Balance of Attraction and Repulsion in Nucleic-Acid Base Stacking: CCSD(T)/Complete-Basis-Set-Limit Calculations on Uracil Dimer and a Comparison with the Force-Field Description

Claudio A. Morgado, †,‡ Petr Jurečka,†,§ Daniel Svozil,† Pavel Hobza,‡,§ and Jiří Sponer*,†,‡

Institute of Biophysics, Academy of Sciences of the Czech Republic, Královolopolská 135, 612 65 Brno, Czech Republic, Institute of Organic Chemistry and Biochemistry, Academy of Sciences of the Czech Republic, Flemingovo náměstí 2, 166 10 Prague 6, Czech Republic, and Department of Physical Chemistry, Palacky University, tr. Svobody 26, 771 46, Olomouc, Czech Republic

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Abstract: We have carried out reference quantum-chemical calculations for about 100 geometries of the uracil dimer in stacked conformations. The calculations have been specifically aimed at geometries with unoptimized distances between the monomers including geometries with mutually tilted monomers. Such geometries are characterized by a delicate balance between local steric clashes and local unstacking and had until now not been investigated using reference quantum-mechanics (QM) methods. Nonparallel stacking geometries often occur in nucleic acids and are of decisive importance, for example, for local conformational variations in B-DNA. Errors in the short-range repulsion region would have a major impact on potential energy scans which were often used in the past to investigate local geometry variations in DNA. An incorrect description of such geometries may also partially affect molecular dynamics (MD) simulations in applications when quantitative accuracy is required. The reference QM calculations have been carried out using the MP2 method extrapolated to the complete basis-set limit and corrected for higher-order electron-correlation contributions using CCSD(T) calculations with a medium-sized basis set. These reference calculations have been used as benchmark data to test the performance of the DFT-D, SCS(MI)-MP2, and DFT-SAPT QM methods and of the AMBER molecular-mechanics (MM) force field. The QM methods show close to quantitative agreement with the reference data, albeit the DFT-D method tends to modestly exaggerate the repulsion of steric clashes. The force field in general also provides a good description of base stacking for the systems studied here. However, for geometries with close interatomic contacts and clashes, the repulsion effects are rather severely exaggerated. The discrepancy reported here should not affect the overall stability of MD simulations and qualitative applications of the force field. However, it may affect the description of subtle quantitative effects such as the local conformational variations in B-DNA. Preliminary calculations for two H-bonded uracil base pairs, including one with a C–H···O H-bond, indicate excellent performance of the tested QM methods for all intermonomer distances. The force field, on the other hand, is less satisfactory, especially in the repulsive regions.

Introduction

Base stacking interactions provide a substantial part of the thermodynamics stability of nucleic acids, shape their structure, and contribute to their dynamics.1–7 QM calculations with the inclusion of electron-correlation effects belong to important tools that help to understand the role of stacking interactions in nucleic acids.8–18 Although the gas-phase QM calculations do not directly correlate with the thermodynamics stability of nucleic acids due to the complex interplay between molecular forces in solvated nucleic acids, they reveal direct and accurate structure-energy relationships which allow for an exhaustive description of the potential
energy surfaces (PES) of stacked structures of dimers of nucleic-acid bases.9 Such calculations characterize the intrinsic forces between the stacked bases, are indispensable to reveal the nature of base stacking, and provide reference data to parametrize/verify other computational methods. Such calculations can also help to partially rationalize experimental data on nucleic acids, albeit direct transferability of the QM data to nucleic acids would require, e.g., inclusion of solvent screening of the gas-phase electrostatics which is specific for distinct nucleic-acid architectures.20,21 The QM description of base stacking in general requires application of electron-correlation techniques and diffuse-polarization basis sets of atomic orbitals. The present golden standard for base-stacking computations is the CBS(T) method,19 which is based on MP2 calculations with complete basis-set (CBS) extrapolation corrected for higher-order electron-correlation contributions using the CCSD(T) Cizek method23 with a medium-size basis set. In the present paper we have carried out an extensive QM characterization of stacked conformations of the uracil dimer, which is the smallest base stacking system found in nucleic acids. For the purpose of comparison we have also investigated two H-bonded dimers.

Uracil-Dimer Stacking in Nucleic-Acid Structures. The stacking of uracil bases can be seen in a wide variety of RNA structures, spanning a large range of different conformations. Besides the intrastrand and interstrand uracil dimers occurring in canonical 5′-UU-3′, 5′-AU-3′, and 5′-UA-3′ A-RNA base-pair steps, many diverse stacking arrangements between two uracil bases are observed in the experimental structures of RNA hairpins,25,26 pseudoknots,27,28 dimeric RNA quadruplexes,29 SRE RNA,30 U2 snRNA stem I from S. Cerevisiae,31 a satellite tobacco mosaic virus/RNA complex,32 and many other RNA systems including obviously the ribosomal structures.33,34 Nevertheless, the main aim of this paper is not to investigate the stacking of uracil bases in particular RNAs but to better describe its physicochemical nature and assess the ability of other methods to evaluate stacking interactions, including a standard MM force field.

Preceding Theoretical Studies. The first electron-correlation characterization of the uracil dimer was reported by Šponer et al.35 using the MP2/6-31G**(0.25) method. The study concluded that the stability of stacking originates in the dispersion attraction and that the orientation dependence stems from the electrostatic term. The calculations ruled out several incorrect models of stacking and showed a surprisingly good performance of common molecular-mechanics force fields combining Lennard-Jones potentials with Coulombic terms with atom-centered point charges. Kratc'hy et al.39 investigated the potential- and free-energy surfaces of the uracil dimer in the gas phase, employing a combination of ab initio, empirical potential, computer simulations, and statistical thermodynamics techniques. They reported eleven low-energy minima in the PES of the uracil dimer: seven H-bonded, one T-shaped, and three stacked structures. The global gas-phase free-energy minimum was predicted to be an H-bonded structure, while stacked structures were found to be less populated than H-bonded ones. N1-methylation shifted the free energy balance in favor of the stacked structures, due to the increased dispersion.
measured the gas-phase infrared spectra of jet-cooled uracil clusters, thymine clusters, and uracil-water clusters. The authors found evidence for the presence of several double hydrogen-bonded uracil dimers and for the formation of a larger highly symmetrical cluster. Interestingly, no evidence was seen for T-shaped and stacked structures, which does not support the predictions made by Kratochvíl et al.\textsuperscript{39} Regarding experimental data in the condensed phase, an estimate of vertical stacking interactions of uridine in aqueous solution can be found in the work of Ts’o et al.,\textsuperscript{37} who reported for this nucleoside a free energy of association of 290 cal mol\textsuperscript{−1}. This number cannot be directly compared to gas-phase calculations either, because it includes the entropy cost of bringing the monomers together and many other terms related to solvation.

