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Structure and Stability of Molecular Crystals with Many Body Dispersion Inclusive Density Functional Tight Binding

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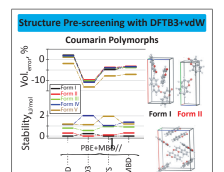
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Abstract

Accurate prediction of structure and stability of molecular crystals is crucial in materials science and requires reliable modeling of long-range dispersion interactions. Semi-empirical electronic structure methods are computationally more efficient than their *ab initio* counterparts, allowing structure sampling with significant speedups. Here, we combine the Tkatchenko-Scheffler van-der-Waals method (TS) and the many body dispersion method (MBD) with third-order density functional tight-binding (DFTB3) via a charge population-based method. We find an overall good performance for the X23 benchmark database of molecular crystals, despite an underestimation of crystal volume that can be traced to the DFTB parametrization. We achieve accurate lattice energy predictions with DFT+MBD energetics on top of vdW-inclusive DFTB3 structures, resulting in a speedup of up to 3000 times compared to a full DFT treatment. This suggests that vdW-inclusive DFTB3 can serve as a viable structural prescreening tool in crystal structure prediction.

Graphical TOC Entry



Introduction

Stability and structure prediction of molecular materials from first-principles electronic structure calculations bears significance to a wide range of problems ranging from pharmaceutical activity of drugs to optical properties of modern organic materials. 1-3 The rugged and complex energy landscapes of molecular crystals give rise to the phenomenon of polymorphism—the ability of molecules to form different crystalpacking motifs—which is a crucial aspect in drug design, food chemistry, and organic semiconductor materials. 4-7 Polymorphic materials exhibit many energetically close-lying minima, which can easily coexist and transform into each other at time scales that are inaccessible by conventional molecular dynamic simulations. Rigorous computational polymorph screening followed by correct stability ranking is therefore a crucial aspect for molecular crystal structure prediction (CSP).⁸⁻¹¹

In recent years, DFT methods have become more reliable in predicting and ranking polymorphic systems due to the incorporation of efficient dispersion correction methods that, 9,12-15 at the same time, ensure computational feasibility. 10,16-19 Especially, the inclusion of beyond-pairwise dispersion interactions through the Many-Body Dispersion (MBD) method coupled with semi-local DFT functionals has proven to be successful in this context. 10,15,20 To address larger length and time scales and more efficient structure prediction, several approximate electronic structure methods have been highly successful including semi-empirical quantum chemical methods such as AM1, PM7 or the DFT-based Density-Functional Tight Binding (DFTB). 21–23 DFTB has been significantly improved recently particularly in its description of charge polarization via third order charge fluctuation corrections (DFTB3)^{24,25} or its description of Hydrogen bonding. 26 Nevertheless, several shortcomings still persist that prohibit its use as standard tool in structure and stability prediction for molecular crystals. 22,27 The most detrimental shortcoming is the lack of long-range dispersion inherited from (semi)-local DFT with which

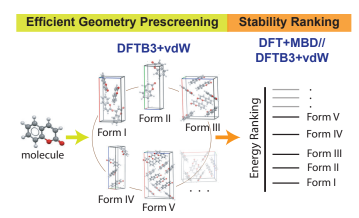


Figure 1: Crystal Structure Prediction: The prescreening step via efficient semi-empirical DFTB3+vdW method (left) prior to accurate stability ranking via DFT+vdW single-point energy on top of DFTB3+vdW geometries (right). Inset (middle) shows the molecular models of the five experimentally observed coumarin polymorphs.

