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Metal-Organic Frameworks Properties from Hybrid Density Functional Approximations

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ABSTRACT

The chemical versatility and modular nature of Metal-Organic Frameworks (MOFs) make them unique hybrid inorganic-organic materials for several important applications. From a computational point of view, ab initio modeling of MOFs is a challenging and demanding task, in particular when the system reaches the size of gigantic MOFs as MIL-100 and MIL-101 with several thousands atoms in the unit cell. Here we show how such complex systems can be successfully tackled by a recently proposed class of composite electronic structure methods revised for solid-state calculations. These methods rely on HF/DFT hybrid functionals (i.e. PBEsol0 and HSEsol) combined with a double-zeta quality basis set. They are augmented with semi-classical corrections to take into account dispersive interactions (D3 scheme) and the basis set superposition error (gCP). The resulting methodologies, dubbed "sol-3c", are cost-effective, yet reach the hybrid functional accuracy. Here, sol-3c methods are effectively applied to predict the structure, vibrational, electronic, and adsorption properties of some of the most common MOFs. Calculations are feasible even on very large MOFs containing more than 2500 atoms in the unit cell as MIL-100 and MIL-101 with reasonable computing resources. We propose to use our composite methods for the routine In Silico screening of MOFs targeting properties beyond plain structural features.

I. INTRODUCTION

Metal Organic Frameworks^{1,2} (MOFs) represent a wide class of hybrid organic-inorganic materials composed by an inorganic clusters (also known as metal knots) interconnected by polytopic organic linkers that self-assembly in a threedimensional, crystalline and microporous structure. During the last two decades these materials have gained considerable attention in industry and academic world thanks to their modular nature, high chemical versatility and tunability. All these features make MOFs promising materials for vast variety of applications ranging from gas storage³ and separation⁴, to (photo)catalysis⁵, mechanical and chemical sensing⁶, and to opto(electronics)^{7,8} and drug delivery^{9,10}. Another promising application area is to use the porous complexes to enable X-ray analysis of molecules that do not natively crystallize or where just few nanograms are available.¹¹ These crystalline sponges for structure elucidation are currently developed into a commercially available technology.¹²

From a computational point of view, theoretical modelling of MOFs is not a straightforward task due to the presence of several levels of complexity that can be summarized as:

(i) The number of synthesized and hypothetical structures is continuously increasing. For instance, more than 70 000 structures of MOFs have been recently identified in the Cambridge Structure Database^{13,14} and the CoReMOF dataset of computation-ready metal-organic frameworks contains more than 100 000 structures¹⁵.

- (ii) The size of the system in terms of number of atoms and unit cell volume, varies remarkably from small MOFs containing less then 40 atoms in the unit cell (e.g. MIL-53¹⁶) to giant MOFs with more than 2000 atoms per unit cell (e.g. MIL-100¹⁷, MIL-101¹⁰, bio-MOFs^{18,19}).
- (iii) The intrinsic complexity of MOFs due to their modular nature with the combination of different Secondary Building Units (SBU) that leads to a large variety of framework topologies, but also to frameworks modifications (e.g interpenetration), host-guest interactions, flexibility and defects.

This scenario represents a challenge for computational studies of MOFs, but also the prediction of their physical and chemical properties is not trivial and requires a proper choice of the adopted level of theory²⁰. Therefore, to tackle the structural and chemical complexity of MOFs, accurate yet cost-effective computational methods are needed.

In the last years, a family of composite electronic structure methods has been proposed.^{21–25} They range from Hartree-Fock to pure and hybrid exchange correlation functionals, which are combined with small to medium Gaussian basis set and augmented with semi-classical corrections to describe London dispersion interactions at long range through the D3²⁶ or D4^{27,28} scheme and remove the basis set superposition error (BSSE) via the gCP^{29,30} method. These composite methods dubbed "3c" have been targeted to molecules and organic crystals³¹ and they have been shown to provide very good results on geometric and thermochemical properties³²

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but they are only partly applicable and transferable to solidstate calculations. Therefore, we recently proposed a revised version of such methods specifically designed for solid state calculations³³. In particular, we adopted exchange-correlation functionals for solids hybridized with 25% of HF exchange and reshaped the molecular basis set to make them suitable for ionic, covalent and molecular solids, inorganic compounds and hybrid materials, such as MOFs. The new revised methods have been renamed "sol-3c"³³ to emphasize a broader range of applicability, including solids. They have also been proved to efficiently tackle large systems of up to thousands of atoms with excellent performance, even with small-scale computing resources.³⁴

In this work, we show how "sol-3c" methods can be successfully applied to model various properties of MOFs, e.g. equilibrium structures, vibrational frequencies and simulated IR/Raman spectra, fundamental band gap, phase stability and adsorption of small molecules. Furthermore, we show that hybrid HF/DFT composite methods can be effectively employed to simulate very large MOFs such as the so-called giant-MOFs (i.e. MIL-100¹⁷ and MIL-101³⁵).

