up to Xe are available and this scheme is probably the most widely used method for accounting for dispersion in DFT at present. Whilst incredibly useful, DFT-D2 is also not free from problems. In particular, for some elements arbitrary choices of dispersion coefficients still had to be made. For example, alkali and alkali earth atoms use coefficients that are averages of the previous noble gas and group III atom. Furthermore, the dispersion energy, $E_{\text{disp}}$, is scaled according to the XC functional used and as a result the interaction energy of two well separated monomers is not constant but sensitive to the choice of XC functional.

With the simple $C_6/r^6$ correction schemes the dispersion correction diverges at short inter-atomic separations and so must be “damped.” The damped dispersion correction is typically given by a formula like

$$E_{\text{disp}} = - \sum_{A,B} f(r_{AB}, A, B) C_6^{AB} / r_{AB}^6,$$  \hspace{1cm} (3)

where the damping function $f(r_{AB}, A, B)$ is equal to one for large $r$ and decreases $E_{\text{disp}}$ to zero or to a constant for small $r$. We illustrate the divergence and a possible damping by the red curves in Fig. 5. How the damping is performed is a thorny issue because the shape of the underlying binding curve is sensitive to the XC functional used and so the damping functions must be adjusted so as to be compatible with each exchange-correlation or exchange functional. This fitting is also sensitive to the definition of atomic size (van der Waals radii are usually used) and must be done carefully since the damping function can actually affect the binding energies even more than the asymptotic $C_6$ coefficients.\textsuperscript{38} The fitting also effectively includes the effects of $C_8/r^8$ or $C_{10}/r^{10}$ and higher contributions, although some methods treat them explicitly.\textsuperscript{19,39} An interesting damping function has been proposed within the so-called DFT coupled cluster approach (DFT/CC), it has a general form that can actually force the dispersion correction to be repulsive.\textsuperscript{40}

**C. Step two – Environment-dependent $C_6$ corrections**

A problem with the simple “DFT-D” schemes is that the dispersion coefficients are predetermined and constant quantities. Therefore the same coefficient will be assigned to an element no matter what its oxidation or hybridization state. However the errors introduced by this approximation can be large, e.g., the carbon $C_6$ coefficients can differ by almost 35% between the $sp$ and $sp^3$ hybridized states.\textsuperscript{9} Thus the emergence of methods where the $C_6$ coefficients vary with the environment of the atom has been a very welcome development. We put

![Energy vs. Distance](image-url)

**FIG. 5.** Schematic illustration of a binding curve ($E_{\text{tot}}$) obtained from a dispersion corrected DFT calculation and the contributions to it from the regular DFT energy ($E_{\text{DFT}}$) and the dispersion correction ($E_{\text{disp}}$). The function $–1/r^6$ used to model the dispersion correction diverges for small $r$ (red dashed curve) and must be damped (solid red curve). Since the details of the damping strongly affect the position of the energy minima on the binding curve, the damping function needs to be fit to reference data. In addition, the damping function will be different for different XC functionals used to obtain $E_{\text{DFT}}$. Before leaving the simple $C_6/r^6$ schemes we note that although the approaches developed by Grimme are widely used, other parameterizations, which differ in, e.g., the combination rules used, are available.\textsuperscript{40–43} Furthermore, functionals such as B97-D\textsuperscript{17} and ωB97X-D\textsuperscript{44} have been specifically designed to be compatible with this level of dispersion correction.
these methods on the second step of our stairway. They still use Equation 3 to obtain the dispersion correction and, as with the step 1 methods, some reference data (such as atomic polarizabilities) is used to obtain the $C_6$ coefficients.

We now discuss three step 2 methods: DFT-D3 of Grimme, the approach of Tkatchenko and Scheffler (vdW(TS)), and the Becke-Johnson model (BJ). The unifying concept of these methods is that the dispersion coefficient of an atom in a molecule depends on the effective volume of the atom. When the atom is “squeezed”, its electron cloud becomes less polarizable leading to a decrease of the $C_6$ coefficients.