DFTB models are parametrized, ²⁸ though early works augmented DFTB with an empirical correction potential. ²⁹

Many recent works have established pairwise dispersion corrections including the D3 and the dDMC methods parametrized for DFTB3. 30,31 Precalculated and tabulated C_6 (dipole-dipole) coefficients are used to calculate pairwiseadditive dispersion energies in these methods. The C_6 coefficients in D3 are environmentdependent via a fractional coordination number and thus do not directly depend on the electronic structure. 32 The Tkatchenko-Scheffler methods, e.g. TS and MBD, use a Hirshfeld partitioning of the electron density which provides a rescaling of free-atom reference dispersion parameters according to the local atomic environment. 15,33,34 All of those methods have been proven to be highly accurate in capturing long-range dispersion interactions for a variety of systems with their strengths in different types of materials. Stöhr et al. have recently proposed a method to replace the Hirshfeld density-partitioning scheme with a charge population analysis that directly correlates atomwise dispersion coefficients and a given Hamiltonian in local basis representation. ³⁵ This enables the incorporation of the TS and MBD methods into DFTB and other semi-empirical methods, where a real-space representation of the electron density is not directly available. Preliminary benchmarks of lattice and interaction energies have shown that these vdW-corrected DFT and DFTB models give comparable accuracy with the original TS/MBD implementation when based on predetermined molecular geometries for a broad range of systems. This suggests the potential application of DFTB+vdW methods in reliable and efficient structural prescreening of molecular CSP as sketched in Fig. 1.

Motivated by this finding and a recently developed implementation of analytical atomic forces in the MBD method, ^{36,37} in this work, we couple the state-of-the-art DFTB3 parameter set 3ob for organic molecules ^{25,27} with the TS and MBD methods for dispersion energy by calculating optimally-tuned range-separation parameters, enabling the standardized use of DFTB3(3ob)+TS/MBD. Thus, we present for the first time a modern semi-empirical Hamiltonian that includes vdW interactions to all atomic dipole orders based on anisotropic polarizabilities. We perform full geometry optimizations for the X23 benchmark database of organic crystals and demonstrate the large-scale applicability of the method on the example of the polymorphic molecular crystals such as coumarin and a flexible pharmaceutical molecule 2-((4-(3,4-Dichlorophenethyl)phenyl)amino)benzoic acid (C₂₁H₁₇Cl₂NO₂) from the 6th CSP blind test organized by the Cambridge Crystallographic Data Centre (CCDC). We find that vdW-inclusive DFTB3, in particular DFTB3+TS/MBD, are viable methods for an accurate description of molecular crystal structure, and identify challenges for the current 3ob parametrization of DFTB3.

The X23 benchmark dataset represents a mixture of molecular crystals dominated by Hydrogen, vdW, and combined Hydrogen-vdW bonding interactions. ^{38,39} We first analyze the quality of the dispersion-corrected DFTB3 geometries in terms of crystal volume and calculated root mean-squared deviation (RMSD) for the X23 dataset. The comparison was made with respect to the experimental crystal structures determined at the lowest possible temperature. ³⁹

Table 1: The mean absolute error (MAE), mean error (ME), and mean absolute relative error (MARE) in volumes and lattice energies of vdW-inclusive DFT(B3) methods on the X23 dataset with respect to the reference experimental values. Mean RMSD of the dataset with respect to experimental structures are also listed.

	Geometry			
-	Volumes			
method	$MAE(Å^3)$	ME(%)	MARE(%)	RMSD(Å)
PBE+D3	6.4	0.02	2.0	0.18
PBE+TS	8.3	-1.12	2.6	0.12
PBE+MBD	5.5	-0.16	1.8	0.12
DFTB3+D3	36.6	-11.23	11.2	0.26
DFTB3+TS	17.8	-5.52	5.6	0.17
DFTB3+MBD	14.8	-4.07	5.0	0.17
	Lattice Energies			
vdW method		$MAE (kJ mol^{-1})$	ME (%)	MARE (%)
	PBE+vdW//PBE+vdW			
D3		4.0	1.7	5.6
TS		14.0	16.2	17.5
MBD		4.8	3.2	6.4
	PBE+vdW//DFTB3+vdW			
D3		7.0	-7.4	8.6
TS		12.8	12.0	16.2
MBD		4.3	-1.1	5.5
•	PBE+MBD//DFTB3+vdW			
D3		6.7	-6.0	8.7
TS		4.8	-0.8	6.3
MBD		4.3	-1.1	5.5