II. METHODS

A. Composite Hybrid HF/DFT

In this section we briefly recall the main ingredients of the composite hybrid HF/DFT methods which are examined in the present work. The total energy provided can be written as^{32}

$$E_{\text{tot}}^{\text{sol-3c}} = E_{\text{tot}}^{\text{DFT/basis}} + E_{\text{disp}}^{\text{D3}} + E_{\text{BSSE}}^{\text{gCP}}$$
(1)

 $E_{\text{tot}}^{\text{DFT/basis}}$ denotes the total energy evaluated with the hybrid XC functional. In the present work, the theoretical methods are the global hybrid PBEsol0 functional³⁶ and the range-separated hybrid HSEsol functional³⁷. The two methods were chosen because they were specifically devised for solids and they use 25% of exact exchange in the hybrid exchange functional, which has been shown to be a good compromise.³⁸ The basis is the revised def2-mSVP for PBEsol0-3c and HSEsol-3c.²² The $E^{\text{DFT/basis}}$ total energy is supplemented with a damped atom-atom two-body dispersion energy as defined in the D3 approach²⁶:

$$E_{\rm disp}^{\rm D3(BJ)} = -\frac{1}{2} \sum_{AB} \sum_{n=6,8} s_n \frac{C_n^{AB}}{R_{AB}^n} f_{\rm damp}^{(n)}(R_{AB})$$
(2)

Here, C_n^{AB} denotes the *nth*-order dispersion coefficient (orders = 6, 8) for each atom pair AB, R^{AB} is their inter-nuclear distances and s_n are the order-dependent scaling factors. The rational Becke-Johnson damping function³⁹ has become the default in combination with D3⁴⁰:

$$f_{\rm damp}^{(n)}(R_{AB}) = \frac{R_{AB}^n}{R_{AB}^n + (a_1 R_0^{AB} + a_2)^n}$$
(3)

The damping function incorporates radii for atomic pairs $R_0^{AB} = \sqrt{C_8^{AB}/C_6^{AB}}$ and functional-specific parameters a_1 and a_2 that have been refitted in the present work for the different composite methods. In addition, the Axilrod–Teller–Muto^{41,42} three-body dipole-dipole-dipole term, hereafter denoted as (ABC), is also included.

The removal of the basis set superposition error (BSSE) due to the use of small basis sets is accomplished through the geometrical counterpoise correction²⁹:

$$E_{\text{BSSE}}^{\text{gCP}} = \frac{\sigma}{2} \sum_{AB} V_A^{\text{gCP}}(R_{AB}) f_{\text{damp}}^{\text{gCP}}(R_{AB})$$
(4)

The difference in atomic energy between a large (nearly complete) basis set and the target basis set for each free atom is used as a measure to generate the repulsive potential $V_A^{\rm gCP}$ with fitting parameters α , β , η .

All functional and basis set specific parameters have been optimized for each revised composite method as discussed in Ref. 33.

B. Computational details

All the calculations have been performed with a development version of the CRYSTAL17^{43,44} code, in its replicated data parallel version. For the evaluation of the DFT exchangecorrelation term an XLGRID (75,974) pruned grid was used. Default convergence criterion for both energy and geometry optimization were employed. The tolerances for one- and two-electron integrals calculation were set to 10^{-7} , 10^{-7} for the Coulomb and to 10^{-7} , 10^{-7} , 10^{-25} for exchange series, respectively. The shrinking factors for the diagonalization of the Hamiltonian matrix in the reciprocal space were set to 2 for the Monkhorst-Pack net and to 2 for the Gilat net respectively, except for MIL-100 and MIL-101 where the Fock matrix in the reciprocal space was diagonalized at the central point of the first Brillouin zone (i.e. Γ point). All the properties discussed in this work have been evaluated on the fully relaxed MOFs structures (i.e. lattice parameters and atomic positions). Geometry optimization was performed with the constraints imposed by the symmetry of the system.