Grimme et al.\textsuperscript{19} capture the environmental dependence of the $C_6$ coefficients by considering the number of neighbors each atom has. When an atom has more neighbors it is thought of as getting squeezed and as a result the $C_6$ coefficient decreases. This effect is accounted for by having a range of precalculated $C_6$ coefficients for various pairs of elements in different reference (hybridization) states. In the calculation of the full system the appropriate $C_6$ coefficient is assigned to each pair of atoms according to the current number of neighbors. The function calculating the number of neighbors is defined in such a way that it continuously interpolates between the precalculated reference values. Therefore if the hybridization state of an atom changes during a simulation, the $C_6$ coefficient can also change continuously. Despite the simplicity of this approach, termed “DFT-D3”, the dispersion coefficients it produces are pretty accurate. Specifically, based on the reference data of Meath and coworkers there is a mean absolute percentage deviation (MAPD) in the $C_6$ coefficients of 8.4%. Moreover, the additional computational cost is negligible since the number of neighbors can be obtained quickly.

In 2009 Tkatchenko and Scheffler\textsuperscript{18} proposed a method which relies on reference atomic polarizabilities and reference atomic $C_6$ coefficients\textsuperscript{45} to calculate the dispersion energy. These quantities are sufficient to obtain the $C_6$ coefficient for a pair of unlike atoms.\textsuperscript{46} To obtain environment dependent dispersion coefficients effective atomic volumes are used. During the calculation on the system of interest the electron density of a molecule is divided between the individual atoms (the Hirshfield partitioning scheme is used) and for each atom its corresponding density is compared to the density of a free atom. This factor is then used to scale the $C_6$ coefficient of a reference atom which changes the value of the dispersion energy. The accuracy of the final (isotropic, averaged) $C_6$ coefficients is quite high, with the MAPD on the data of Meath and coworkers being only 5.4%. However, it is not yet clear if the scaling of $C_6$ coefficients with volume will be accurate for more challenging cases such as different oxidation states in e.g. ionic materials.

The most complex step 2 method is the BJ model,\textsuperscript{13–16,39} which exploits the fact that around an electron at $r_1$ there will be a region of electron density depletion, the so-called XC hole. Even for symmetric atoms, this creates asymmetric electron density and thus non-zero dipole and higher-order electrostatic moments, which causes polarization in other atoms to an extent given by their polarizability. The result of this is an attractive interaction with the leading term being dipole-induced dipole. The biggest question in the BJ model is how to quantify the XC hole and in the method it is approximated as the exchange-only hole (calculated using the Kohn-Sham orbitals). An average over the positions $r_1$ of the reference electron then gives the average square of the hole dipole moment, denoted $\langle d^2\rangle$ which together with the polarizabilities $\alpha$ enters the formula for the dispersion coefficient

$$C_6^{AB} = \frac{\alpha_A \alpha_B \langle d^2 \rangle_A \langle d^2 \rangle_B}{\langle d^2 \rangle_A \alpha_B + \langle d^2 \rangle_B \alpha_A}. \quad (4)$$

This is, in fact, very similar to the vdW(TS) formula but in vdW(TS) precalculated $C_6$ coefficients are used instead of the hole dipole moments. The BJ model is very intriguing and several authors have studied how it relates to formulae derived from perturbation theory.\textsuperscript{47–49}

In the BJ model the $C_6$ coefficients are altered through two effects. First, the polarizabilities of atoms in molecules are scaled from their reference atom values according to their effective atomic volumes. Second, the dipole moments respond to the chemical environment through changes of the exchange hole, although this effect seems to be difficult to quantify precisely. The details of how to obtain atomic volumes and the exchange hole are known to affect the results obtained to some extent,\textsuperscript{50–52} but overall the accuracy of the asymptotic $C_6$ coefficients obtained from the BJ model is quite satisfactory, with a MAPD of 12.2% for the data of Meath and coworkers.\textsuperscript{19} Compared to the two previous step 2 approaches, however, the computational cost is relatively high, in the same ballpark as the cost of a hybrid functional.\textsuperscript{26}

### D. Step three – Long-range density functionals

The methods discussed so far require predetermined input parameters to calculate the dispersion interaction, either the $C_6$ coefficients directly or the atomic polarizabilities. Now we discuss approaches that do not rely on external input parameters but rather obtain the dispersion interaction directly from the electron density. This, in principle, is a more general strategy and thus we place these methods on step 3 of our stairway. The methods have been termed non-local correlation functionals since they add non-local (i.e. long range) correlations to local or semi-local correlation functionals.