The crystal volume evaluates the overall crystal lattice description, whereas the RMSD provides a more detailed evaluation of molecular orientation and alignment within the corresponding molecular crystal. The lattice energies were compared against experimental measurements (via back-corrected experimental sublimation enthalpies). 10,39 Revised experimental and/or high-level calculation data was used for benzene, 40 naphthalene, 41 and cytosine. 42 We also made sure the lowest energy conformer of gasphase (isolated molecule) was selected for calculations of 0 K lattice energies.³⁹ Extra care was taken to choose the lowest energy conformation of isolated molecules cytosine, ⁴³ oxalic acid, 44 succinic acid, 45 and urea 46 due to their conformational flexibility. A summary of statistics of the volumes, lattice energies, and RMSD of the X23 dataset is given in Table 1 with more details to be found in Figs. S1, S2 and Tables S1 and S2 in the Supporting Information (SI).

The three studied PBE+vdW methods are quite comparable in describing the crystal volumes with overall MARE values ranging from 1.8% to 2.6% with PBE+MBD yielding the best performance when compared to experi-

ment. PBE+D3 provides similar accuracy with a MARE of 2.0%. When combining the same vdW methods with DFTB3, we find that crystal volumes are systematically contracted compared to the PBE+vdW methods as can be seen from mean of volume errors in Table 1 (also can be seen in Fig. 2b). The resulting relative errors of 5.6% and 5.0% with respect to experiment for the DFTB3+TS and DFTB3+MBD methods dominantly originate from a strong underestimation of the volume of Ammonia and the three organic acid crystals in the X23 set (vide infra), whereas the DFTB3+D3 method yields a systematic volume underestimation across the dataset and an over fivefold increase in relative error of 11.2% compared to experiment. This observed volume contraction persists across the vdW-inclusive methods and partly appears to originate in the description of Pauli repulsion in the 3ob parametrization of DFTB. The 3ob parametrization has been developed to correct the known overbinding of covalent bonds in the MIO basis set, however intermolecular distances still seem to suffer from insufficient Pauli repulsion at longer distances. This issue is known to originate from the underlying basis confinement.²⁷

When combined with DFTB3, TS and MBD significantly outperform D3 in their description of crystal volume. One can attribute this twofold difference in relative error to the fitting of damping parameters of D3 in favor of energetics³⁰ rather than geometries as opposed to the optimally-tuned range-separation parameters based on energetics and geometries adopted for DFTB3+TS/MBD. For the sake of comparison, we revisited the D3 range-separation parameter of D3 for DFTB3+D3(3ob) based on a balanced description of energetics and geometries (i.e. S66x8, ^{47,48} similar to TS/MBD). We find a reduced volume MARE of 8.2%, which still corresponds to a larger underestimation of crystal volumes than found with DFTB3+TS/MBD.

A few systems in the X23 set, specifically CO₂ and Ammonia, are persistently described poorer than others regardless of the choice of dispersion method. PBE+vdW methods fail to give reasonable volumes for CO₂ and Am-

monia. The volume of the former is overestimated while the latter is underestimated by 8-10 %. ^{39,49} When moving from DFT to DFTB, this error becomes larger regardless of the employed method for the dispersion energy. In the case of the relatively strong H-bound Ammonia, it is indeed more relevant to compare the optimized structure with the cubic deuterated ammonia (ND₃) geometry at 2 K (with 128.6 $Å^3$ versus 135.1 $Å^3$ for Ammonia at 180 K), as isotope effects can be neglected at very low temperature. ¹⁰ Also, organic acid groups i.e. oxalic and succinic acids are still poorly described due to an insufficient description of charge polarization within the carboxyl groups in the 3ob parametrization. 22,27 These systems represent particular challenges for future parametrization work and further developments in charge polarization treatment within DFTB, whereas larger crystals are described consistently better with existing parametrizations.