III. RESULTS AND DISCUSSION

In the following, along with the results obtained with the hybrid HF/DFT composite methods, we also report data computed with the HFsol-3c method³³, for comparison. It combines HF with a minimal basis set along with the two semiempirical terms mentioned above and a third correction for short-ranged basis incompleteness effects (see Ref.³² for further details). As an additional comparison, present results for sol-3c methods are benchmarked against the more expensive B3LYP-D3(ABC)/pob-TZVP-rev2⁴⁵ level of theory.

A. Equilibrium geometries

For any simulation study, obtaining reasonable structures is a mandatory starting point for further calculation such as adsorption, electronical, vibrational, and mechanical properties. This is particularly true for MOFs because of the variability of organic and inorganic building units.

Lattice parameters and bond lengths have been computed with sol-3c composite methods for a set of 15 well known MOFs (hereafter referred to as the MOF-LC15 dataset) that differ for metal ion, organic linker, topology and SBUs. Some of the crystal structures of the MOFs under examination are shown in Figure 1 (see Figures S2-S8). Results are analyzed graphically in Figure 2, which illustrates the relative percentage error of the predicted unit cell volume with respect to experimental reference data. The latter have been collected from low-temperature X-ray diffraction measurements. In addition, B3LYP-D3(ABC)/pob-TZVP-rev2 results are also shown for comparison. Detailed computed results are reported Tables S1-S6.

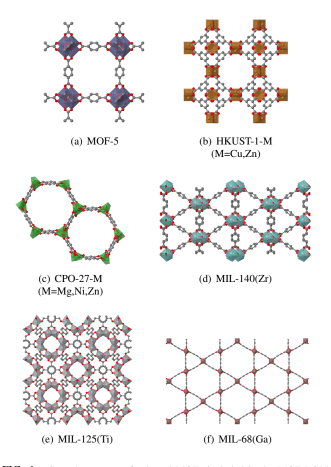


FIG. 1. Crystal structure of selected MOFs included in the MOF-LC15 dataset. The inorganic building units are represented by translucent coordination polyhedra of different colors. Color code for atoms: O in red, C in grey, Hydrogens are omitted for clarity.

PBEsol0-3c and HSEsol-3c give results similar to each other thus indicating that the screening of the HF exchange does not impact the property under consideration. They are both in excellent accord with B3LYP-D3(ABC)/pob-TZVPrev2 results with an overall good agreement with the experimental values. Hybrid sol-3c composite methods show remarkably small relative error of $\pm 2\%$ expect for MIL-53(Al) in the narrow pore (NP) phase for which a significant underestimation of the unit cell volume is observed. This is related to the peculiar structure of MIL-53(Al) NP (vide infra) in which aromatic rings of the linkers are in close contact and dispersive contributions play a relevant role thus suggesting that the D3 correction could be partly overestimated. HFsol-3c tends to underestimate the volume of conventional cell of CPO-27- $(M)^{46}$ (M=Zn, Mg, Ni), HKUST-1(M)^{47,48} (M=Zn, Cu), MIL-53(Al)¹⁶ (NP, LP), MIL-125(Ti)⁴⁹ and UiO-66⁵⁰ dehydroxylated, while for the other systems it gives lattice parameters in reasonable agreement with the experimental data. However, it overestimates the lattice volume of MOF-5⁵¹ and MIL- $47(V)^{52}$ with a Relative Error% (RE%) greater than 4% and 8%, respectively. This large discrepancy is probably caused by the combination of the HF method with a minimal basis set.

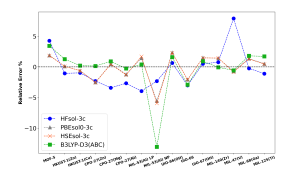


FIG. 2. Relative Error % (RE%) from experimental unit cell volumes computed at HFsol-3c, PBEsol0-3c, HSEsol-3c and B3LYP-D3(ABC)/pob-TZVP-rev2 for a dataset containing 15 well-known MOFs.