**The Scope of the Present Study.** The present study is aimed differently than the preceding ones. We have shown that AMBER, the leading molecular-mechanics force field for nucleic acids, provides a surprisingly good description of base stacking.\textsuperscript{35} However, this does not mean that the force field provides an exact description of stacking. In fact, the MM force fields, albeit often providing a very insightful description of nucleic-acid structure and dynamics, are known to have limitations. Besides the overall topologies, nucleic-acid structures and functions are affected by subtle structural details. In canonical double helices these variations are known as local conformational variations, i.e., modest deviations from the average helices that are determined by base sequence and other molecular interactions, such as those caused by protein or drug binding, or crystal packing forces in X-ray experiments.\textsuperscript{58–61} Accurate local positioning of bases is also important elsewhere, for example in the catalytic centers of ribozymes.\textsuperscript{62,63} Local conformational variations are assumed to be primarily caused by base stacking forces and are of primary importance for indirect readout of proteins, sequence-dependent DNA elasticity, etc. Studies, experiments as well as theory, of local conformational variations turned out to be very difficult, as local conformational variations are associated with very subtle energy changes. Computational studies ranging from extensive analysis of the base-stacking PESs up to full-scale MD simulations could provide insights into the sequence dependence of NA structure.\textsuperscript{64–72} However, such quantitative studies would require an exceptionally high accuracy of the energy description of direct base—base interactions, solvent screening effects, and the conformational space of the sugar—phosphate backbone. Regarding base stacking, local conformational variations are often associated with interactions involving nonparallel (mutually tilted) bases, where close contacts (steric clashes) between nucleobase edges or exocyclic functional groups are combined with local unstacking in other parts of the stacked base-pair steps. Steric effects associated with the amino groups of guanine in the minor groove of CpG B-DNA steps\textsuperscript{50,69} or helical-twist/base-pair-roll redistribution in alternating pyrimidine-purine A-RNA sequences belong to the best documented examples.\textsuperscript{58,73,74} Nonparallel stacked bases are obviously very common in complex noncanonical RNA regions. When the bases are not parallel, the stacking geometry typically reflects a competition between a segment of the dimer where the monomers are locally unstacked and another segment where the monomers are sterically clashing. An accurate description of this local compression (clash) is important for a proper description of the whole stacked system. We have recently investigated several geometries of the CpG B-DNA steps using the CBS(T) reference method, and these calculations indeed suggested that once geometries with nonparallel bases are considered, the differences between the benchmark calculations and the force field (as well as other methods) can be significant.\textsuperscript{75} For a proper description of local conformational variations the accuracy of the van der Waals (vdW) term of the force field is critical, as it determines the balance between the steric clashes and the partially unstacked regions. The electrostatic part of the stacking energy with its \( r^{-1} \) dependence is not contributing significantly to the energy changes associated with the steric contacts. In addition, the electrostatic components of stacking in DNA are in general effectively attenuated due to solvent screening. The solvent screening actually limits the direct applicability of gas-phase QM calculations in studies of DNA local conformational variations, because the dominant role of electrostatics for the gas-phase stacking-energy dependence on helical twist vanishes in solvated nucleic acids.

Herein we report an interaction-energy analysis of the PES of the uracil dimer in stacked conformations, covering a wide range of about 100 structures with specific emphasis given to geometries of dimers in tilted conformations and with unoptimized vertical separation. The reference points are obtained with the CBS(T) method. We first scan a four-dimensional space considering twist between parallel bases, displacements in \( x \) and \( y \) directions, and the vertical separation between the monomers. Then, we investigate several geometries with tilted bases, i.e., structures with competing clashing and unstacking. While purely compressed or extended dimers with parallel bases cannot occur in real nucleic-acid structures, structures with nonparallel bases can be accompanied with close interatomic contacts (steric clashes) and local unstacking even upon the overall optimization of the vertical separation between the bases or base-pair steps.\textsuperscript{76} The reference CBS(T) calculations serve as a benchmark for several other computational methods: the AMBER force field,\textsuperscript{38} DFT-D,\textsuperscript{53} SCS(MI)-MP2,\textsuperscript{77} and DFT-SAPT.\textsuperscript{78} The choice of the AMBER force field is dictated by the need to evaluate the performance of its nonbonded empirical potential—whose vdW term is very similar to the vdW term of most other MM force fields—in a wider region of the PES of nucleic-acid base dimers. The DFT-D and the SCS(MI)-MP2 methods have been chosen because they have proven a cost-efficient way for obtaining interaction energies in good agreement with CCSD(T) or CBS(T) benchmarks. However, both methods have been parametrized against the S22 training set, which only contains noncovalent complexes optimized to a minimum, and their performance in regions other than the minimum has not been as widely tested. We have performed DFT-SAPT energy calculations and decompositions for all of the structures in order to gain some insight into the nature of the interaction.
Among other results we show that the MM force-field calculations are basically capable of providing a satisfactory description of base stacking for the uracil dimer. However, the agreement between the force field and the reference CBS(T) calculations breaks down in the repulsive regions of the PES. The observed differences are caused by the vDW term of the force field and are large enough to substantially affect the description of the fine local conformational variations in nucleic-acid duplexes. Therefore, the correct description of such geometries may represent one of the challenges for future refinements of MM force fields. The DFT-D, SCS(MI)-MP2, DFT-SAPT, and CBS(T) methods are mutually much more consistent in the repulsive region, although the differences between them also increase upon incrementing the repulsion. We have also investigated the dependence of the interaction energy on the intermonomer distance for two H-bonded uracil dimers. The mutual agreement between the QM methods is surprisingly good. On the contrary, the discrepancy between the AMBER force field and the QM methods for short intermonomer distances is much more accentuated, with the force field severely exaggerating the repulsion as the H-bonding distance is decreased. Nevertheless, it is important to point out that, considering the reliability of molecular modeling, the inaccuracy of the force-field description for the H-bonded base pairs is not as painful as the differences reported for stacking. The H-bonded base pairs, in contrast to stacking, usually represent interactions that are well separated from the other interactions. Thus the exaggeration of short-range repulsion in base pairs is likely to lead to mere overestimation of H-bond lengths with no significant effects on the local conformational variations. We would like to note here that our present study focuses on the stacking interactions and that the H-bonded data are more limited. The balance of forces in H-bonded base pairs is very different from that in the stacked structures. This topic requires a thorough analysis, and work in this direction is currently underway. For the information concerning the accuracy of the DFT-D methods for both H-bonded and stacked complexes we refer the reader also to studies that are available in the literature. For instance, Sherrill showed that Grimme’s DFT-D (PBE-D) performs very well for both H-bonded and stacked complexes. A feeling about the spin-component-scaled methods can be obtained from refs 13 and 80. Finally, the DFT-SAPT method has been shown to give highly accurate results for both H-bonded and π-bonded complexes, and the accuracy of the intermolecular components has also been studied for both cases.