Contrary to the modest description of crystal volume, the internal orientation and conformation of molecular crystals is described well by DFTB3+vdW methods as shown by RMSD errors in Fig 2a. TS/MBD yield geometries with substantially lower RMSD than D3 when compared to experimental crystal structures (0.1 Å for predominantly vdW-bound crystals and 0.2 Å for other systems). The enhanced treatment of geometries by DFTB3+TS/MBD methods combined with their significant speedup, compared to their DFT counterpart, can be advantageous in exhaustive structural search in material science. In contrast to the N^3 scaling of generalised gradient approximation (GGA) functionals like PBE, DFTB scales as $N \log(N)$ in sufficiently sparse systems.⁵⁰ Our performance analysis on selected X23 crystals shows that a speedup of up to a factor of 3000 can be achieved for DFTB3+TS compared to all-electron PBE+TS in FHI-aims (tight basis set), 51 whereas the performance gain for DFTB3+MBD is more moderate with a speed up of only 100 times. This essentially suggests that DFTB3+TS/MBD methods are optimally suited for the study of large complex molecular crystals, while smaller systems, such as the ammonia crystal, can be well treated by accurate

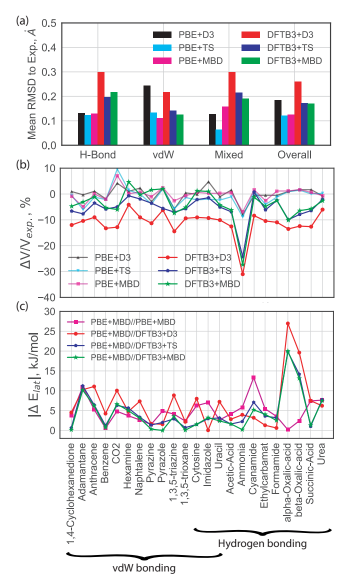


Figure 2: The X23 benchmark dataset: (a) Mean of RMSD values as calculated with respect to experimental geometries, (b) relative error of volumes with respect to experimental geometries, (c) absolute error of lattice energies (in kJ mol^{-1}), both with respect to the experimental values. For comparison of PBE+vdW//DFTB3+vdW combination with respect to the PBE+vdW energies see Fig. S3. The abbreviations used in the legends is described in the footnote.

DFT+vdW methods.

Turning to the description of lattice energies and crystal stability for the X23 dataset, we find PBE+vdW methods to yield relative errors of 5.6%, 6.4%, and 17.5% for PBE+D3, PBE+MBD, and PBE+TS, respectively (see Table 1 and Fig. S2). The comparably large error for PBE+TS compared to PBE+D3 might

seem surprising, considering that both methods describe the dispersion energy using a pairwise approximation. However, range separation or damping parameters in vdW methods can be chosen such that interaction energies for a broad range of systems are optimized.³² This, can lead to incorporation of contributions, which at this pairwise level of treatment should not be included, and could potentially negatively affect transferability across different systems. TS van der Waals functional, on the other hand is based on free atom reference data with functional-specific range-separation tuned to exclusively represent interaction energies of small intermolecular complexes (S22), which minimizes effects beyond pairwise contributions. ³⁹

When replacing PBE+vdW with DFTB3+vdW in the description of lattice energies, regardless of vdW method, we find lattice energies with minimum MARE as high as 15% corresponding to minimum MAE of 13 kJ mol⁻¹ compared to experiment (see second column in Fig. S2a of SI). While this may disqualify the DFTB3+vdW methods in their current formulation as outright stability prediction methods, following the scheme of Fig. 1, we can use the higher quality of crystal structure prediction at the DFTB3+vdW level in order to perform structural prescreening. Thereby, we identify stable structures using DFTB3+vdW and evaluate improved energetics at the DFT+vdW The evaluation of lattice energies at the DFT level i.e. PBE+MBD on top of DFTB3+MBD structures improves the lattice energy prediction significantly as shown in Table 1 and Fig. 2c. In fact, the relative error of PBE+MBD//DFTB3+TS/MBD energies is comparable in performance with the full DFT+MBD i.e. PBE+MBD//PBE+MBD when compared to experiment. This means that replacing optimized PBE+MBD crystal structures with DFTB3+TS/MBD structures does not significantly affect the lattice energy prediction compared to experiment.