Another important structural feature for MOFs is the metal-ligand bond length that characterizes the link between the inorganic and the organic building blocks, This is particulary relevant to properly describe the structure of hybrid organic-inorganic materials as MOFs and usually quite hard to be predicted even by well-tuned force fields^{53–57}. Results are gathered in Table S7-S11 of the Supporting Information. Overall, metal-ligand bonds computed with PBEsol0-3c and HSEsol-3c are in good agreement with B3LYP-D3(ABC)/pob-TZVP-rev2 data used as reference. Instead, HFsol-3c systematically underestimates the metal-linker distance for all the systems except for MIL-47(V), where a lengthening is observed. Again, this could be related to the use of a minimal basis set in the composite method.

Overall, sol-3c HF/DFT composite methods predict structures of similar quality as dispersion-corrected B3LYP with triple-zeta basis sets, or even better, but at a lower cost.

B. Vibrational frequencies and simulated IR and Raman spectra

As an example of the ability of sol-3c methods to provide high quality vibrational frequencies and to simulate IR and Raman spectra, we selected the well-known UiO-66 Zirconium-MOF for which accurate experimental measurements are available.^{50,58}

Its framework is built of $Zr_6O_4(OH)_4$ units linked by 12fold connection through 1,4-benzene-dicarboxylate (BDC) ligands. Vibrational frequencies were computed at the Γ point on the fully optimized structure. Infrared and Raman intensity were calculated via a linear response CPHF/KS approach as detailed in Ref.^{59,60} A Lorentzian broadening of 5 cm⁻¹ was adopted for both the spectra. As common practice^{61,62} computed vibrational frequencies were scaled for a better comparison with experimental spectra. Here, a factor of 0.95 was used. The spectra were simulated at the PBEsol0-3c level of theory.

As shown in Figure 3(a) and 3(b) for IR and Raman spectra, respectively, the overall agreement between experimental and theoretical results is excellent in both cases. This result offer an example of how composite methods can be used in supporting the experimental evidence and interpreting measured spectra. Indeed, several spectral regions may be highlighted and assigned to vibrational modes, concisely: i) the first one around 3839⁻¹ related to the OH stretching of isolated hydroxyls of the inorganic cluster; ii) the second one around 3200⁻¹ where CH stretching of benzene-dicarboxyl moiety are observed; iii) the third one between 1700 and 1200 ⁻¹ characterized by bands due to asymmetric and symmetric modes of carboxylate; iv) the fourth is the region 1200-600 ⁻¹ characterized by in-plane and out-of-plane deformation modes of aromatic rings and C-H groups; v) the fifth region (600-200 $^{-1}$) characterized by modes due to inorganic cluster deformation as the Zr-O symmetric and asymmetric vibrations and μ_3 -OH stretching; vi) a last interval concerns the mode below 200⁻¹, where collective vibrations and lattice modes are present. Recently, the good performance of the PBEsol0-3c method in the prediction of vibrational frequencies and IR/Raman spectra has been also confirmed in the study of encapsulated molecules in HKUST-1(Cu)⁶³ and ZIF-8⁶⁴.

C. Fundamental band gap and dielectric constant

MOFs have also emerged as interesting materials for their electronic and dielectric properties as low band gap materials with very small dielectric constant (k) with promising applications as optical and optoelectronic materials.^{7,20,65–69} Therefore, an accurate prediction of the band gap is important for a precise estimation of the dielectric response⁸.

For this properties, HFsol-3c is not of interest because of the well-known overestimation of the band gap of the HF method, while the hybrid sol-3c composite methods have

(i) UiO-66(Zr) IR spectrum

10

(b) UiO-66(Zr) Raman spectrum

enumber (cm-1)

1000

1400

1600

600

FIG. 3. (a) Comparison between experimental and simulated (a) Infrared and (b) Raman spectra of hydroxylated UiO-66(Zr) as computed with the PBEsolO-3c composite method. Simulated Raman spectrum refers to 298.15 K with an incoming laser frequency of 785 nm. The unit cell of hydroxylated UiO-66(Zr) is also shown as inset.