The non-local correlation energy $E_{c}^{nl}$ is calculated from

$$E_{c}^{nl} = \iint dr_1 dr_2 n(r_1) \phi(r_1, r_2) n(r_2). \quad (5)$$

This is a double space integral where $n(r)$ is the electron density.
density and \( \phi(r_1, r_2) \) is some integration kernel. The kernel is analogous to the classical Coulomb interaction kernel \( 1/|r_1 - r_2| \) but with a more complicated formula used for \( \phi(r_1, r_2) \) with \( O(1/|r_1 - r_2|^6) \) asymptotic behavior. The formula has a pairwise form and thus ignores the medium between points \( r_1 \) and \( r_2 \). Various forms for \( E_{\text{c}}^{\text{nl}} \) were proposed in the nineties\(^4\)\(^6\) but were restricted to non-overlapping fragments. This rather severe limitation was removed by Dion et al. in 2004\(^1\)\(^0\) who proposed a functional form which can be evaluated for overlapping molecules and for arbitrary geometries. Within this approach the XC energy \( E_{\text{xc}} \) is calculated as

\[
E_{\text{xc}} = E_{\text{x}}^{\text{GGA}} + E_{\text{c}}^{\text{LDA}} + E_{\text{c}}^{\text{nl}},
\]

where the terms on the right hand side are the exchange energy in the revPBE approximation,\(^5\)\(^3\) the LDA correlation energy, and the non-local correlation energy term, respectively. This method has been termed the van der Waals density functional (vdW-DF). vdW-DF is a very important conceptual development since it adds a description of dispersion directly within a DFT functional and combines correlations of all ranges in a single formula.

Since the development of the original vdW-DF a number of follow up studies have aimed at understanding and improving the performance of the method. First, it turns out that vdW-DF tends to overestimate the long range dispersion interactions: Vydrov and van Voorhis have shown that when the \( C_6 \) coefficients are evaluated the errors can be as large as \( \sim 30\% \).\(^5\)\(^4\) To address this these authors proposed (computationally cheaper) functionals that reduce the average errors by approximately \( 50\% \).\(^5\)\(^4\)\(^5\)\(^7\)\(^8\) The developers of the original vdW-DF tried to address its tendency to overbind at large separations by proposing vdW-DF\(^2\)\(^5\)\(^8\)\(^9\) This functional, which involves changes to both the exchange and non-local correlation components tends to improve the description of the binding around energy minima, however, the \( C_6 \) coefficients it predicts are considerably underestimated.\(^5\)\(^6\) Second, aside from the particular form of \( E_{\text{c}}^{\text{nl}} \), the choice of the exchange functional used in Equation 6 has received considerable attention. The original revPBE exchange functional chosen for vdW-DF often leads to too large intermolecular binding distances and inaccurate binding energies. To remedy these problems alternative “less repulsive” exchange functionals have been proposed\(^1\)\(^0\)\(^6\)\(^3\)\(^6\) which when incorporated within the vdW-DF scheme (Equation 6) lead to much improved accuracy. Of these the “optB88” and “optPBE” exchange functionals have been shown to offer very good performance for a wide range of systems.\(^6\)\(^0\)\(^6\)\(^1\)\(^4\)\(^6\)\(^5\)

Initially the non-local correlation functionals came, to a lesser or greater extent,\(^6\)\(^6\)\(^7\) with a higher computational cost than GGAs or hybrid functionals. However, thanks to the work of Román-Pérez and Soler the computational cost of vdW-DF is now comparable to that of a GGA.\(^6\)\(^8\) In addition, self-consistent versions of vdW-

DF and several of its offspring are now implemented in widely distributed DFT codes such as Siesta,\(^6\)\(^8\) VASP,\(^6\)\(^1\) QuantumESPRESSO,\(^6\)\(^9\) and QChem.\(^6\)\(^6\) Other ways to reduce the cost of vdW-DF calculations have also been reported, for example, Silvestrelli has shown\(^7\)\(^0\)\(^7\)\(^1\) that utilizing Wannier functions to represent the electron density allows for an analytic evaluation of the functionals proposed in the nineties.\(^5\)\(^6\) Particularly interesting is the local response dispersion (LRD) approach of Sato and Nakai\(^7\)\(^2\)\(^7\)\(^3\) which yields very accurate \( C_6 \) coefficients. Overall, by not relying on external reference data, the step 3 approaches are less prone to fail for systems outside of the reference or fitting space of step 2 methods. However, very precise calculations of the dispersion energy are difficult and the formulae underlying vdW-DF and similar approaches can be somewhat complicated.

