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New Model for Predicting Adsorption of Polar Molecules in Metal–Organic Frameworks with Unsatuated Metal Sites

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Supporting Information

ABSTRACT: Conventional molecular models fail to correctly describe interactions of adsorbates with coordinatively unsaturated sites (CUS) present in a large number of metal–organic frameworks (MOFs). Here, we confirm the failure of these models for a prototypical polar adsorbate, carbon monoxide, and show that simply adjusting their parameters leads to poor agreement with experimental isotherms when outside the fitting conditions. We propose a new approach that combines quantum mechanical density functional theory (DFT) with Monte Carlo simulations to rigorously account for specific interactions at the CUS. By explicitly including electrostatic interactions and employing accurate DFT functionals that describe dispersion interactions, our modeling approach becomes generally applicable to both polar and nonpolar molecules. We demonstrate that this CUS model leads to substantial improvement in carbon monoxide adsorption isotherm predictions, and correctly captures the coordination binding mechanism. This paper represents a major stepping stone in the development of a robust, transferable and generally applicable approach to describe the complex interactions between gas molecules and CUS, with great potential for use in large-scale screening studies.

Metal–organic frameworks (MOFs) are the focus of many recent studies due to their potential role in a variety of applications. They can have extremely high porosities and surface areas, as well as unique binding sites. These attributes, combined with the ability to tailor MOFs through metal and ligand selection, promote them as ideal candidates for challenging gas separations. The focus of this research is on a subset of MOFs that contain coordinatively unsaturated sites (CUS), also known as open metal sites (OMS). These sites form when the metals are not fully coordinated to ligands but are also bound to guest molecules, e.g., solvent, in the as-synthesized form. Upon preadsorption activation, these molecules can be removed, leaving free CUS. These sites are then able to form strong coordination bonds with certain adsorbates that are able to donate electrons to the metal atoms. This selectivity feature of CUS has already been exploited to separate mixtures of physically similar gases, e.g. ethane/ethylene, which are hard to separate using other means.

Due to the sheer number of potential combinations of linkers and metals, the number of available MOFs is extremely high. Screening this huge number of possible structures for a particular gas separation using experimental techniques is prohibitively costly and time-consuming. For this reason, high-throughput computational screening approaches that have recently emerged could play a pivotal role in assessing performance of MOFs for gas separations prior to more detailed experimental studies. A caveat to the success of these strategies is the well-documented failure of conventional molecular models in accounting for the coordination interaction between CUS and certain gas molecules. This has led to large underestimations of simulated adsorption isotherms when compared to experiment for those adsorbates in which this interaction plays a key role, e.g. ethylene. In recent years, a few approaches that make use of quantum mechanical (QM) calculations in combination with classical grand-canonical Monte Carlo (GCMC) simulations were recently reviewed, and the reader is referred to those articles for a more detailed discussion. Our particular strategy restricts the use of QM to describe the local interaction at the CUS, while relying on standard molecular models to describe interactions with the remainder of the framework. This enhances the transferability of the model in comparison with alternative approaches, and we believe this is essential for a CUS model to be viable for high-performance of MOFs. DOI: 10.1021/acs.jpclett.8b00967

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throughput screening simulations. Indeed, our previous work showed that the model is both transferable to chemically similar adsorbates (ethylene to propylene) and to different MOFs with the same CUS unit. No such transferability has yet been demonstrated for other CUS modeling approaches.