been designed with a certain percentage of exact exchange that makes them guite effective in the description of the electronic structure of solids. Figure 4 shows the fundamental band gap calculated with the hybrid HF/DFT composite methods for a series of Isoreticular Metal-Organic Frameworks (IRMOF) and other selected MOFs for which experimental data are available^{70–73}. PBEsol0-3c tends to systematically overestimate the band gap for all of the systems considered as well as B3LYP-D3(ABC)/pob-TZVP-rev2. The only exception is represented by HSEsol-3c, which thanks to its screened Coulomb exchange potential gives results in very good agreement with the experimental values, in particular for UiO-66-OH-NH₂ and MIL-125-NH₂. HSEsol-3c seems to be the most promising method for describing fundamental band gap of MOFs with a Mean Absolute Relative Error % (MARE%) of 8.8% followed by B3LYP-D3(ABC) and PBEsol0-3c with a MARE% of 14.1% and 27.1% respectively (see Table S12). For PBEsol0-3c, we also computed the unit cell polarizability and the corresponding dielectric constant at the static limit. Results are reported in the supporting information (see Table S12). As discussed by some of us⁸, the polarizability of the different frameworks largely depends on the type of ligand and topology, whereas the dielectric constant that inversely depends on the unit cell volume becomes very small and covers a range of 1.2-2.0. Present results confirm that trend with the values of k being close to

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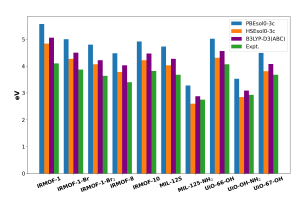


FIG. 4. Comparison between experimental value taken from Ref.^{70–73} and the computed band gaps by using different level of theory.

the one computed with B3LYP-D3(ABC)/pob-TZVP-rev2.

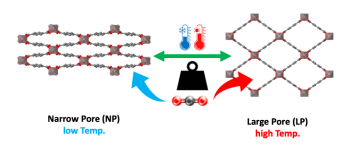


FIG. 5. Reversible switch of the MIL-53 framework between narrow-pore phase (NP) and large-pore phase (LP). This effect is mediated by external stimuli such as pressure, temperature and gas adsorption.

D. The MIL-53(AI) bistability

MOFs are stimuli-responsive materials⁷⁴. One of the most striking evidences of this property is the breathing behavior of the isoreticular family of MIL-53⁷⁵ and other MOFs as MIL-68 (Ga) and MIL-47 $(V)^{52,76}$. Indeed, depending on the temperature, pressure or interaction with adsorbed gases the framework undergoes a reversible phase transition between a narrow pore (NP) structure and a large pore (LP) phase. This peculiar behaviour can be very important for selective adsorption processes^{77–80} or as molecular sieve but also for tuneable dielectrics⁸¹.

Here, we focus on the temperature-driven phase bistability of MIL-53(Al)^{16,82} between the low temperature monoclinic NP structure and the high temperature orthorhombic LP one. Concerning structural parameters for both systems (see Table I for unit cell volumes and Table S3 for lattice parameters), overall results obtained with composite sol-3c methods are in reasonable agreement with the experimental values measured at 77K by neutron diffraction technique. As mentioned before, for MIL-53(Al)¹⁶ LP, HFsol-3c yields slightly underestimated lattice parameters and volume, while the composite HF/DFT methods provide results in good agreement with the experimental data. For MIL-53(Al)⁷⁵ NP, all composite methods provide a volume that is underestimated with respect to the experimental one.

The relative stability of the two phases of MIL-53(Al) for all of the examined methods are also reported in Table I). As expected, MIL-53(Al) NP is the most stable structure at all levels of theory.

TABLE I. Unit cell volumes (in $Å^3$) of the LP and NP structures of MIL-53(Al) and electronic energy difference between the two phases (in kJ/mol).

Method	Vol. LP	Vol. NP	ΔE_{lp-np}
HFsol-3c	1363.1	843.9	3.9
PBEsol0-3c	1439.7	815.9	13.4
HSEsol-3c	1443.3	812.6	11.7
B3LYP-D3(ABC)/VTZP ⁸¹	1436.5	802.2	19.4
RPA+EC ⁸³	1455	860	7.4
Exp. (T=77K) ¹⁶	1419.2	863.9	-

Hybrid composite methods, in particular HSEsol-3c, provide a relative stability of the two phases close to the recently reported RPA+EC value of 7.4 kJ/mol^{75,83}, while HFsol-3c tends to underestimate the relative stability of the two phases.

E. Gas adsorption

MOFs represent very promising materials for gas separation and storage thanks to the combination of high surface area with coordinatively unsaturated metal sites, which can act as Lewis acid and directly interact with the adsorbate. In this section we discuss the adsorption of two small molecules as CO and CO₂ on CPO-27-M (M=Mg, Zn) and HKUST-1(Cu).(see Figures S9 and S10) The calculated interaction energies are compared with the experimental values obtained from calorimetric measurements at room temperature. It is worth noting that the former should be corrected to include the zero-point vibrational energy and a thermal correction to enthalpy at the given temperature. This contribution has been estimated to be small and to amount around 3.5-5.0 kJ/mol for CPO-27(M)⁸⁴ at room temperature, therefore it has not been explicitly taken into account in the present work.