### E. Higher steps – Beyond pairwise additivity

The main characteristic of the methods discussed so far is that they consider dispersion to be pairwise additive. As a consequence the interaction energy of two atoms or molecules remains constant no matter what material separates them and all atoms interact “on their own” with no consideration made for collective excitations.\(^7\)\(^4\) While such effects don’t seem to be crucially important in the gas phase, especially for small molecules, they are important for adsorption or condensed matter systems where the bare interaction is screened.\(^7\)\(^5\) We now briefly discuss some of the methods being developed which treat dispersion beyond the pairwise approximation. The range of approaches is quite wide, from methods based on atom centered interactions, to methods involving density, to methods using electron orbitals. Because of the freshness of most of the methods we discuss them together.

#### FIG. 6. The long-range electron correlations of two isolated atoms can be described by an effective \( -1/r^6 \) formula. In a condensed system the interaction is modified (screened) by the presence of the other electrons.

Recently, the Axilrod-Teller-Muto\(^7\)\(^6\)\(^7\)\(^7\) formula has been used to extend the atom-centred pairwise approaches to include three-body interactions.\(^1\)\(^9\)\(^7\)\(^8\) The triple-dipole interaction between three atoms A, B, and
where \( \alpha, \beta, \) and \( \gamma \) are the internal angles of the triangle formed by the atoms A, B, and C. The dispersion coefficient \( C_{ABC} \) is again obtained from reference data. Notably, von Lilienfeld and Tkatchenko estimated the magnitude of the three body terms to be \( \sim -25\% \) (destabilizing) of the two body term for two graphene layers. Since the three-body interaction is repulsive for atoms forming an acute angled triangle, the interaction was found to be destabilising for stacked configurations of molecular clusters such as benzene dimers.

The atom-centered approach can be used to approximate the many-body dispersion interaction using a model of coupled dipoles. Here quantum harmonic oscillators with characteristic frequencies occupy each atomic position and the dispersion interaction is obtained from the shifts of the frequencies of the oscillators upon switching on their interaction. The model has been applied outside DFT for some time and has recently been used to show the non-additivity of dispersion in anisotropic materials. The initial applications of this approach termed many-body dispersion (MBD) are quite promising and suggest that the higher-order dispersion terms can be important even for systems such as solid benzene. However, getting accurate relations between atoms and oscillator models seems to be rather challenging, especially when describing both localized and delocalized electrons.

In the context of DFT, orbitals can be used to calculate the correlation energy using the adiabatic-connection fluctuation-dissipation theorem (ACFDT). The particular approach which has received the most attention recently is the so-called random phase approximation (RPA). Results from RPA have been very encouraging and it exhibits, for example, a consistent accuracy for solids and the correct asymptotic description for the expansion of graphite, a feature which the pairwise methods fail to capture. RPA is analogous to post-HF methods and indeed direct links have been established. Unfortunately, it also shares with the post-HF methods a high computational cost (the cost increases approximately with the fourth power of the system size) and slow convergence with respect to basis set size. Approaches going beyond RPA are also receiving attention, for example, the second-order screened exchange or single excitations corrections.

Another method involving orbitals to obtain the correlation energy is to combine DFT with post-HF methods in the hope of exploiting the benefits of each approach. While the post-HF methods can describe long range interactions accurately (albeit expensively) using orbitals, DFT is efficient for the effective description of the short range part of the interactions. So called range separated methods are an example of this approach where long range correlations are treated by, e.g., second-order perturbation theory, the coupled cluster method, or RPA. Range separated functionals can be very accurate, can account for many-body interactions, and can deliver relatively fast convergence of the correlation energy with respect to basis set size. Another example from this class are the so called double hybrid functionals, which include Fock exchange and a second order perturbation theory correlation contribution. Generally the coefficients for these contributions are fitted to reproduce reference data. Since the original double hybrid functionals were more concerned with reaction energies and barrier heights than dispersion, dispersion interactions were actually underestimated. Newer functionals, e.g., mPW2PLYP-D, add a dispersion correction in the sense of step 1 or 2 methods, however, they are computationally very expensive and at present prohibitively expensive for most condensed phase and surface studies.