Our previous work focused solely on nonpolar adsorbates, ignoring electrostatic interactions for simplicity. Here, we generalize our approach to deal with polar adsorbates for which electrostatic interactions are important. We have chosen carbon monoxide as a test case due to its simplicity and importance in practical applications. Carbon monoxide is a gas present in a variety of industrial sectors. It is commonly formed as an unwanted byproduct in incomplete combustion reactions of various oil and gas mixtures. Therefore, its removal from these streams is an important step in reducing pollution. Carbon monoxide can, however, also be a valuable feedstock. It is often produced from either coal gasification or steam reforming, as a component of synthesis gas mixtures (mainly made up of H$_2$CO). This is because for many industrial uses, such as the industrial production of methanol, aliphatic alcohols and aldehydes, it is advantageous to have both gases present; also CO is an important reducing agent within metal refining. However, the correct ratio of H$_2$:CO must first be obtained for many of these applications, requiring cost-effective gas separations, in which MOFs could play a role. Furthermore, for many applications, the use of pure carbon monoxide is required or industrially preferable, importantly, as a primary component in the production of acetic acids. The separation of pure carbon monoxide from a gas mixture, such as synthesis gas, becomes much more demanding when attempting to remove any nitrogen components. This is due to the similar boiling points of carbon monoxide ($\approx -192 ^\circ$C) and nitrogen ($\approx -196 ^\circ$C), making distillation separation costly and difficult.

The main challenge faced in simulating polar gases, like CO, with our DFT/GCMC coupled approach is that we will now need to account for the effect of electrostatics in the isolation of the local CUS interaction, as discussed in more detail later. We will also improve our approach by applying a DFT method that accurately accounts for dispersion interactions through the use of the vdW-DF2 exchange-correlation functional. This is an important step in reducing the number of approximations required in our previous work, which was based on energies obtained with the PBE exchange-correlation functional that were assumed to include no dispersion contributions. The step of extending the model to polar adsorbates is crucial to fully establishing the transferability and generality of our approach. To this end, we also performed calculations for ethylene to demonstrate that transferability is retained with the new procedure. This paper will show that our model can indeed provide good quantitative agreement with experiment and can potentially be used as an important tool for predicting MOF selectivity in gas mixtures that include both polar and nonpolar components.

As carbon monoxide has orbitals with electron-donating capabilities, it will interact strongly with CUS sites. For example, Bordiga et al. experimentally studied CO adsorption in HKUST-1 MOF, and found that electrostatics alone was unable to account for the interaction with the copper atoms, and that a $\pi$–metal interaction was taking place. We computed the adsorption energy profiles for CO adsorption at the unsaturated Cu site of HKUST-1 using DFT. The binding energies and distances at the energy minima are shown in Table 1 for each of the DFT calculations (the PBE functional was included to enable direct comparison with our previous approach). The adsorption energy found for carbon monoxide is nearly as strong as that of ethylene, confirming that the CUS plays a crucial role in carbon monoxide adsorption. Additionally, the binding energy of $\approx -29$ kJ/mol for carbon monoxide in HKUST-1, found with vdW-DF2, is promising as it matches the experimental enthalpy of adsorption found by Rubes et al. (29 kJ/mol). Furthermore, an increase in the DFT adsorption energy for ethylene from PBE, $-23.3$ kJ/mol, to the vdW-DF2 case, $-36.4$ kJ/mol, should be noted, suggesting that dispersion is being accounted for in the latter case.

The binding orientations for the vdW-DF2 energy minima can be seen in Figure 1a,b. The orientations shown for carbon monoxide and ethylene at the minimum energy are the same for both exchange-correlation functionals. We have examined several other possible orientations for the CO molecule at the CUS (see the Supporting Information, Figure S1). The most favorable orientation, shown in Figure 1b, involves adsorption parallel to the Cu–Cu vector, with the carbon atom closest to the metal. This agrees closely with a relatively recent DFT study conducted by Supronowicz et al.

Together with previous studies, our DFT results show that accurately modeling CO adsorption in MOFs will require CUS interactions to be accounted for within GCMC simulations. In the literature there is surprisingly little in terms of adsorption simulations of carbon monoxide in MOFs, especially on CUS-containing MOFs. Fischer et al. investigated the performance of existing CO models in GCMC simulations, and used the best performing model to predict adsorption in MOFs for CO/H$_2$ separation. Karra et al. also used GCMC simulations to predict CO/N$_2$ separations in HKUST-1 without explicitly accounting for any CUS interaction, and appeared to obtain good agreement with experiment. However, more recent work by Martín-Calvo et al. illustrated that using the same model led to underestimation by simulation in the low pressure region, in which solid–fluid interactions dominate. Those authors subsequently developed a new CO model that was fitted to experimental vapor–liquid equilibrium data.

