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Density functional for van der Waals forces accounts for hydrogen bond in benchmark set of water hexamers

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A recent extensive study1 investigates how various exchange-correlation (XC) functionals treat hydrogen bonds (HBs) in water hexamers, a critical HB benchmark set. Accurate wave-function studies including perturbative (MP2),1,2 quantum-Monte Carlo (DQMC),3 and coupled-cluster [CCSD(T)] (Ref. 2) calculations agree on the relative energetics of the low-lying isomers:4 The prism has lowest energy followed by (in order) cage, book, and cyclic isomers (Fig. 1). The study1 shows that traditional generalized gradient approximation (GGA) and hybrid functionals used in density-functional theory (DFT) give the wrong dissociation-energy trend of the four isomers (Fig. 1) and that van der Waals (vdW) dispersion forces give key contributions to the dissociation energy and asks whether functionals that incorporate vdW forces implicitly into the XC functional predict the correct trend and yield accurate total dissociation-energy values.1 Here an affirmative answer is given for the vdW-DF functional.3

Typically, a HB is an intermolecular bond between an intramolecular bond of an H atom on a donor molecule and an acceptor molecule. The directionality is the angle between these bonds. Studies on various HB-ed dimers suggest that vdW forces can have a significant effect on the strength of the HB, including an indicated directionality effect.4 The vdW-DF describes vdW effects approximately3,9 but differs significantly from another method doing so, DFT-D,10 being derived from first principles and thus nonempirical and informing about the electron structure.

All calculations are made with the GPAW software.11 The vdW-DF (Ref. 3) implementation is the same real-space method as in Ref. 14, which adds vdW forces perturbatively3 to a self-consistent revPBE15 calculation. The latter uses a well converged grid spacing of 0.13 Å and Eq. (2) is evaluated on a grid twice as dense (a grid spacing of 0.15 Å results in binding-energy differences smaller than 1%). The hexamer is in a cubic cell with side 18 Å and 35 electronic bands are used. The dissociation energy is calculated as in Ref. 1 but with the vdW-DF.

Figure 1 shows trends in dissociation-energy values of the water-hexamer isomers relative to the prism configuration. Even within the narrow energy scale, the trends in the results from wave-function methods [MP2 (Ref. 1), DQMC (Ref. 1), and CCSD(T) (Ref. 2)] are very distinct and close to each other and to earlier CCSD(T) results.16 These trends cannot be accounted for by DFT with traditional GGA [PBE (Ref. 17), RPBE (Ref. 18), and revPBE (Ref. 15)] and hybrid [PBE0 (Ref. 13)] XC functionals.1 Clearly, the vdW-DF and DFT-D methods give both the right stable configuration and the proper energetic order of the lowest isomers.

To follow a trend is not a sufficient criterion for a good XC functional. For instance, LDA gives trend but is ignored due to lack of basic nonlocal interactions and its known overestimate of dissociation of water clusters by more than 50%.1,19 The vdW-DF dissociation-energy values of the low-lying isomers show an underbinding of about 50 meV/H2O (Table I). At typical vdW separations (about 3–5 Å) the accuracy of vdW-DF might be about 0.01 eV,14 but it is expected to be worse at the short separations of HBs (O–H separations being 1.9 Å in the water dimer and, e.g., 1.7–2.0 Å in the water hexamer cage,1 with vdW-DF and MP2 numbers given in Table I). The revPBE exchange functional15 used in vdW-DF lies close to exact exchange [Hartree–Fock (HF)] results for, e.g., argon, krypton,3 and benzene dimers but gets repulsive at shorter separations, depending nonlocally on the density $n(\vec{r})$ with a general kernel $\phi$ derived from first principles.9 Already used successfully for a wide spectrum of systems,5 its virtues, e.g., biological molecules, are highlighted by its account of stacking and HB interactions between nucleobases7 and DNA intercalation.8

The vdW-DF dissociation energy and asks whether functionals that incorporate vdW forces implicitly into the XC functional predict the correct trend and yield accurate total dissociation-energy values. Here an affirmative answer is given for the vdW-DF functional.3

For calculation of electronic structure and total energy, DFT offers an alternative to wave-function methods that is particularly powerful for extended systems, a well-proven excellent tool for strongly bound systems. For also abundant sparse systems that are bound by weak nonbonding forces, including HBs, nonlocal correlation functionals have been proposed recently. One such, the vdW-DF (Ref. 3) includes vdW forces in a seamless fashion with a derived XC functional,

$$E_{\text{vdW-DF}} = E_{\text{X}}^{\text{revPBE}} + E_{\text{C}}^{\text{LDA}} + E_{\text{C}}^{\text{nl}},$$

(1)

where

$$E_{\text{C}}^{\text{nl}} = \frac{1}{2} \int d\vec{r} d\vec{r}' \phi(\vec{r}, \vec{r}') n(\vec{r})$$

(2)
where other flavors of GGA might be closer as for the water dimer (Fig. 2). The relatively small underbinding by use of other less repulsive exchange functionals such as PBE, HF, and a hopefully soon developed systematically improved exchange functional. The latter should be accurate in regions where the HB applies and where vDW and covalent bonds change functional. The Center for Atomic-Scale Materials Design is funded by the Lundbeck Foundation.

We thank Angelos Michaelides, Biswajit Santra, and Matthias Scheffler for providing the hexamer structures calculated in Ref. 1. The Center for Atomic-Scale Materials Design is funded by the Lundbeck Foundation.

![FIG. 1. Dissociation energy per H2O molecule of the water hexamer in low-lying isomers shown below calculated with wave-function [MP2 (Ref. 1), DQMC (Ref. 1), and CCSD(T) (Ref. 2)] and DFT methods [with XC functionals in the PBE, RPBE, and revPBE flavors of GGA and in PBE0 (Ref. 1), DFT-D (Ref. 1), and vDW-DF] relative to the prism isomer.](image)

![FIG. 2. Exchange part of the dissociation energy for the water dimer at varying separations (defined by inset) with exact exchange (HF) calculated by the EXX(X) method (Ref. 12) and with DFT using PBE, RPBE, revPBE, and LDA XC functionals, each calculated self-consistently and then subtracting the correlation energy. LDA exchange is shown to illustrate its tendency to overbind. The nonself-consistent (PBE densities) exchange part of PBE0 is also included by mixing of PBE and EXX exchange as in Ref. 13.](image)

TABLE I. Calculated dissociation-energy values in meV/H2O and shortest separations between hydrogen and oxygen atoms (R–O–H) of the low-lying water-hexamer isomers. Absolute-energy values $E_{\text{diss}}$ calculated with the vDW-DF (Ref. 3) allows comparison with MP2 result (Ref. 1) (numbers in parentheses), and relative energy values $E_{\text{rel diss}}$ with results from a recent CCSD(T) calculation (Ref. 2) (numbers in square brackets).

<table>
<thead>
<tr>
<th></th>
<th>Prism</th>
<th>Cage</th>
<th>Book</th>
<th>Cyclic</th>
</tr>
</thead>
<tbody>
<tr>
<td>$E_{\text{diss}}$</td>
<td>$-280(-332)$</td>
<td>$-279(-332)$</td>
<td>$-277(-330)$</td>
<td>$-269(-324)$</td>
</tr>
<tr>
<td>R–O–H (Å)</td>
<td>$1.8(1.7)$</td>
<td>$1.9(1.9)$</td>
<td>$1.8(1.8)$</td>
<td>$1.8(1.7)$</td>
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