Encouraged by these results, we proceed to

^{*}We use the following abbreviations: 'level of theory for energy evaluation'//'level of theory for geometry optimization'

study two highly polymorphic systems. First we focus on coumarin with five experimentally observed polymorphs (see the inset of Fig. 1), which have been recently studied by vdW-inclusive DFT. ¹¹ The structures of all coumarin polymorphs have been refined at room temperature, and low-temperature structures (90 K) are available for polymorphs I, III, IV. ¹¹

Figures 3a and b show the RMSD results for crystal structures calculated with respect to experiment at 300 and 90 K, respectively. PBE+vdW methods yield equally good performance with a mean RMSD of just below 0.2 Å (room temperature) and 0.1 Å at 90 K. DFTB3+TS/MBD yield geometries as good as PBE+vdW with RMSD of just above 0.1 Å at 90 K. Figure 3c shows the optimized unit cell volumes of coumarin polymorphs with respect to experimental structures. PBE+vdW methods slightly overestimate the crystal volume, whereas DFTB3+D3 strongly underestimates by more then 10% compared to experiment. DFTB3+TS/MBD methods provide an average relative volume error with respect to the experiment of 4.8%, and 4.2%, respectively. The larger relative volume error of polymorph V, compared to other polymorphs, is due to comparison with an experimental structure measured at 300 K. This example highlights the significant impact of thermal expansion, which is up to 4 % between 300 K and 90 K.¹¹

We compare the stability rankings of coumarin polymorphs based on lattice energies as shown in Fig. 3d. Fig. S6a shows the stability rankings based on DFTB3+vdW energies. It can be seen that generally they are not sufficient for accurate energy determination of polymorphic systems, although DFTB3+MBD//DFTB3+MBD capture the first two polymorphs correctly. As established herein we calculate PBE+MBD energies on top of DFTB3+vdW geometries. Experiment guides the stability rankings of coumarin forms as: Form I < Form II < Form IV < Form V, with the Form I being the most stable phase. It is quite encouraging that the full PBE+MBD and PBE+MBD//DFTB3+MBD both yield the correct energy ranking for the first three polymorphs, considering the narrow

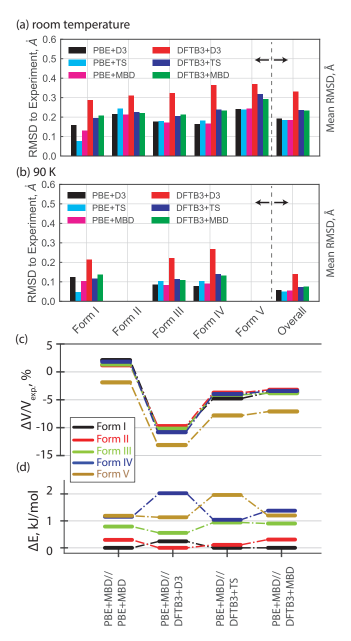


Figure 3: Comparison of DFT(B3)+vdW methods for coumarin polymorphs in terms of: (a) RMSD with respect to experimental structures at room temperature, (b) RMSD with respect to experimental structures 90 K (complete experimental data were not available for Form II and V) , (c) optimized unit cell volumes $\Delta \text{V/V}_{exp}$ in% in which 90 K experimental structures were used for comparison (expect for polymorph V with structure at room temperature), (d) stability rankings based on lattice energies ΔE in kJ mol^{-1} .

energy range within which the five polymorphs are ranked. Notice that the geometry used for final polymorph stability ranking has an impact of up to 1 kJ mol⁻¹ per molecule (see Figure 3d), which is large enough to qualita-

tively affect the polymorphic energy landscape.