In this work, several existing models will be tested for CO adsorption in MOFs to choose the most suitable model for further study. Martín-Calvo et al. carried out a comprehensive comparison of CO models for their ability to predict vapor–liquid equilibrium, and we will use their study as a basis for our assessment. In particular, they examined four different models, UFF, Straub and Karplus (S&K), Piper et al., and their own model, which we will call the M-C model. Of these, the UFF and M-C models showed the best agreement...
with liquid−vapor equilibrium data. The S&K model uses the same geometry as M-C, but the different parameters lead to a much poorer performance in vapor−liquid equilibrium simulations. The Piper et al. model was not investigated further here, as it uses a complex four-site point charge geometry, making it challenging for later combination with our CUS model, and also performed poorly in liquid−vapor equilibrium simulations. The Lennard-Jones (LJ) parameters and point charges for all CO models considered can be found in Table S1, and a visual representation of the site layouts is shown in Figure S2.

Martín-Calvo et al. also proposed a modified model in which the LJ \( \epsilon \) values for the CO−MOF interaction were increased by a factor of 1.2 relative to the standard mixing rules, and some of the mixed \( \sigma \) values were very slightly altered, in order to match CO adsorption in HKUST-1 at 298 K. The CO-MOF parameters for this modified M-C model are shown in Table S2 together with parameters obtained with the standard Lorentz−Berthelot combining rules to enable direct comparison. Here we will compare the performance of these models for predicting adsorption of CO in HKUST-1, but also in MOF-5, or IRMOF-1, as an example of a widely studied MOF without CUS. In particular, we will assess the performance of the models against high pressure experimental adsorption data of CO in HKUST-1, which was not available at the time of the Martín-Calvo study.

In agreement with Martín-Calvo et al., both the UFF and M-C models largely underestimate adsorption at all conditions when compared with both sets of HKUST-1 experimental data (Figure 2a,b). As both models replicate vapor−liquid equilibrium data very well, this is unlikely due to the gas−gas interactions and is caused instead by not accounting for specific interactions with the CUS of the MOF, which these force fields were not designed to describe. Unsurprisingly, the modified M-C model performs well in comparison with experimental adsorption at 298 K (Figure 2a), since the scaling parameters for CO−MOF interactions were designed to fit this particular data set. This scaling uniformly enhances dispersion interactions between CO and all the framework atoms (see Table S2), leading to higher adsorption. The S&K model also shows good agreement with experiment at 298 K, but this is now due to the much higher values of the CO point charges (see Table S1), leading to stronger electrostatic interactions with the framework (see Figure S3). The price of these enhanced electrostatics is a much poorer performance in vapor−liquid equilibrium simulations. Therefore, error cancellation between gas−gas and gas−solid interactions is the most likely explanation for the rather fortuitous adsorption agreement at 298 K in the case of the S&K model.

Although the modified M-C model closely agrees with the Wang experimental data set, to which it was fitted, it is not able to match the isotherm of Rubes et al. at 303 K (Figure 2b). In fact, Figure 2a clearly shows that the Wang et al. isotherm has a lower adsorbed amount than that of the more recent work of Rubes et al., despite the latter corresponding to a slightly higher temperature (303 K). This is true even after
Figure 2. Simulated (lines) and experimental (symbols) carbon monoxide adsorption isotherms in HKUST-1 [(a) 298 K; (b) 303 K] and IRMOF-1 [(c) 298 K]. In panel a, the experimental data are from Wang et al. In panel b, the experimental data are from Rubeš et al. 