Computational Details

Geometries. The structures have been obtained in the following way. First, a uracil monomer is optimized using the RI-MP2 method along with the cc-pVTZ basis set. Then, it is placed in the xy plane with the center of mass coinciding with the origin. The N1=H1 “glycosidic” bond is parallel to the y-axis and is pointing to the direction of negative y values, and the Watson–Crick face of the base is oriented toward the direction of negative x values (Figure 1), with a minor modification for the structures derived to study interatomic clashes (NP1-NP4, see below and Supporting Information). This first monomer is always fixed, and the second monomer is initially superimposed on the first one. Then the position of the second monomer is determined via 6 independent parameters. There are three angles that are applied counterclockwise and consecutively via rotational matrices: rotation around the x axis (γ), rotation around the y axis (α), and rotation around the z axis (ω). The first two angles are applied to introduce the tilting between bases. When comparing to B-DNA conformational parameters, the α, γ, and ω angles are roughly analogous to propeller twist, buckle, and helical twist angles. Then, the second monomer is shifted along the x and y axes by the parameters Δx and Δy. Finally, the vertical distance is adjusted by Δz. This means that in all cases Δz is equal to the distance between the center of mass of the second monomer and the xy plane, while the first monomer is located (note that Δz ≡ r in the figures of the stacked dimers). As the stacking energy is very sensitive to vertical compression or extension of the dimer and the vertical dependence of the stacking energy reflects the balance of vdW interaction terms, we have scanned the stacking-energy dependence on Δz for most combinations of α, γ, ω, Δx, and Δy.

The structures are grouped into two different sets, P and NP. The set P contains 9 different uracil dimers (P1, P2, ..., P9), in conformations where the planes of the rings are parallel (P) to each other (α, γ = 0), whereas the set NP contains 4 different dimers (NP1, NP2, NP3, and NP4) in conformations in which one monomer is tilted with respect to the other one, that is, nonparallel (NP) conformations. A dimer is defined by those structural parameters that are kept fixed during the scan, where a free parameter is varied in order to build a potential-energy curve for each dimer. This way we have generated a subset of structures for each dimer, and these subsets add up to a total of 105 distinct geometries. With one exception (P3, for which we vary the twist angle) we have scanned the Δz dependence of the stacking energy for a fixed combination of the other five geometrical parameters.

The four tilted dimers NP1-NP4 have been selected manually based on visual inspection of a number of α, γ, ω, Δx, and Δy combinations in order to obtain four vertical scans with diverse clashes. Obviously, one could imagine...
many other such geometries that would be worthy of study; however, the calculations remain expensive, and thus we have limited the number of studied structures.

For the H-bonding calculations the WC and “Calcutta” dimers have been selected (denoted HB1 and HB2, respectively), and the scans have been carried out by varying the O2···N3 and the O4···N3 distances, respectively. Note that the Calcutta base pair contains a C−H···O H-bond. The initial (equilibrium) geometries of both H-bonded dimers are taken from ref 84.

The Cartesian coordinates of all structures are given in the Supporting Information, while Figures 2 and 3 show the geometrical arrangement of the stacked and H-bonded dimers respectively.

**Figure 2.** Top and side views of the parallel and nonparallel structures of the uracil dimer, respectively. The first uracil monomer is shown in gray. Some atoms have been labeled in order to aid in the visualization.

**Interaction Energies.** The interaction energy $\Delta E^{AB}$ of a stacked or H-bonded complex is calculated according to the supermolecular approach as the difference in electronic energy between the complex and the isolated monomers (eq 1).$^{85,86}$ Since we have used only rigid monomers, we do not consider monomer deformation energies. In the case of the SCS(MI)-MP2 and CBS(T) methods, all of the interaction energies have been corrected for the basis set superposition error (BSSE) using the counterpoise (CP) technique.$^{87,88}$ This correction is not applied within the DFT-D formalism as this method was parametrized with respect to BSSE-corrected data.$^{53}$

$$\Delta E^{AB} = E^{AB} - E^{A} - E^{B}$$ (1)
Then the force-field term is derived at the same computational level as used of the force field requires that the force-field electrostatic demonstrated in the literature, meaningful benchmarking nonbonded potential of the AMBER force field. As paper is to validate the physical correctness of the deserves an explanation. One of the main purposes of this methods used in the present work, except for DFT-SAPT, where the interaction energy is given by the sum of selected perturbation contributions. In an intermolecular perturbation theory like DFT-SAPT the BSSE takes place only in the $\delta HF$ term (see below).