### F. Summary

The higher steps of the stairway with the promising schemes that go beyond the pairwise approximation close our overview and classification of methods for treating dispersion within DFT. In Table I we summarize some of the key aspects of the most relevant schemes, such as what properties they use to obtain the dispersion contribution (via, e.g., the \( C_{6} \)) and the approximate relative computational cost. The computational cost of a method is, of course, a key factor since no matter how accurate it is, it will not be used if the computational cost is prohibitive. In this regard the cost of the simple pairwise corrections on step 1 is essentially zero compared to the cost of the underlying DFT calculation and these methods can be recommended as a good starting point for accounting for dispersion. Because of the correct asymptotic behavior (at least for gas phase molecules) they are preferable to methods from ground. Compared to the step 1 methods the DFT-D3 and vdW(TS) schemes don’t add a significant computational cost, whereas the vdW-DF approach increases the computational time by about 50% compared to a GGA calculation. Since the accuracy of these approaches tends to be higher than that of the step 1 methods, they should be preferred over the step 1 corrections. However, when applied to a particular problem, the accuracy of any method used should be tested or verified against experimental or other reference data since any approach can, in principle, fail for a specific system. The methods on higher steps are mostly in development and currently the ACFDT or range separated functionals require a computational cost two to four orders of magnitude higher than a GGA calculation which limits their applicability.
TABLE I. Summary of some of the most widely used methods for capturing dispersion interactions in DFT, ordered according to the “Step” they sit on in the stairway to heaven (Fig. 4). Information on the reference used for the C₆ coefficients, what the C₆ depend on, and the additional computational cost of each approach compared to a regular GGA calculation is reported.

| Method     | Step | Reference for C₆ | C₆ depend on          | Additional computational cost a | Ref. |
|------------|------|------------------|-----------------------|---------------------------------|------|
| Minnesota  | 0    | None             | N/A                   | None                            | e.g. 34 |
| DCACP      | 0    | None             | N/A                   | Small                           | 11   |
| DFT-D      | 1    | Various          | Constant              | Small                           | e.g. 110 |
| DFT-D3     | 2    | TDDFT            | Structure             | Small                           | 19   |
| vdW(TS)    | 2    | Polarizabilities | Atomic volume         | Small                           | 18   |
|            |      | and atomic C₆    |                       |                                 |      |
| BJ         | 2    | Polarizabilities | Atomic volume, X hole | Large                           | 14   |
| LRD        | 3    | C₆ calculated    | Density               | Small                           | 72   |
| vdW-DF     | 3    | C₆ calculated    | Density               | ≈50%                            | 10   |
| Dbl. hybrids | 4   | None or as “-D”  | Orbitals              | Large                           | 110  |

a The BJ model and the double hybrids (labelled “Dbl. hybrids”) are more computationally expensive than the simpler “DFT-D” methods and the calculation of the correlation energy in vdW-DF leads to a ≈50% slow-down compared to a GGA calculation when done efficiently.
III. GENERAL PERFORMANCE

We have commented in passing on how some of the methods perform, mainly with regard to the dispersion coefficients. Now we discuss accuracy in more detail by focussing on binding energies for a few representative gas phase clusters, a molecular crystal, and an adsorption problem. Before doing so, it is important to emphasise two points. First, establishing the accuracy of a method is not as straightforward as it might seem since it must be tested against accurate reference data (e.g. binding energies). Experimental data can be inaccurate and is rarely directly comparable to theory since quantum nuclear and/or thermal effects, which affect the experimental values, are often neglected in simulation studies. Theoretical reference data, on the other hand, is more appropriate but rather hard to come by since accurate reference methods such as post-HF methods (MP2, CC) or quantum Monte Carlo (QMC) are so computationally expensive that they can generally be applied to only very small systems (tens of atoms).

Second, the results obtained with many dispersion based DFT approaches are often strongly affected by the fitting procedure used when combining the long range dispersion interaction with the underlying exchange or XC functionals.