The CO CUS model is built by isolating the CUS local interaction from the DFT adsorption energy profile and that enhancing electrostatics or the LJ parameters fails to adequately correct for this. Furthermore, these techniques also lose backward compatibility for MOFs without CUS. There-fore, we will investigate if isotherm agreement can be improved by explicitly accounting for the CUS interaction through combining QM and GCMC. We will compare simulations only with experiment across the full isotherm. This, in conjunction with the HKUST-1 results, further reinforces the idea that these models are capturing the gas–gas interaction correctly, as well as the standard van der Waals and electrostatic interactions with the solid. They fail when CUS are present in the MOFs, which is to be expected, and this makes both of them good candidates for combining with the CUS model proposed here. The failure of the modified M-C model shows that, although strengthening the LJ parameters improves agreement in HKUST-1 at low pressure, the parameters cannot be transferred to IRMOF-1. This raises doubts about the possibility of generally transferring this model to other MOFs (both with and without CUS) without further parameter fitting.

Based on the above results, we are now in a position to select the most appropriate CO model to be combined with the QM-based CUS interaction. When looking at the performance of the UFF and M-C models, both replicate vapor–liquid equilibrium data well and also both appear to capture the van der Waals and electrostatic interaction with IRMOF-1 correctly. The decision was made to use the M-C model for the CUS method due to it being able to better replicate the experimental dipole moment of the carbon monoxide molecule, with the M-C model exactly matching experiment at 0.112 D, while UFF is significantly higher at 0.58 D.

It was clearly shown that none of the existing CO models can fully describe the adsorption mechanism on HKUST-1, and that enhancing electrostatics or the LJ parameters fails to adequately correct for this. Furthermore, these techniques also lose backward compatibility for MOFs without CUS. Therefore, we will investigate if isotherm agreement can be improved by explicitly accounting for the CUS interaction through combining QM and GCMC. We will compare simulations only to the Rubeš et al. data set, as it appears to be more reliable as discussed previously, and covers a wider pressure range.

The CO CUS model is built by isolating the CUS local interaction from the DFT adsorption energy profile, with this CUS local interaction profile then fitted to a modified Morse
potential for use in GCMC simulations. The CUS binding energies are isolated by removing the standard force field energies used within the GCMC simulations, i.e., the LJ potential and electrostatic potential (assumed to be negligible for ethylene). In our previous work’s approach (PBE-based), the DFT calculations were assumed to contain no dispersion and therefore only the repulsive contribution of the LJ potential needed to be removed from the DFT profile. In the new procedure (vdW-DF2-based) the DFT profile contains dispersion interactions, and therefore the full LJ potential can be subtracted from the DFT profile, rather than requiring the repulsive contribution to be decoupled from the LJ potential. Further details of the CUS model can be found in the Computational Methods.

When our previous CUS model is used, based on DFT calculations with the PBE functional, we observe a significant improvement relative to the standard models, but simulations now significantly overestimate adsorption (Figure 3). This was initially surprising, as this procedure had been successfully used to predict ethylene adsorption in the same MOF, and suggests that the assumption that PBE energies contain no dispersion contribution is not entirely valid in the case of CO. As can be seen in Figure 3, the CUS model derived from the vdW-DF2 functional, based on an accurate treatment of dispersion interactions, provides much better agreement with experiment throughout the entire pressure range.

Figure 4 shows selected simulation snapshots of CO adsorption in HKUST-1. At very low pressures, as expected from our DFT calculations, the CUS is the preferred binding site for CO molecules (Figure 4a). This is in contrast to predictions using standard models, where CO was seen to adsorb first in the small cages. As the pressure increases, the small cages begin to be populated as they also present strong adsorption energies (Figure 4b). At higher pressure (Figure 4c,d), CO molecules occupy a variety of sites, with a relative preference in the order CUS > small cages > large cages.