**Methods**

**AMBER Nonbonded Empirical Potential.** The empirical-potential calculations of the interaction energies have been carried out with a local code, employing the vdW and Coulombic terms of the AMBER force field. The atom-centered point charges were derived by means of an electrostatic potential (ESP) fitting at the MP2/aug-cc-pVDZ level of theory using the Merz–Kollman methodology as implemented in the Gaussian 03 software package. This nonbonded potential is analogous to that in AMBER, with the only exception that condensed-phase simulations with AMBER are performed with charges derived in a RESP fitting with the Hartree–Fock (HF) method and the 6-31G* basis set. Given that the HF method overestimates dipole moments, the use of HF charges implicitly accounts for polarization.

The choice of the MP2/aug-cc-pVDZ ESP charges instead of the original AMBER HF/6-31G* RESP charges deserves an explanation. One of the main purposes of this paper is to validate the physical correctness of the nonbonded potential of the AMBER force field. As demonstrated in the literature, meaningful benchmarking of the force field requires that the force-field electrostatic term is derived at the same computational level as used in the reference QM calculations. Then the force-field electrostatic parameter set mimics as close as possible the respective Coulombic term of the full QM calculation. When using standard HF/6-31G* AMBER charges, we would inevitably bias the force field vs QM comparison. It would be often difficult to judge whether the observed differences reflect the basic limitation of the force-field model, the presence of interactions not included in the force-field model, or just the fact that the force-field and the QM calculations consider uncorrelated and correlated electrostatics, respectively. In addition, the present paper is focused on evaluating the capability of the van der Waals term of the force field to describe steric clashes. When introducing steric clashes, the electrostatic energy is basically constant. Note that a comparison of HF and MP2 ESP charges for base stacking is available in the literature. In addition, in one of our earlier stacking papers we have also demonstrated that the use of out-of-plane charges or distributed multipoles does not improve the performance of the force field for base stacking compared with the set of atom-centered ESP-derived point charges. Thus, the use of atom-centered MP2/aug-cc-pVDZ charges minimizes the difference between the force-field and QM description caused by the ESP electrostatics, allowing unambiguous assessment of the van der Waals part of the force field.

**DFT Augmented with an Empirical Dispersion Term (DFT-D).** The DFT-D calculations have been performed with the TurboMole 5.8 software package, in combination with a local code that computes the empirical dispersion correction. For the DFT part of the calculation we have used the RI approximation also known as density fitting. In the DFT-D approach as developed by Jurečka et al., a pairwise additive potential of the form $C_{ij}r^{-6}$ is used to account for long-range dispersion effects that are poorly described with common density functionals. The dispersion-corrected energy is given by

$$E_{DFT-D} = E_{KS-DFT} + E_{disp}$$

where $E_{KS-DFT}$ is the self-consistent Kohn–Sham energy, and $E_{disp}$ is a term containing the empirical dispersion correction:

$$E_{disp} = -\sum_i f_{damp}(r_{ij}) R_{ij}^6 C_{6,ij} r_{ij}^{-6}$$

$$f_{damp} = \frac{1}{1 + e^{-d/r_{ij}([S_R\lambda])^{-1}}}$$

In eq 3 $r_{ij}$ is the distance between atoms $i$ and $j$, $R_{ij}^6$ is the equilibrium vdW separation between atoms $i$ and $j$ (derived from the atomic vdW radii), and $C_{6,ij}$ is the composite dispersion coefficient for the pair of atoms $i$ and $j$ (calculated from the corresponding atomic $C_6$ coefficients). The damping function (eq 4) is needed because the $r^{-6}$ form is not valid at short distances, and because some short-range correlation effects are already present in the density functional. In this equation $d$ is a parameter determining the steepness of the damping function, and $S_R$ is a scaling coefficient that adjusts the magnitude of the vdW radius and that has been determined for several density-functional/basis-set combinations. The values of the $C_6$ atomic coefficients have been taken from the work of Grimme, whereas the $d$ and $S_R$ parameters as well as the combination rules for the vdW...
radii and the composite dispersion coefficients are the same as those in the work of Jurečka et al. Here we use the TPSS functional along with the 6-311++G(3df,3pd) basis set. This level of theory complemented with the dispersion correction will hereafter be referred to as DFT-D.

**SCS-MP2 for Molecular Interactions (SCS(MI)-MP2).** The SCS(MI)-MP2 calculations have been performed with the Molpro 2006.1 software package, applying the frozen-core and density-fitting approximations. This method is a reparameterization of the original SCS-MP2 method, in which the same- (SS) and opposite-spin (OS) components of the MP2 energy are empirically scaled (\(E_{\text{MP2}} = c_{\text{SS}}E_{\text{OS}} + c_{\text{SS}}E_{\text{SS}}\)) in an attempt to overcome the deficiencies of the MP2 theory, like the overestimation of the dispersion contribution to the correlation energy. The original SCS-MP2 method reduces the overestimation of the dispersion energy for stacked structures and thus provides very good estimates of the stabilization energy for these systems. Unfortunately, the method also reduces the dispersion for H-bonded structures, which results in this case in an underestimation of the stabilization energy. Using multivariate linear least-squares analysis and the CCSD(T) data in the S22 training set, Distasio and Head-Gordon found the optimal parameters that minimized the error between SCS-MP2 theory and CCSD(T), for the cc-pVXZ (\(X=T, Q\)) and extrapolated cc-pV(\(X\)Y)Z (\(X=\text{DT}, \text{TQ}\)) levels. The resulting method, known as SCS(MI)-MP2, provides very good estimates of stabilization energies for both planar H-bonded and stacked structures. Here we calculate the SCS(MI)-MP2 interaction energies using the cc-pV(\(X\)Y)Z extrapolation (cc-pV(\(X\)Y)Z→cc-pVXZ), for which the same- and opposite-spin optimized parameters are 1.46 and 0.29, respectively. The MP2/CBS limit is approximated according to the following extrapolation scheme

\[
E_{\text{XY}} = E_{\text{SCF},XY} + \frac{X^3E_{\text{CORR},X} - Y^3E_{\text{CORR},Y}}{X^3 - Y^3} \quad Y > X (5)
\]

where \(X = 2\) and \(Y = 3\) for the D→T extrapolation used in this work.