Adsorption of CO_2 causes a decrease of both lattice parameters and unit cell volume for all the systems herein discussed with respect to pristine material (see Tables S13, S15 and S17), The compression is greater for CPO-27(Zn) rather than HKUST-1(Cu) and CPO-27(Mg).

CO₂ adsorption energies calculated with composite methods are shown in Figure 6 in comparison with experimental data used as reference and B3LYP-D3(ABC)/pob-TZVP-rev2 results. HFsol-3c strongly overestimates the interaction energy while both PBEsol0-3c and HSEsol-3c give similar results in good agreement with experiment. Interestingly, the accord would be even better when including the estimated ZPE and thermal correction to enthalpy at room temperature.

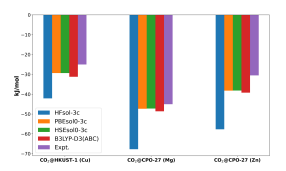
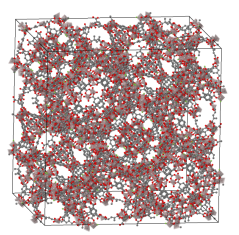


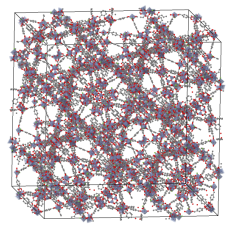
FIG. 6. Adsorption energy for CO_2 in HKUST-1(Cu) and CPO-27-M (M=Mg,Zn), comparison between calculated electronic energy at different level of theory and experimental heat of adsorption.

As regards the adsorption of CO, the host-guest interaction causes again a contraction of the cell with respect to the pristine material (see Tables S14, S16 and S18). The trend is similar to the adsorption of CO_2 with the decrease of the lattice parameters and unit cell volume being greater for CPO-27(Zn) and smaller for HKUST-1(Cu), while for CPO-27(Mg) there is no sizeable modification of the structure. At variance with CO₂, both HF and hybrid composite methods give a largely overestimated interaction energy between CO and the metal centre for all examined MOFs as clearly shown in Figure S11. In this case, B3LYP-D3(ABC)/pob-TZVP-rev2 results nicely agrees with experiment thus suggesting some not well balanced basis set dependence for the composite methods. Preliminary results indicates that the proper description of the interaction of CO with the adsorption site requires a more flexible basis set for carbon. Work is in progress to further check it.

We also analyzed the interatomic distance and the angle between the metal center and the adsorbed molecules (see Tables S19-S24). For CO, HFsol-3c provides both distances and angles in agreement with B3LYP-D3(ABC)/pob-TZVPrev2 data, while PBEsol0-3c and HSEsol-3c underestimate the M—CO distance. Concerning CO₂, composite HF/DFT methods provides bond length in good agreement with B3LYP-D3(ABC)/pob-TZVP-rev2 data, while HFsol-3c underestimates the interaction distance. In this case, CO₂ forms a bent complex with MOFs that has been explained in terms of the electrostatic interaction between O atoms of the carboxylic groups around the metal and the quadrupole moment of CO_2^{84} .



(a) MIL-100-M (M=Al,Sc,Cr,Fe)



(b) MIL-101-M (M=Al,Cr)

FIG. 7. Crystallographic unit cell of (a) MIL-100-M (M=Al,Sc,Cr,Fe) and (b) MIL-101-M (M=Al,Cr). The inorganic building units are represented by translucent coordination polyhedra of different colors. Color code for atoms: O in red, C in grey, Hydrogens are omitted for clarity.