To further validate our improved protocol for including the CUS interaction, it was also applied to the previously studied case of ethylene adsorption in HKUST-1. The ethylene parameters obtained with the new protocol will also be transferred to a different copper paddlewheel MOF (PCN-16) to confirm that the CUS model’s transferability demonstrated by Campbell et al. is maintained. As shown in Figure 5a, the new procedure, utilizing vdW-DF2, leads to practically identical results as obtained with PBE for this system. This arises directly from the similar minimum energy binding distances obtained with the two DFT approaches (binding distances are PBE 2.61 Å and vdW-DF2 2.65 Å) and similar CUS specific interaction (PBE −38 kJ/mol, after adding dispersion, and vdW-DF2 −39 kJ/mol). The good agreement with experiment demonstrates the generality of our new approach over both polar and nonpolar adsorbates. Furthermore, using the exact same parameters for the CUS interaction obtained in HKUST-1 leads to very good predictions of adsorption in PCN-16 (Figure 5b), again with negligible differences from previous work. This highlights that the new procedure retains the same transferability demonstrated for copper paddlewheel MOFs. In Figures S5 and S6, it is also shown that the CUS model is relatively insensitive to the choice of framework point charges, and the adsorption isotherm is only slightly affected.

In summary, in this paper we have shown that existing carbon monoxide models are unsuitable for capturing the orbital behavior of CUS-containing MOFs, and consequently agree poorly with experimental adsorption isotherms for HKUST-1. Furthermore, attempts to improve agreement through adjustment of LJ parameters fail to find agreement across the full isotherm, and also lose backward compatibility with non-CUS containing MOFs. This work builds upon our group’s CUS model, which has been shown to be transferable across adsorbates (ethylene to propylene) and adsorbents (copper paddlewheel MOFs). It has now been successfully extended to polar adsorbates (carbon monoxide) and, through removal of dispersion-related assumptions, agrees closely with experiment across the full isotherm. Importantly, it also captures the correct binding mechanism at the CUS, which is required for accurate simulation of competitive binary adsorption. The updated procedure was also back-validated successfully against previous work, to ensure that transferability was indeed retained. Overall, this work highlights the robust nature of this CUS approach and its flexibility across different adsorbent and adsorbate types, making it an ideal candidate for use in large-scale computational screening of MOFs for gas separations.

**Computational Methods**

Dispersion and repulsion interactions were described through the standard Lennard-Jones 12−6 potential. Parameters from the DREIDING force field were used to describe all the MOF framework atoms apart from copper. Parameters for copper atoms are not present in DREIDING and therefore the Universal force field (UFF) was used. This combination of parameters has been successfully used to model adsorption in MOFs without CUS. The point charges used to describe the framework’s electrostatic interaction were those of Castillo et al., which have been used previously for carbon monoxide adsorption simulations in MOFs. For ethylene, a united atom approach was used for each CH group, with parameters taken from the TraPPE force field. This force field was selected as it is fitted against vapor−liquid equilibrium data, helping to ensure that the gas−gas interactions are correctly captured and thus enabling the focus of this research to be solely on the gas−solid interactions. Furthermore, the TraPPE force field has been used previously to describe adsorption of alkenes and alkanes in MOFs. With the exception of the modified M-C model, all cross-species LJ interactions were estimated using the standard Lorentz–Berthelot combining rules.

![Figure 3. CUS model simulated (lines), non-CUS model simulated (dashed line), and experimental (points) carbon monoxide adsorption isotherms in HKUST-1.](image3.png)