A. Gas phase clusters

Because of growing computer power and algorithmic improvements, systems of biological importance such as DNA and peptides can now be treated with DFT. As discussed in the introduction, dispersion plays an important role in stabilising the folded state and thus dispersion corrected functionals are required. However, since accurate reference data can only be obtained for systems with tens of atoms, it’s just not possible to obtain reference data for DNA or a protein directly and fragments of the large molecules are used to build test sets of binding energies and geometries instead. One such test set is the popular S22 data set of Jurečka et al. It is useful since it contains 22 different dimers with a range of weak bonding types and with a wide range of interaction energies. Results for some of the methods are reported in Table II; specifically we report mean absolute deviations (MAD) and mean absolute percentage deviations (MAPD). Included in Table II are results from second order Møller-Plesset perturbation theory (MP2) – a widely used post-HF method – as are results from the LDA and PBE functionals as examples of methods that don’t treat dispersion explicitly.

| Method          | MAD  | MAPD | Ref. |
|-----------------|------|------|------|
| Ground          |      |      |      |
| LDA             | 8.66 | 30.5 | 116  |
| PBE             | 11.19| 57.6 | 116  |
| M06-2X          | 2.52 | 11.3 | 117  |
| LAP             | 2.51 | 7.8  | 35   |
| Step 1          |      |      |      |
| B97-D2          | 1.51 | 7.3  | 44   |
| ωB97X-D         | 0.76 | 5.6  | 44   |
| Step 2          |      |      |      |
| BLYP-D3         | 0.96*| –    | 19   |
| PBE-vdW(TS)     | 1.25**| 9.2**| 18   |
| rPW86-BJ        | 1.50 | 6.1  | 51   |
| Step 3          |      |      |      |
| LC-BOP+LRD      | 0.86 | 4.6  | 73   |
| vdW-DF          | 6.10 | 22.0 | 67   |
| optB88-vdW      | 1.18 | 5.7  | 60   |
| vdW-DF2         | 3.94 | 14.7 | 118  |
| VV10            | 1.35 | 4.5  | 118  |
| Higher steps    |      |      |      |
| B2PLYP-D3       | 1.21*| –    | 19   |
| ωB97X-2         | 1.17 | 8.8  | 119  |
| PBE-MB          | –    | 5.4**| 80   |
| RPA             | 3.29**| –    | 101  |
| post-HF         |      |      |      |
| MP2             | 3.58 | 19.4 | 115  |

* using the original reference values of Jurečka et al. (Ref. 22)
** using the values of Takatani et al. (Ref. 120) which are very similar to those of Podeszwa et al. (Ref. 115)

Small water clusters, in particular water hexamers, are an interesting system where dispersion plays a significant role (see, e.g. Refs. 121–125). The four relevant isomers (known as “book”, “cage”, “cyclic”, “prism”, Fig. 7) all have total energies that differ by \(< 1.3 \text{ kJ/mol per molecule}\) according to accurate post-HF methods. While the “prism” and “cage” isomers are preferred by post-HF methods, standard XC steps of the stairway can surpass those from methods on higher steps. This is not that surprising since some of the methods reported in Table II were actually developed by fitting to the S22 data set itself.
functionals find the more open “cyclic” and “book” isomers to be more stable. Recent work reveals that dispersion impacts profoundly on the relative energies of the isomers and improved relative energies are obtained when dispersion is accounted for. Indeed the water hexamers have become an important test system with techniques such as DFT-D, \textsuperscript{121} vDW(TS),\textsuperscript{121} vDW-DF,\textsuperscript{126} optB88-vdW,\textsuperscript{60} and the modified vDW-DF approach of Silvestrelli (BLYP-vdW(Silv.))\textsuperscript{127} all having been applied. The performance of these schemes in predicting the relative energies of the water hexamers is shown in Fig. 7, where one can also see the improvement over a standard functional such as PBE. One general point to note, however, is that even without a dispersion correction certain functionals can already give quite accurate absolute binding energies for systems held together mainly by hydrogen bonding and adding dispersion corrections can actually lead to too large absolute binding energies. This is indeed the case for the water hexamers where, for example, the vdw(TS) correction to PBE gives binding energies for the hexamers that are too large by \(\approx 4\) kJ/mol per molecule.

**B. Molecular crystals – solid benzene**

Molecular crystal polymorph prediction is another important area where dispersion forces can play a critical role, and even for molecular crystals comprised of small molecules several polymorphs often exist within a small energy window.\textsuperscript{128,129} Identifying the correct energetic ordering of the polymorphs can therefore be a stringent test for any method. A large number of molecular crystals have been characterised experimentally (see, e.g. Ref. 130) and this, therefore, could serve as a rich testing ground for DFT-based dispersion schemes.