F. Pushing the limit of sol-3c methods: Giant MOFs

In the introduction we mentioned the very large variability in the number of atoms in the unit cell of MOFs that can make the quantum-mechanical modeling very challenging. So far, we have reported and discussed results for small-to-medium MOFs with tenths-to-hundredths atoms in the unit cell. In the following, we show how "sol-3c" methods can be effectively applied to large and very large MOFs with thousands of atoms in the unit cell. In particular, we focused on the so-called giant-MOFs, namely: MIL-100 and MIL-101 (see Figure 7). They are comprised of oxo-centered trinuclear units (denoted as trimeric units or TU) of trivalent metals, namely Al and Cr, connected with 1,3,5 BTC (benzene-1,3,5-tricarboxylic acid) and 1,4 BDC (benzene-1,4-dicarboxylic acid), respectively. Starting from the experimental cubic crystalline structure of MIL-100 and MIL-101 we removed all water molecules and positioned a counterion on one of the metal of each TU. For sake of simplicity, we decided to model MIL-100 and MIL-

TABLE II. Predicted lattice parameters and unit cell volumes of MIL-100 (M^{III}) optimized at HFsol-3c, PBEsol0-3c and HSEsol-3c level of theory in comparison with B3LYP-D*/TZVP⁸⁵ results and experimental data^{86–89} along with percentage deviation of unit cell volume ($\Delta V\%$)

		1		0 1 0		· · ·
Al ^{III}	a(Å)	c(Å)	c/a	volume(Å ³)	mean LP(Å)	ΔV%
HFsol-3c	70.701	70.563	0.9980	352723.1	70.655	-4.26
PBEsol0-3c	71.484	71.284	0.9972	364258.7	71.417	-1.12
HSEsol-3c	71.462	71.262	0.9972	363922.8	71.395	-1.22
B3LYP-D*/TZVP ⁸⁵	71.531	71.494	0.9995	365813.0	71.519	-0.70
Exp. ⁸⁶ Sc ^{III}	71.678	-	-	368401.0	-	-
	a(Å)	c(Å)	c/a	volume(Å ³)	mean LP(Å)	$\Delta V\%$
HFsol-3c	73.990	74.051	1.001	405390.7	74.010	-1.37
PBEsol0-3c	74.283	74.319	1.000	410090.8	74.295	-0.22
HSEsol-3c	74.270	74.280	1.000	409732.7	74.273	-0.31
B3LYP-D*/TZVP ⁸⁵	74.676	74.823	1.002	417248.2	74.725	1.52
Exp. ⁸⁷	74.350	-	-	411001.0	-	-
Cr ^{III}	a(Å)	c(Å)	c/a	volume(Å ³)	mean LP(Å)	$\Delta V\%$
HFsol-3c	72.410	72.452	1.001	379876.3	72.424	-1.97
PBEsol0-3c	72.470	72.497	1.000	380749.0	72.479	-1.75
HSEsol-3c	72.456	72.479	1.000	380503.3	72.464	-1.81
B3LYP-D*/TZVP ⁸⁵	72.602	73.249	1.009	386097.1	72.818	-0.36
Exp. ⁸⁸ Fe ^{III}	72.906	-	-	387511.4	-	-
Fe ^{III}	a(Å)	c(Å)	c/a	volume(Å ³)	mean LP(Å)	$\Delta V\%$
HFsol-3c	72.677	72.738	1.001	384197.7	72.697	-2.61
PBEsol0-3c	72.940	72.724	0.997	386913.6	72.868	-1.92
HSEsol-3c	72.919	72.712	0.997	386622.8	72.850	-1.99
B3LYP-D*/TZVP ⁸⁵	73.158	73.302	1.002	392311.9	73.206	-0.55
Exp. ⁸⁹	73.340	-	-	394481.1	-	-

101 with fluoride as counterion to save as many symmetry operators as possible. The final model structures result in a tetragonal space group that still retains 16 symmetry operators with primitive unit cells containing about 2800 and 3600 atoms, respectively.

In Table II we report the optimized lattice parameters and unit cell volume for MIL-100 computed with sol-3c composite methods along with experimental data and B3LYP-D*/TZVP results (hereafter we use TZVP as shortcut for the combination of pob-TZVP and 6-311G(d,p) basis sets) taken from Ref⁸⁵ Although the computed unit cell is tetragonal, the difference between a and c parameters is very small with the c/aratio being close to one. Therefore, a very small deviation from cubic lattice is predicted for MIL-100 so that the optimized structures can be considered as pseudo-cubic, with a mean lattice parameter remarkably close to the experimental values. The overall structural results obtained with our sol-3c methods are consistent with both B3LYP-D*/TZVP calculations and experimental data, in particular the DFT composite methods provide a percentage deviation of the unit cell volume from experimental data below 1.5%, while HFsol-3c shows a slightly larger deviation of about 4.2%.