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Adsorption isotherms were obtained through GCMC simulations using the open source code Music. All framework and adsorbate atoms were kept rigid, and an interaction cutoff distance of 13 Å was applied for the LJ interactions. Ewald summations were applied in all solid–fluid electrostatic interactions, while fluid–fluid electrostatic interactions were described using the Wolf summation method. We made use of pretabulated grids (PMAPs) to speed up the calculation of solid–fluid LJ and electrostatic interactions, with a grid spacing of 0.15 Å. For all GCMC simulations that did not include CUS interactions, cavity bias based on the LJ PMAP was used for insertion and deletion trials. In the case of simulations explicitly including CUS interactions, insertion and deletion were done randomly. In addition to insertion and deletion trials, molecules were allowed to rotate and translate, using optimized maximum displacements. 600 000 000 steps were used for all non-CUS model GCMC simulations and 100 000 000 were used for all CUS model GCMC simulations. [Note: in the case of the S&K model for CO (see Table S1) due to a memory allocation error within the Music version, bias insertion could not be used; to compensate for this, the number of steps was increased to 1 000 000 000.] The first 50% of steps were ignored to ensure equilibration, and the remaining steps were split into 20 equal blocks for error analysis. Pressure values were converted to fugacities for input into the simulation code using the Peng–Robinson equation of state. The final absolute adsorbed amounts from the simulations were converted to excess, for comparison with experiment, using the Myers and Monson method. Furthermore, except where noted, experimental adsorption isotherms were scaled by the ratio of the theoretical over experimental nitrogen pore volumes, so that they can be directly compared to GCMC simulations on a perfect crystal. A more detailed discussion of the pore volume scaling procedure can be found in the Supporting Information, and previous work.

Figure 4. Diagram showing the filling of the HKUST-1 framework at different pressure points throughout the GCMC CUS model simulations (303 K). Color code for atoms is blue - copper, red - oxygen, brown - carbon, white - hydrogen, yellow - CO binding to CUS, black - CO adsorption on large cage, and green - CO adsorption on small cage. We note that the simulation box is periodic in all three directions of space.
The CUS approach has been described in detail in previous work,\textsuperscript{5,11} therefore here we will focus on the changes to the procedure that generalize it to handle polar molecules. The procedure comprises 5 key stages:

1. Quantum-mechanical calculations to obtain the adsorption energy profile between the adsorbate and the CUS.
2. Isolation of the CUS contribution from the DFT profile.
3. Fitting the resulting profile to a modified Morse potential.
4. Including the new CUS interaction site in GCMC simulations.
5. Validating GCMC adsorption isotherms against experiment.

For the DFT calculations, in our previous work\textsuperscript{5} we used the PBE exchange-correlation functional,\textsuperscript{20} which was assumed to account for no dispersive interactions. Here, we relax this assumption by applying a DFT functional that accurately accounts for dispersion. There are two commonly used approaches to account for dispersion interactions within DFT calculations, using an explicit correction term or implicitly accounting for it within the exchange-correlation functional.\textsuperscript{50} The former approach can be successfully applied with functionals assumed to have little to no dispersive interaction, and can even be combined with dispersion-including functionals if they are thought not to fully capture the long and short-range of the dispersive interaction.\textsuperscript{54} However, the success of this approach is very dependent on the system being studied and can also involve partially scaling the correction\textsuperscript{55} when applied to exchange-correlation functionals already partially capturing dispersion, which adds an empirical nature to the procedure. As such, the decision was made to focus on exchange-correlation functionals that implicitly account for dispersion. The vdw-DF2\textsuperscript{52} was selected as it has been successfully employed in force field development for MOFs, and provided better agreement with MOF experimental isotherms than an approach utilizing PBE-D\textsuperscript{52}-corrected DFT calculations.\textsuperscript{53} We also carried out calculations with PBE for comparison with our previous approach.

DFT calculations with the PBE\textsuperscript{20} exchange-correlation functional were carried out with CP2K software\textsuperscript{53} using a similar protocol as in our previous work.\textsuperscript{5} The basis sets used for all but the Cu atoms were triple-ζ plus polarization (TZVP) with PBE-optimized Goedecker pseudopotentials.\textsuperscript{54–56} These basis sets were unavailable for copper, and therefore the double-ζ plus polarization (DZVP) sets were used. The energy cutoff selected for PBE was 600 Ry, and spin polarization was included. Furthermore, a single point counterpoise correction\textsuperscript{57} was used to account for basis set superposition error (BSSE). The importance of BSSE for carbon monoxide calculations was much higher than that of the previous ethylene DFT calculations for PBE optimization calculations at the minimum, the BSSE corrections for ethylene and carbon monoxide were ≈2 kJ/mol and ≈14 kJ/mol, respectively.