Solid benzene is one of the most widely examined test systems, with studies focussing on the experimental density and the cohesive energy of the crystal.\textsuperscript{131} The latter is obtained from the experimental sublimation energy from which the effects of temperature and quantum nuclear effects must be subtracted giving a value in the 50 to 54 kJ/mol per molecule range.\textsuperscript{132} Calculations with PBE give an abysmal cohesive energy of only 10 kJ/mol per molecule and a volume \(\approx 30\%\) too large compared to experiment.\textsuperscript{133} This huge difference between theory and experiment suggests that dispersion is important to the binding of the crystal and indeed the error is greatly reduced when even rather simple schemes are used. For example, PBE-D2 and the DCACP potentials give estimates of the lattice energy (55.7 kJ/mol\textsuperscript{133} and 50.6 kJ/mol,\textsuperscript{36} respectively) and densities that are within 10\% of experiment. While the PBE-vdW(TS) scheme overbinds slightly (66.6 kJ/mol), it has been suggested that the many-body interactions are quite important for this system and their inclusion reduces the PBE-vdW(TS) value by 12 kJ/mol.\textsuperscript{80} Non-local vdw-DF overestimates the cell volume by \(\approx 10\%\) but gives a rather good cohesive energy of 58.3 kJ/mol.\textsuperscript{116} VDW-DF2 reduces both the binding energy (to 55.3 kJ/mol) and the error in the cell volume, which turns out to be \(\approx 3\%\) larger than the experimental value.\textsuperscript{116} Using functionals proposed to improve upon the overly repulsive behavior of vdw-DF, such as optB88-vdW leads to an improvement of the reference volume (\(\approx 3\%\) smaller than experiment), but the cohesive energy is overestimated, being 69.4 kJ/mol.\textsuperscript{116} Again, the overestimated cohesive energy may result from missing many-body interactions. The RPA has also been applied to this system and while the density is in very good agreement with experiment, the cohesive energy is slightly underestimated, at 47 kJ/mol.\textsuperscript{132} Overall, the improvement over a semi-local functional such as PBE is clear but more applications and tests on a wider range of systems are needed to help establish the accuracy of the methods.

**C. Adsorption – benzene on Cu(111)**

Adsorption on solid surfaces is another area where great strides forward have recently been made with regard to the role of dispersion. Dispersion is of obvious importance to the binding of noble gases to surfaces but it can also be important to chemisorption\textsuperscript{134} and, e.g., water adsorption.\textsuperscript{64} Indeed for weakly chemisorbed systems dispersion forces attain an increased relative importance and one can estimate from various studies (e.g., Refs. 135 and 136) that dispersion contributes about 4 to 7 kJ/mol to the adsorption energy of a carbon-sized atom; not a negligible contribution.

Of the many interesting classes of adsorption system, organic molecules on metals have become a hot area of re-
search and for such systems dispersion must be included if reasonable adsorption energies and structures are to be obtained. Benzene on Cu(111) is an archetypal widely studied system for organic adsorbates on metals. It is also an interesting system because it illustrates how difficult it is sometimes to use experimental reference data, which for this system has proved to be somewhat of a “moving target”. Indirect estimates obtained from temperature programmed desorption initially placed the adsorption energy at 57 kJ/mol\(^{137}\) however, a recent reinterpretation of the experiment moved the adsorption energy up to the 66 to 78 kJ/mol range.\(^{138}\) When looking at this system with DFT, not unexpectedly, PBE gives very little binding (5 kJ/mol).\(^{139}\) Most of the step 1 and 2 methods overestimate the binding,\(^{140}\) for example, PBE-D2 and PBE-vdW(TS) give 97 and 101 kJ/mol, respectively. The exception is the work of Tonigold and Groß\(^{135}\) where a value of 59 kJ/mol was obtained, based on dispersion coefficients obtained by fitting to post-HF data of small clusters. Recently, Ruiz et al.\(^{138}\) approximately included the many-body effects in calculating the \(C_6\) coefficients in the vdW(TS) scheme, which reduced the predicted adsorption energy to 88 kJ/mol. vdW-DF gives adsorption energies of about \(~53\) kJ/mol which underestimates both the old and new reference data.\(^{139,141}\) It is clear from this and other systems that adsorption is very challenging for dispersion-based DFT methods at present. The “DFT-D” methods face the problem of obtaining the \(C_6\) coefficients for the atoms within the surface of the solid and the pairwise methods neglect many-body effects. Both of these issues will require much more consideration in the future.