The same excellent agreement is also obtained for the even larger MOF MIL-101, as reported in Table III. Experimental data are available solely for MIL-101(Cr) for comparison. For lattice constant and volume a remarkable accord with predicted results is observed. A similar correspondence is obtained for MIL-101(Al), too, but in this case the lattice parameters has been estimated by considering the similarity between the structure of MIL-100 and MIL-101 and scaling the volume of MIL-101(Cr) with the ratio of the volumes of MIL-100(Cr) and MIL-100(Al).

Overall, present results for giant-MOFs are quite striking if one considers the size and complexity of those systems. Furthermore, it is worth noting that B3LYP-D*/TZVP calculations on MIL-100 required supercomputing resources with 1024 CPU cores, at least, whereas with hybrid sol-3c methods a full relaxation of the structures can be easily carried out on just 60 CPU cores in less than a week⁹⁰. Therefore, accurate large-scale calculations become affordable on smallto-medium computing resources that are commonly available in many research laboratories³⁴.

IV. CONCLUSIONS

In this work, we reported on the application of a family of ab initio composite methods designed for solid-state calculations to study several properties of Metal-Organic Frameworks.

In particular, we focused on composite methods based on hybrid density functional approximations. The combination of both global and range-separated hybrid functionals as PBEsol0 and HSEsol, combined with a double-zeta quality basis set and augmented with properly tuned semi-classical corrections turns out to be quite effective and reliable in modelling MOFs. They show a remarkably good performance in predicting structures, band gaps, vibrational frequencies and partly adsorption energies.

We also benchmarked the simpler HFsol-3c composite

Al ^{III}	a(Å)	c(Å)	c/a	volume(Å ³)	mean LP(Å)	$\Delta V\%$
HFsol-3c	86.514	86.588	1.0009	648082.8	86.539	-2.87
PBEsol0-3c	87.464	87.497	1.0004	669341.9	87.475	0.31
HSEsol-3c	87.442	87.476	1.0004	668858.8	87.454	0.24
Exp. ^a	87.383	-	-	667247.6	-	-
Cr ^{III}	a(Å)	c(Å)	c/a	volume(Å ³)	mean LP(Å)	$\Delta V\%$
HFsol-3c	88.677	88.684	1.0001	697374.4	88.679	-0.64
PBEsol0-3c	88.843	88.792	0.9994	700848.0	88.826	-0.14
HSEsol-3c	88.821	88.772	0.9994	700340.0	88.805	-0.22
Exp ⁹¹ .	88.869	-	-	701860.3	-	-

TABLE III. Predicted LPs of MIL-101 (M^{III}) optimized at HFsol-3c, PBEsol0-3c and HSEsol-3c level of theory in comparison with experimental data⁹¹ along with percentage deviation of unit cell volume ($\Delta V\%$)

^{*a*} Estimated values, see text for details.

method. Although it shows some drawbacks due to the limitation of the Hartree-Fock method combined with a minimal basis set, the smaller computational cost makes it promising for either high-throughput screening of MOFs or as an entry level for very large-scale calculations on MOFs that contain thousands of atoms in the unit cell.

Overall, hybrid sol-3c composite methods provide results of comparable or with even better quality than the more demanding B3LYP-D3(ABC)/pob-TZVP-rev2 method. As an example, the cost of a single SCF cycle and gradient calculation for MIL-127(Al) a MOF with more than 400 atoms in the unit cell and 8 symmetry operators, is 1.5-2.0 times larger than the hybrid sol-3c composite methods (see Fig.s S12-S13). This proves that they are of practical use also for complex solids as MOFs. Furthermore, the effective deploy of hybrid functionals reduces the delocalization error that plagues DFT approximations in solid state calculations, in particular when transition metals are involved.

We are confident that hybrid sol-3c composite methods are valuable and cost-effective computational tools for MOFs and can be used for In Silico prediction of properties beyond plain structural features⁹².

CONFLICTS OF INTEREST

There are no conflicts of interest to declare.

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SUPPLEMENTARY MATERIAL

See Supplementary Material for: lattice parameters, conventional cell volumes, metal-linker bond lengths and additional figures of the MOFs in the MOF-LC15 dataset; fundamental band gaps, unit cell polarizabilities and dielectric

constants of selected MOFs; conventional cell parameters, metal-linker bond lengths, adsorption energies and figures of CO and CO₂ adsorbed in CPO-27(M) (Mg,Zn) and HKUST-1(Cu); cost efficiency methods comparison for MIL-127(Al).

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