We further extend the applicability of the methods by studying the C₂₁H₁₇Cl₂NO₂ molecule from the 6th CCDC CSP blind test (molecule XXIII)—a former drug candidate molecule with five known crystalline polymorphs (see the optimized structures in Fig. S4). The preliminary results for XXIII's five known stable structures are shown in Fig. S5. While all methods systematically underestimate the volumes, we show that the structures of polymorphs are best described by DFTB3+MBD with MARE of 4% w.r.t. experimental geometries close to full PBE+MBD calculations with MARE of 1.9% (see Fig. S5a). The stability rankings of XXIII polymorphs are shown in Fig. S5b (see Fig. S6b for the evaluation of stability rankings based on DFT(B3)+vdW energies). We notice that the polymorphic energy landscape for flexible pharmaceutical molecular crystals is strongly dependent on the optimized geometry. Using less accurate DFTB3+D3 geometries, i.e. large volume underestimations (see Fig. S5a), for the final energy ranking can lead to changes of 10 kJ mol⁻¹ per molecule in the polymorphic energy differences (see Figure S5b). We observe that PBE+MBD//DFTB3+MBD ranks the first two polymorphs equal to the full PBE+MBD This is an encouraging result, calculation. which suggests that crystal structures obtained with DFTB3+vdW are more accurate when vdW interactions play a prominent role in intermolecular interactions as compared to hydrogen bonds. Since vdW interactions become most prominent in the case of flexible molecules, we anticipate vdW-inclusive DFTB to become a valuable structural prescreening tool for molecular crystals, supramolecular complexes, and systems of biological interest.

In summary, we coupled pairwise dispersion corrections and the many-body dispersion method with DFTB3 using charge population analysis and optimally-tuned range-separation parameters for the current state-of-the-art 3ob parameter set for organic molecules. We examined the applicability of this approach for organic crystals using the X23 benchmark set of molecular crystals and two highly polymor-

phic systems, finding encouraging results. The proposed method yields significantly improved geometries compared to bare DFTB, whereas energetics can still be improved.⁵² We suggest to improve the lattice energy prediction by calculating single-point PBE+MBD energies on top of DFTB3+vdW geometries, which were found to be very close to the full DFT calculations. We identified remaining issues in the DFTB description potentially stemming from the parametrization of the 3ob parameter set. As more suitable DFTB parametrizations become available, the here presented approach will become even more effective for complex molecular materials. Further studies are necessary to confirm the transferability of our results to other systems, such as carbon nanostructures, larger flexible molecules, and hybrid organic-inorganic materials. Additionally, applications beyond structure search, such as the calculation of thermal corrections and phonon spectra⁵³ are an important field with the urgent need of efficient electronic structure methods such as the ones presented here.

Computational Details

We have interfaced previously published modules for the TS³³ and MBD³⁴ methods with the latest development version of the DFTB+ code. 50 DFTB3+D3/TS/MBD calculations were performed using DFTB+ and the 3ob parametrization.²⁷ We have optimized specific damping parameters for the TS/MBD methods using S66x8 dissociation curves ^{47,48} balancing the accuracy of intermolecular geometries and energies at the same time (see Fig. S6 for the fitting procedure). The optimized damping parameters are 1.05 (the s_R parameter) for TS, 1.0 (the β parameter) for MBD. All geometry optimizations were done using the FIRE algorithm ⁵⁴ in the Atomic Simulation Environment . 55,56 Root mean-squared deviation was calculated by constructing a 15-molecule supercell followed by a calculation of the RMSD15 value as implemented in the Mercury package. ⁵⁷ For RMSD calculations, only heavy atom positions were considered (Hydrogen positions were ignored). DFT calculations were performed using the FHI-aims code 51 with PBE functional 58 together with D3/TS/MBD dispersion interactions. For all DFT calculations, the light basis set in FHI-aims was used for optimization and energies were obtained with the converged tight basis using the optimized structures. The automatic k-points mesh for sampling the Brillouin zone was selected such that $n_i \times a_i = 30 \text{ Å}$, where n_i is the k-points sampling for

the corresponding a_i lattice parameter.

Supporting Information

The Supporting Information contains additional supporting data and detailed tabulations of calculated lattice volumes and energies as well as details concerning the fitting procedure for the range-separation parameters and further computational details.

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