FIG. 8. In molecular crystals the molecules can be packed in a range of orientations and at a range of distances from each other. This makes molecular crystals a more challenging class of system than gas phase clusters, and, indeed, even for the relatively simple case of benzene shown here, few methods are able to very accurately describe the cohesive properties of the crystal. The benzene molecules in the central unit cell are shown with large balls, all other benzenes in the neighboring unit cells are shown in wireframe.

IV. FINAL REMARKS

An enormous amount of progress has been made over the last few years with the treatment of dispersion forces within DFT. It is now one of the most exciting and thriving areas of development in modern computational materials science and an array of methods has been developed. Here we have introduced and classified some of the main, often complementary, approaches. We have also discussed how some schemes perform on a selection of systems, including gas phase clusters, a molecular solid and an adsorption problem. These examples, and the many others in the literature, demonstrate that the range of systems which can now be treated with confidence with DFT has been greatly extended. Connected with this, the variety of systems and materials to which dispersion forces are now thought to be relevant has grown substantially. As a result the mantra that “dispersion forces are not important” is heard less often now and there is much less of a tendency to sweep dispersion forces under the rug.

We have seen how many methods have been applied and tested on gas phase molecular clusters, for which there are now several approaches that can yield very high accuracy. However, a key contemporary challenge is the need to develop methods that will be accurate for both gas phase molecular systems and problems involving condensed matter such as adsorption. Here, much work remains to be done with regard to the development of DFT-based dispersion techniques for condensed matter as well as in simply better understanding how current techniques perform in e.g. adsorption systems where there may be strong polarization effects. In this regard, approaches based on the ACFDT such as RPA look promising. However, given their high computational cost and complex set-up it is likely that for the foreseeable future such methods will mainly be useful for tour de force reference style calculations rather than for routine studies.\(^{142}\)

Looking to the future, the efficient description of many-body correlation effects in metals and other solids is an important unresolved issue. Even simply better understanding their importance for different systems would be useful. For example, although many body correlation effects should be important in solids, the vdW-DF method which neglects them can perform surprisingly well for solids.\(^{61}\) Another problem closely related to dispersion is the issue of screening which differs substantially in solids and molecules. For molecules, approaches that neglect screening of exchange, e.g., the so-called long-range corrected (LC) exchange functionals\(^{143}\) are beneficial. Indeed, the LC functionals improve many properties of molecules such as electrostatic moments which in turn decreases an important source of errors in vdW bonded systems.\(^{55,119,144}\) In solids, especially metals or semiconductors, the interaction of electrons is significantly modified by the presence of the other electrons and it is essential to capture this XC effect. Indeed, no screen-
Anisotropy can, however, become an issue for highly anisotropic and polarizable objects. Isotropic dispersion coefficients seem to be a good first approximation since the anisotropy for molecules is on the order of 10%. Since screening is system dependent it is unclear how to treat solids and molecules on the same footing in a computationally efficient manner. A further issue, which is often of minor importance but should be accounted for when high accuracy is being sought is anisotropy of the dispersion coefficients. Isotropic dispersion coefficients seem to be a good first approximation since the anisotropy for molecules is on the order of 10%. Anisotropy can, however, become an issue for highly anisotropic and polarizable objects and to establish the overall importance would be useful. The anisotropy can be included in some schemes, as has been done, for example, in the BJ model and the vdw(TS) approach. Many factors have prompted the recent progress with DFT, with one key factor being the parallel development of post-HF methods. This has provided the accurate reference data which has served to both shine light on problems with existing XC functionals and against which new methods can be proved. The fact that some of this reference data has been easily accessible – such as the S22 data set – has also helped. However, as stressed above, an important challenge nowadays is to develop methods that are accurate for solids and for adsorption. Unfortunately for these systems accurate reference data are scarce and, indeed, urgently needed either from experiment (e.g. microcalorimetry for adsorption) or higher level electronic structure theories (e.g. post-HF methods, approximations of ACFDFT, QMC). It is encouraging that progress is being made in both of these areas. It is also very encouraging that condensed phase reference systems are beginning to emerge, such as ice, LiH, water on LiH and water on graphene. By tackling these and other reference systems with the widest possible range of techniques we will better understand the limitations of existing dispersion-based DFT approaches, which will aid the development of more efficient and more accurate methods for the simulation of materials in general.

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