**DFT-Symmetry Adapted Perturbation Theory (DFT-SAPT).** The DFT-SAPT calculations have been performed with the Molpro 2006.1 software package. DFT-SAPT is a method that uses molecular properties from density functional theory in order to calculate intermolecular interaction energies by means of symmetry-adapted perturbation theory (SAPT). In this method the interaction energy is given as the sum of the first- and second-order energies, plus the \(\delta HF\) term. The first-order energy includes the electrostatic and exchange-repulsion contributions, while the second-order energy includes the induction, exchange-induction, dispersion, and exchange-dispersion contributions. The \(\delta HF\) term is an estimate of higher-order Hartree–Fock contributions and is determined as the difference of the HF interaction energy and the sum of the first- and second-order contributions, with the exception of the dispersion and exchange-dispersion energies. Since the HF interaction energy is calculated with BSSE-corrected monomer energies, the \(\delta HF\) term is BSSE dependent. The interaction energy is given by eq 6, and the electrostatic, induction, dispersion, and exchange contributions are defined in eqs 7–10.

\[
E_{\text{int}} = E_{\text{pol}}^{(1)} + E_{\text{ex}}^{(1)} + E_{\text{ind}}^{(2)} + E_{\text{ex-ind}}^{(2)} + E_{\text{disp}}^{(2)} + E_{\text{ex-disp}}^{(2)} + \delta HF (6)
\]

\[
E_{\text{elec}} = E_{\text{pol}}^{(1)} (7)
\]

\[
E_{\text{ind}} = E_{\text{ind}}^{(2)} + E_{\text{ex-ind}}^{(2)} (8)
\]

\[
E_{\text{disp}} = E_{\text{disp}}^{(2)} + E_{\text{ex-disp}}^{(2)} (9)
\]

\[
E_{\text{exch}} = E_{\text{ex}}^{(1)} (10)
\]

For the energy decomposition we have employed the LPBE0ACXC potential with the pure ALDA kernel for both the static and dynamic response, along with the aug-cc-pVXZ basis set. We have also utilized the density-fitting approximation. The ionization potential (IP) of the monomer and the energy of the highest occupied molecular orbital (HOMO), which are required to evaluate the shift parameter, have been calculated at the PBE0/aug-cc-pVXZ level of theory.

The relatively small aug-cc-pVXZ basis set has been used because our primary goal is to get an idea about the relative magnitude of the interaction energy components. DFT-SAPT interaction energies are usually underestimated when using this basis set, mainly because the dispersion component is underestimated by about 10–20%. This underestimation does not change the conclusions regarding the relative importance of the individual contributions to the interaction energy given below. With a larger basis set DFT-SAPT was shown to provide very accurate total interaction energies in good agreement with the most accurate CCSD(T) calculations. Using the aug-cc-pVXZ basis set for the IP calculations does not affect the shift values dramatically, and resulting errors should be smaller than the SAPT basis-set-size errors.

**MP2/CBS Corrected for Higher-Order Correlation Effects (CBS(T)).** The MP2 and CCSD(T) calculations have been performed with the Molpro 2006.1 software package, applying the frozen-core approximation. In the case of MP2, we have also applied the density-fitting approximation. We have carried out the MP2/CBS calculations by extrapolating the Hartree–Fock and the correlation energies separately, following the Helgaker et al. extrapolation scheme (eqs 11 and 12), in which \(E_X\) is the energy of the basis set with the largest angular momentum \(X\), \(E_{\text{CBS}}\) is the energy for the complete basis set, and \(\alpha\) is a parameter that was fitted in their original work. Herein we have used the aug-cc-pVXZ→aug-cc-pVXZ extrapolation.

\[
E_{\text{HF}}^{\text{CBS}} = E_{\text{HF}}^{\text{CBS}} + A e^{-\alpha X} (11)
\]

\[
E_{\text{CORR}}^{\text{CBS}} = E_{\text{CORR}}^{\text{CBS}} + BX^{-3} (12)
\]

Given that higher-order correlation-energy contributions cannot be neglected, we have approximated the CCSD(T)/CBS interaction energies according to the following scheme:

\[
\Delta E_{\text{CBS}}^{\text{CCSD(T)}} = \Delta E_{\text{CBS}}^{\text{MP2}} + (\Delta E_{\text{CCSD(T)}}^\text{CBS} - \Delta E_{\text{MP2}}^\text{CBS}) \bigg|_{6-31+G^{**}} (13)
\]
The use of eq 13 is based on the assumption that the difference between the CCSD(T) and MP2 interaction energies is basically basis-set independent and can therefore be evaluated with a small- or medium-size basis set. This assumption was validated for several H-bonded and stacked clusters, including H-bonded and stacked structures of the uracil dimer. A CCSD(T)/CBS interaction energy estimated according to eq 13 is also known as CBS(T). 15

Data Analysis. The interaction-energy data have been analyzed in terms of root-mean-squared (rms) errors, maximum absolute deviations (MAX), and XY plots. rms errors are calculated using the standard formula. In this work we define a deviation as the difference between the value obtained with a given method (AMBER, DFT-D, SCS(MI)-MP2, or DFT-SAPT) and the value obtained with CBS(T), and the MAX value for a given dimer simply corresponds to the absolute value of the largest deviation. All of the plots have been generated with the Matplotlib graphics package in combination with the SciPy and NumPy packages. Energies and plots are given in the Supporting Information.

Results and Discussion

The Overall Performance of the Methods. Let us first assess the data qualitatively. Figure 4 shows the dependence of the stacking energy on twist at \( r = 3.3 \ \text{Å} \) (P3), which is primarily determined by the electrostatic term. The twist energy variation is rather modest due to the lower polarity of uracil compared to cytosine and guanine.35 The DFT-D method achieves a close to exact match to the CBS(T) curve. The AMBER force field is more stabilizing than CBS(T), and also the profile of the twist dependence somewhat deviates from the CBS(T) data. The SCS(MI)-MP2 and DFT-SAPT methods systematically underestimate the binding, while the shapes of the curves basically match the CBS(T) one. In the case of the DFT-SAPT method the underestimation is mainly due to the small basis set used (aug-cc-pVDZ); with a larger basis set yet closer match with the CBS(T) data is expected.