Because vdw-DF2\textsuperscript{52} was not available in CP2K, Quantum Espresso\textsuperscript{58} DFT calculations using the nonlocal correlation vdw-DF2 functional were performed with the periodic PWSCF v.5.3.0 code, using the same DFT optimization procedure as outlined in Campbell et al.\textsuperscript{5} Plane-wave basis sets were used to describe the valence electrons together with norm-conserving pseudopotentials within the Troullier-Martins approach for the core electrons.\textsuperscript{59} It should be noted that as plane-wave basis sets are being used there will be no BSSE error present. The Kohn–Sham orbitals were expanded with 50 Ry cutoff for the kinetic energy and 200 Ry cutoff for the charge density. The first Brillouin zone integrations were performed with the Marzari-Vanderbilt smearing method at the gamma point. The convergence criteria were as follows: convergence threshold for self-consistency was 1 × 10\textsuperscript{-6} Ry (using a local-density-dependent Thomas-Fermi screening mixing mode with a factor of 0.7 for self-consistency) and convergence on forces was 1 × 10\textsuperscript{-3} Ry/au.

Once each adsorption energy profile for the adsorbate with respect to the CUS has been determined, the CUS contribution must be isolated. The exact method for this isolation depends on the adsorbate and exchange correlation functional being used. In the case of ethylene, it is assumed that electrostatics do not play a significant role in adsorption in MOF frameworks.\textsuperscript{54} This assumption is not valid for carbon monoxide, as discussed, and therefore the electrostatic energies must also be subtracted from the DFT profile. Furthermore, the PBE functional is assumed to not account for dispersion interactions and therefore only the repulsive contribution from the LJ potential must be subtracted from the DFT profile. This is achieved by applying the Weeks–Chandler–Andersen (WCA) approximation\textsuperscript{60,61} on the LJ potential. With vdw-DF2, the dispersion term should also be removed from the DFT profile. Therefore, the dispersion and repulsion contributions can simply be removed together through the standard LJ potential, eliminating the need for the WCA approximation. The variations in the interaction forms for each of the systems can be seen in eqs 1–4.

![Graph](image-url)
For carbon monoxide, the vdW-DF2 was used for the CUS model and shown in Table 2. In the case of carbon monoxide, a new interaction site was placed on the center of the ethylene double bond.

The now isolated Cu–π interaction represents the specific attraction between each adsorbate and the CUS site of the MOF. However, the interaction site used within GCMC to account for the CUS interactions must be decided upon. In the case of ethylene, a new interaction site was placed on the center of the ethylene double bond. For carbon monoxide, the carbon atom is the nearest binding site to the copper (see Figure 1b) and therefore will be the CUS interaction site used in GCMC simulations. The CUS interaction site selected was the M-C model, with framework point charges from Castillo et al. The same cutoff scheme as in previous work was used for the CUS model and can be found in the Supporting Information (eq S2 and Table S6).

## SUPPORTING INFORMATION

### Supporting Information

The following files are available free of charge on the University of Strathclyde’s data repository: Input files for all GCMC simulations and DFT calculations (http://dx.doi.org/10.15129/9aceaaff-a12f-421d-8317-0fa073ed35b4). The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.jpclett.8b00967.

CO orientations - DFT calculations; CO GCMC model details; CO electrostatic interaction profiles; effect of framework point charges; experimental pore volume scaling details; CUS cut-off scheme (PDF)

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**Notes**

The authors declare no competing financial interest.

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