Figure 5 shows the vertical scans for P1 and P2, i.e., the untwisted and antiparallel undisplaced dimers (untwisted and antiparallel refer to \( \omega \) angles of 0 and 180°, respectively). When assessing the methods using the vertical scans, the most important region is that \( \sim 0.2–0.3 \ \text{Å} \) around the CBS(T) minimum, while the most relevant descriptor is the slope (gradient) of the PEC at a given distance within the repulsive region. This is related to the force associated with the repulsion. Two findings are apparent. First, all methods show very flat energy dependence around the vertical-separation minimum for the untwisted P1 dimer, which has the most repulsive electrostatic arrangement. The AMBER force field gives the shortest optimal vertical separation of the monomers, which has the most repulsive electrostatic arrangement. The AMBER force field gives the shortest optimal vertical separation of the monomers for this arrangement. In contrast, the force field has an excessively steep onset of the repulsion in the short-range repulsion region of the P2 dimer which is also associated with the overestimation of the optimal intermonomer distance. The data for the P2 dimer also suggest that DFT-D is little overestimating the repulsion in the short-separation region, while SCS(MI)-MP2 provides a curve with almost the same slope as the CBS(T) one, albeit the method underestimates the interaction energy. DFT-SAPT also underestimates the interaction energy.

Figure 6 shows the vertical scans for the two untwisted-displaced dimers (P4 and P5). These are still in the electrostatically repulsive orientation, but the geometrical overlap of the bases is reduced. The most visible result is the overestimation of the short-range repulsion and of the optimal intermonomer distance by the force field. Modest exaggeration of the repulsion by the DFT-D method is also seen. The SCS(MI)-MP2 and DFT-SAPT methods (in particular the latter, due to the relatively small basis set used) are typically shifted toward higher stacking energies for all intermonomer distances. Interestingly, the difference between DFT-SAPT and CBS(T) widens visibly upon reducing the intermonomer separation.

Figure 7 summarizes the vertical scans on the four antiparallel displaced dimers (P6–P9). There is again a large exaggeration of the short-range repulsion by the force field. Figure 8 gives the data associated with the vertical scans of the four nonparallel (tilted) dimers with steric clashes (NP1–NP4). The NP1 geometry brings the O4 region of the.
first monomer into contact with the N1 region of the second one. In the NP2 geometry the N1/O2 region of the second monomer is in close contact with the aromatic ring of the first one. The NP3 geometry leads to the closest contact between the O2 atom of the second monomer and the C4 atom of the first one, and the NP4 geometry has the N3–H3 bond of the second monomer tilted to the center of the aromatic ring of the first one. The twist angle is 180° for all of the four clashed dimers. Obviously, these four dimers do not guarantee an exhaustive sampling of all possible steric contacts between the two uracils; however, we hope that they are diverse enough to allow us to make correct conclusions. The calculations are quite consistent with the results derived from the calculations on the P1–P9 dimers. The large deviation (excessive repulsion) of the AMBER force field in the short-range repulsion region is very clear, and the difference between DFT-D and CBS(T) is also visible. DFT-D overestimates the binding at larger separations, but then it has a faster and steeper onset of the repulsion upon vertical compression of the dimers. SCS(MI)-MP2 and DFT-SAPT underestimate the interaction energy and the difference between either of these methods, and CBS(T) increases as the intermolecular distance decreases. Both effects are more pronounced for DFT-SAPT, mainly because of the basis set chosen for these calculations.

Finally, Figure 9 gives the energy scans for the WC (HB1) and Calcutta (HB2) H-bonded uracil dimers. The problems of the AMBER force field to describe the repulsive region of the PES are even more visible than for the stacked dimers, in particular for the Calcutta structure. On the QM side there is essentially a very good agreement between the CBS(T), DFT-D, and SCS(MI)-MP2 methods. DFT-SAPT underestimates the strength of the H-bonding interaction at all distances. However, we need to reiterate that DFT-SAPT calculations were done with a relatively small aug-cc-pVDZ basis set. Specifically for H-bonding, very large basis sets with higher-angular momentum functions are vital. The repulsion appears to be well captured by DFT-SAPT. We would like to point out again that the H-bonding data presented here is still preliminary. The issue of the description of the short-range repulsion in H-bonding is even more complex than it is for stacking and will be addressed in the future as it is beyond the scope of the present stacking study.

**Interaction-Energy Statistics.** Rms errors with respect to the CBS(T) reference data are given in Table 1 along with the corresponding MAX values. For the twist-angle scan (P3) the best performer is the DFT-D method, with a rms error of only 0.15 kcal mol⁻¹ and a MAX value of 0.23 kcal mol⁻¹. Both SCS(MI)-MP2 and DFT-SAPT underestimate the strength of the interaction across the entire range of twist angles, showing rms errors of 0.74 and 1.39 kcal mol⁻¹ respectively. On the contrary, AMBER overestimates the attraction with a rms error of 0.61 kcal mol⁻¹.
Turning now to the performance of the QM methods in the vertical scans of the parallel dimers, we see that the DFT-D method is again the best performer with regards to the chosen statistical descriptors, giving for example the smallest rms deviation (0.52 kcal mol$^{-1}$). DFT-D overestimates the strength of the interaction at long distances and underestimates it at short distances (refer to Figures 5–8). The DFT-D deviations around the PEC minimum are typically very small. This is not surprising as the parametrization of DFT-D relies mostly on a training set containing noncovalent complexes optimized to a minimum. The fact that DFT-D always underestimates the strength of the interaction at short distances may be rooted in the inability of the density functional to completely account for short-range correlation effects, which are not compensated for by the empirical correction. SCS(MI)-MP2 also shows a small rms deviation with respect to the CBS(T) data (0.86 kcal mol$^{-1}$), but, unlike DFT-D, the shape of the SCS(MI)-MP2 curve closely resembles the CBS(T) one for all of the scans. This means that in general this method may produce better energy gradients, in particular in the repulsive regions of the PES. This method, however, systematically underestimates the strength of the interaction. Finally, DFT-SAPT exhibits the largest rms error among the QM methods (1.37 kcal mol$^{-1}$), and the method systematically underestimates the strength of the interaction. Note, however, that the DFT-SAPT data were obtained with a relatively small basis set (aug-cc-pVDZ), while the other methods are calculated (directly or effectively) with a CBS extrapolation.

The AMBER force field shows larger deviations with respect to CBS(T) than the QM methods, with a rms error of 1.53 kcal mol$^{-1}$. AMBER overestimates the strength of the interaction at long distances and underestimates it at short distances, the only exception being the untwisted undisplaced P1 dimer, where the AMBER energies are lower along the entire range of distances. It partially could be due to increased polarization effects in this particular geometry that would not be captured by a nonpolarizable force field. However, the DFT-SAPT data below do not indicate a significant role of induction. Most likely, the overestimation of the strength of the interaction is due to the underestimation of the repulsion by AMBER, which in turn might be a consequence of an improper description of the anisotropy caused by a particularly unfavorable Pauli repulsion in the P1 conformation. The force field assumes isotropic interactions and spherical atoms, which might not be accurate enough in some specific geometries.

Similar findings are seen in the case of the vertical scans for the nonparallel dimers. With a rms error of 0.55 kcal mol$^{-1}$ DFT-D is again the best performer with respect to this statistical descriptor. The method remains to overestimate the strength of the interaction at long distances and underestimate it at short distances. The performance of SCS(MI)-MP2 is also good, bearing the smallest MAX value and a
Figure 7. Potential energy curves (left) and energy differences with respect to the CBS(T) data (right) for P6 through P9, calculated with the AMBER, DFT-D, SCS(MI)-MP2, DFT-SAPT, and CBS(T) methods.
Figure 8. Potential energy curves (left) and energy differences with respect to the CBS(T) data (right) for NP1 through NP4, calculated with the AMBER, DFT-D, SCS(MI)-MP2, DFT-SAPT, and CBS(T) methods.
rms error of 0.81 kcal mol\(^{-1}\). The same error amounts to 1.13 kcal mol\(^{-1}\) in the case of DFT-SAPT. Moreover, the shapes of the SCS(MI)-MP2 curves again resemble the CBS(T) ones, and, particularly when going from the minimum to the repulsive region (see the NP4 curve in Figure 8), this method produces better energy gradients than DFT-D. AMBER shows a larger rms error of 1.77 kcal mol\(^{-1}\), and its deviations are more pronounced in the repulsive region of the PECs. In order to assess the importance of the results, we need to consider which geometries can be

![Figure 9. Potential energy curves (left) and energy differences with respect to the CBS(T) data (right) for HB1 and HB2, calculated with the AMBER, DFT-D, SCS(MI)-MP2, DFT-SAPT, and CBS(T) methods.](image)

**Table 1.** Root-Mean-Squared Deviations (rmsd, kcal mol\(^{-1}\)) and Maximum Absolute Deviations (MAX, kcal mol\(^{-1}\)) of the AMBER, DFT-D, SCS(MI)-MP2, and DFT-SAPT Interaction Energies with Respect to the CBS(T) Reference Data

<table>
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<tr>
<th>dimer</th>
<th>AMBER</th>
<th>DFT-D</th>
<th>SCS(MI)-MP2</th>
<th>DFT-SAPT</th>
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* Some extreme MAX values are not visualized in the figures because they are out of the range of intermolecular distances in those figures. Refer to Tables S2, S3, and S4 in the Supporting Information for the complete data. *P3 was not included in the calculation of rmsd and MAX values, i.e., the values in this row correspond to the group of dimers for which vertical scans have been performed.
populated in real structures. It is clear that severely compressed structures are not likely to be populated, but structures with shortened intermonomer separations with energies 1−2 kcal mol$^{-1}$ above the minima on the CBS(T) curves are certainly accessible. For these geometries the force field repulsion is already visibly exaggerated.

With respect to the H-bonding scans the best performance is achieved by the DFT-D and SCS(MI)-MP2 methods, with rmsd values of about ∼0.5 kcal mol$^{-1}$. Both methods overestimate the strength of the interaction at short distances. DFT-SAPT evaluated with the medium-sized basis set underestimates it at all distances, showing a rmsd value of 1.78 kcal mol$^{-1}$. The large rmsd associated with the AMBER data is due to the huge deviations toward more repulsive values that occur in the regions of short H-bonding distances, in particular for the HB2 structure. The very large repulsion given by the force field for this dimer indicates an excessive repulsion for the C−H···O contact. This can be due to an excessive steepness of the repulsion with the Lennard-Jones 6−12 potential and most likely is also due to a too large atomic radius of that particular hydrogen atom in the AMBER force field. The data show that the force-field performance is better for the stacked dimers than for the H-bonded ones. However, as noted in the introduction, the overestimation of short-range repulsion for H-bonded base pairs may have less serious consequences for the molecular-mechanical studies than errors in the short-range repulsion for stacking. The interaction energy of HB1 at the minimum of the AMBER curve amounts to ∼10.7 kcal mol$^{-1}$, whereas for the same geometry Šponer et al. reported an AMBER energy of −12.1 kcal mol$^{-1}$. This difference is because the latter value was computed with the original AMBER HF charges, which differ from the MP2 charges used in the present study. The HF charge distribution is more polar, which produces a better stabilization.

**Interaction-Energy Gradients.** Regions of interatomic contacts are associated with interaction energy gradients, and their inaccurate description may affect the outcome of the calculations, including the population of conformations in MD simulations. Nevertheless, simulations sample also all the additional degrees of freedom which should, in most cases, attenuate the impact of the incorrect description of energy gradients in the clashed regions. In contrast, potential energy scans that were often used to study the local conformational variations can be severely distorted by the incorrect description of the repulsion as in such scans one does not vary enough degrees of freedom to overcome the exaggerated repulsion. It is thus promising to see that the QM methods perform well for the clashes. For instance, for P8 the difference between the interaction energies at $r = 2.9$ and $r = 3.1$ Å (a range that is within 0.3 Å from the CBS(T) minimum) is 1.92, 2.19, 2.50, and 2.92 kcal mol$^{-1}$, calculated with CBS(T), SCS(MI)-MP2, DFT-SAPT, and DFT-D, respectively. On the contrary, AMBER yields an interaction-energy difference of 4.57 kcal mol$^{-1}$. Clearly, the SCS(MI)-MP2 value is the closest to the CBS(T) one, and this level of accuracy might be necessary to fully understand the intrinsic interactions in nucleic-acid systems such as those mentioned above, where variations in the geometry are often accompanied by very subtle energy differences.

Figure 10 presents an estimate of the interaction-energy gradient for the AMBER and CBS(T) PECs of the NP3 dimer. The force field clearly deviates from the CBS(T) data. The NP3 geometry with vertical distance of 3.5 Å is easily accessible on the CBS(T) PEC. At this distance, however, the force field energy gradient is ∼3 times larger than the CBS(T) one, so this geometry would be penalized when attempting a force field calculation. The NP3 dimer shows probably the most severe differences among the stacked structures; similar plots for the rest of the stacked dimers can be found in the Supporting Information.

The H-bonding dimers show even more drastic differences between the AMBER and CBS(T) gradients (see Figures S12 and S13), with the same consequences as those described above for the stacked dimers.

**DFT-SAPT Interaction-Energy Decomposition.** Figures 11 and S14−S23 in the Supporting Information show the DFT-SAPT decompositions for the stacked dimers. Some general trends have emerged from the decomposition analysis. In basically all the stacked structures the dispersion energy is the leading stabilizing contribution, with the electrostatic term being almost always stabilizing or even repulsive, as in the case of the P1 dimer at large intermolecular distances. Interestingly, for P1 the electrostatics switches from repulsion to attraction at short distances. This can be easily explained by a competition of the multipolar and overlap electrostatic components at short intermonomer distances. The multipolar part is repulsive here due to the very unfavorable dipole orientation. At short distances, however, the attractive overlap electrostatic interaction prevails. The DFT-SAPT electrostatics thus cannot be directly compared with the force field electrostatics. The latter takes into consideration only the electrostatic potentials (ESP), while the overlap electrostatic effects are effectively included in the vdW term of the force field. Therefore, the utilization of DFT-SAPT calculations in parametrizations of simple biomolecular force fields does not appear to be straightforward. It is to be noted that for biomolecular recognition the ESP part of the electrostatics is the most important term. Note that the overlap electrostatics is always
attractive, and the repulsion in the intermolecular complexes is due to Pauli exchange.

For P6, NP1, and NP2 the electrostatic interaction contribution is stabilizing and very significant, with a magnitude that is closer to that of the dispersion term, when compared to the rest of the dimers. For each dimer there is a region where the dispersion and exchange terms tend to cancel each other, usually around the minimum, which is characteristic for vDW complexes. When going into the repulsive part of the PECs the magnitude of the exchange term quickly outweighs that of the dispersion term. It thus should be noted that when assessing the DFT-SAPT decompositions, it is very important to have the dimers at proper vertical separation. This is straightforward for geometries with parallel bases but may be more complex for geometries with tilted bases. DFT-SAPT decompositions for geometries with unrelaxed vertical separation or with unrealistic clash (e.g., due to errors in the experimental structure) can be biased. Taking the variability due to the intermolecular separation into account our results are consistent with those from previous DFT-SAPT studies.\textsuperscript{48,120,121}

For most of the stacked dimers the magnitude of the $\delta HF$ term is comparable to that of the induction term, and both terms represent the smallest contributions to the interaction energy. The horizontal twist scan (P3) simply confirms that the electrostatic term exhibits the largest twist dependence.

\textbf{Figure 11.} DFT-SAPT energy components (left) and comparison between DFT-SAPT and AMBER electrostatic energies (right) for P1, P2, and P3.
(followed by the exchange term to a lesser extent) and that the dispersion term is basically independent of the twist angle. Also, the dispersion and exchange terms basically cancel each other in the range of twist angles from 90 to 180°.

The DFT-SAPT data of the HB1 and HB2 structures basically confirm that the most relevant stabilizing contribution in the case of H-bonding is the electrostatic term and that the induction and $\delta HF$ terms play a more important role than they do for the stacked dimers. Although smaller than the electrostatic term, the dispersion contribution is not negligible, as it amounts to about $-8$ and $-6$ kcal mol$^{-1}$ at the CBS(T) minimum of the HB1 and HB2 dimers, respectively.

**Summary and Conclusions**

We have carried out reference CBS(T) calculations for about 100 geometries of the uracil dimer in stacked conformations. The calculations have been specifically aimed at geometries with unoptimized distances between the monomers including geometries with mutually tilted monomers that are characterized by a delicate balance between local steric clashes and local unstacking. Until now such geometries had not been investigated using reference QM methods and were not included in the parametrization of methods such as DFT-D. Such geometries, however, often occur in nucleic acids and are of decisive importance for local conformational variations in B-DNA as well as in some cases for the exact positioning in B-DNA as well as in some cases for the exact positioning of bases in compactly folded RNAs. Errors in the short-range repulsion region would have a major impact on potential energy scans which were often used in the past to investigate local geometry variations in DNA.
the excessive clash with not much effect on the geometry. It has been noticed that an excessive repulsion associated with too large hydrogens of the thymine methyl group affects substantially 2D stacking energy scans of propeller twisting in ApT and ApA B-DNA steps. However, the reduction of the hydrogen radius, which markedly improved the quality of the empirical force-field stacking energy scans, had no visible effect in test simulations. An exaggerated short-range repulsion in the force field description was also reported for cation-solute interactions and affects for example simulations of quadruplex DNA where the cations residing in the channel of the quadruplex stem look oversized. This effect was originally attributed to the lack of polarization in the force field, but it is more likely caused by the short-range repulsion imbalance discussed in the present paper. Also, when using MP2/aug-cc-pVDZ ESP charges the AMBER force field seems to provide a better gas-phase description for stacking than for H-bonding, at least for the uracil dimer. Despite the reported difference we would like to clearly state that the overall description of base stacking and H-bonding by the nonbonded potential of the AMBER force field is good, and, as we pointed out elsewhere, stacking is probably the best approximated term in the force field. The discrepancy reported here should not affect the overall stability of the simulations and qualitative applications of the method. However, it may affect the description of very subtle quantitative effects such as the local conformational variations in B-DNA.

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Supporting Information Available: Studied geometries (xyz coordinates), tables of interaction energies, plots of energy gradients, and plots of DFT-SAPT energy contributions. This material is available free of charge via the Internet at http://pubs.acs.org.

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