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High-efficiency prediction of water adsorption performance of porous adsorbents by lattice grand canonical Monte Carlo molecular simulation†

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Water adsorption has come under the spotlight for its tremendous potential in numerous environment- and energy-related applications. Given the vast adsorbent space, computational studies play a critically significant role in facilitating the discovery of potential candidates. However, large-scale computational deployment by conventional grand canonical Monte Carlo (GCMC) to identify optimal water adsorbents is challenging due to its extreme computation time and expense. In this work, a lattice GCMC method (LGCMC) with hierarchically constructed discretized interaction of host-guest and guest-guest driven by atomistic potentials was attempted to accurately and rapidly simulate the water adsorption performance of adsorbents using a coarse-grained Molinero water (mW) model. Nevertheless, given the monatomic nature of the mW model, leading to different phase behaviors in nanoscale confinement, a remarkable discrepancy in the primitive LGCMC-predicted isotherms, especially different step positions, compared with experiments was observed. Thus, a general correction strategy of water adsorption isotherm by tuning the saturation pressure was adopted. Taking metal-organic frameworks (MOFs) as examples, simulated water adsorption isotherms consistent with experimental results were obtained by the correction strategy using LGCMC. It is worth highlighting that the simulation of water adsorption in adsorbents by LGCMC can be accomplished within a few hours, which yields a significant acceleration of two to three orders of magnitude compared to conventional GCMC simulations. Therefore, the corrected LGCMC is a powerful tool to screen a huge number of adsorbents to facilitate the discovery of potential adsorbents for water adsorption-related applications, and this study provides microscopic insights into water adsorption mechanisms in porous adsorbents.

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Introduction

Water is one of the necessities to guarantee the survival and development of mankind as well as the best green working medium in numerous environment- and energy-related applications owing to its abundance, non-toxicity and high enthalpy of evaporation.¹ The adsorption phenomenon of water in porous adsorbents gave birth to a series of energy conservation and environmental protection thermal-driven applications, for example, adsorption water harvesting, desalination and purification for obtaining fresh water^{2–4} and

adsorption heat pumps and chillers for producing heat or cold.^{5,6} The water adsorption performance of porous adsorbents including water stability, deliverable capacity and kinetics play a vital role in determining the application performance.^{7,8} In recent years, fast-growing novel adsorbents showing outstanding water adsorption performance such as metal-organic frameworks (MOFs)⁹ and covalent organic frameworks (COFs)¹⁰ have attracted much more attention than conventional adsorbents including zeolites, silica gels and porous carbons.¹¹ The best candidates are various in different adsorption applications, for which the achievable performance of adsorbent/water working pairs not only is affected by operational conditions¹² but also depends on diverse water adsorption characteristics such as hydrophilicity, water uptake, step position of adsorption isotherms and desorption hysteresis.^{1,13} Fundamentally, all these water adsorption characteristics can be acquired from water adsorption isotherms, which highlights the importance

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of acquiring water adsorption isotherms of various adsorbents for evaluating their potential in different applications.

From the perspective of experiment, water adsorption isotherms of synthesized adsorbents can be tested using gas adsorption equipment and then be used as a primary basis for evaluating application performance. In recent years, novel adsorbents with excellent water adsorption performance have been continuously reported in various adsorption applications. For example, soc-MOF-1-Cr with a record-breaking water uptake of about 2 g g^{-1} at more than 70% relative humidity was regarded as a paragon in dehumidification applications.¹⁴ Subsequently, the reported MOF-801-Zr used for water harvesting from desert air, an Al-based MOF-303 delivering more than twice the amount of water ($\sim 0.7 \text{ L kg}^{-1}$ per day in the Mojave Desert) was developed, which benefits from its high deliverable water capacity ($\sim 0.45 \text{ g g}^{-1}$) at a low relative humidity of 10–30%.^{15,16} The MOFs ZJNU-30-Zr¹⁷ and CUK-1-Co¹⁸ and the COF TpPa-1 (ref. 19) with attractive water uptake and stepwise adsorption isotherms were recognized as potential candidates for adsorption chillers driven by a low-grade heat source. It is not difficult to obtain the water adsorption isotherms of several adsorbents, but it is impractical to synthesize and test thousands of adsorbents to identify the best performer for water adsorption. An effective machine learning (ML) model was successfully developed based on an experimental water adsorption isotherm database to predict water adsorption isotherms of various adsorbents. However, three experimental structural features (accessible surface area, available pore volume and pore diameter) as input descriptors were indispensable for executing the prediction procedure.²⁰ It is still extremely challenging to comprehensively characterize all existing adsorbents, especially since around 100 000 synthesized records of MOFs were documented in the Cambridge Structural Database.²¹

From the perspective of computation, gas adsorption isotherms of a large number of adsorbents may be obtained from molecular simulation with the crystal structure of adsorbents, which can greatly reduce the experimental time and cost in the high-efficiency discovery of top performers, and have been successfully practiced in various applications such as carbon capture, methane storage and hydrogen storage.^{22–24} The conventional and most widely used approach for simulating water adsorption in porous materials is the grand canonical Monte Carlo (GCMC) simulation based on a grand canonical ensemble (μ VT), which has been used to simulate water adsorption in porous carbon, zeolites and MOFs.^{25–27} For example, GCMC assists the exploration of the water adsorption mechanism in ZIF-8-Zn,²⁸ the temperature effect on water adsorption and desorption processes in MIL-101-Cr,^{29,30} and the influence of structural defects on the water adsorption properties of Zr-based MOFs such as MOF-801-Zr (ref. 31) and UiO-66-Zr.³² In each GCMC cycle, the addition or deletion of water molecules is implemented and the total potential energy of the simulation system is calculated, and these processes are repeated till the system

reaches equilibrium to output accurate water adsorption results. However, the hard truth is that an extremely long time is required to simulate water adsorption in adsorbents, up to tens of months of computational wall clock time on a single central processing unit (CPU) core to yield results, which is linked to the ultra-slow equilibrium convergence of the adsorbent/water system.^{28,31,33} Tracing it to its cause, except for time-consuming potential energy calculation in each cycle, the formation of water clusters during the water adsorption process due to the strong hydrogen-bonding networks causes a local free-energy minimum trap and thus leads to millions of cycles or a long time to reach equilibrium.^{28,34} Therefore, large-scale GCMC computational deployment to identify optimal water adsorbents can be very challenging due to its extremely high computation expense.

Several strategies have been proposed to accelerate the GCMC simulations. For example, the interactions between atoms of adsorbents and water molecules were precomputed on a grid to reduce the computation time of potential energy during the simulations. Paranthaman *et al.* executed water adsorption simulation in an Al(OH)(1,4-NDC) MOF and obtained consistent results with experimental water adsorption isotherms, in which 2×10^9 cycles were carried out.³⁵ Another possible remedy is using a more efficient insertion and deletion strategy such as the continuous fractional component Monte Carlo (CFCMC),³⁶ in which water molecules have a higher chance of being accepted by adding or removing molecules gradually rather than all of a sudden in regular GCMC. It has been shown to achieve convergence more rapidly than the regular GCMC in the case of water adsorption in MOF-806-Zr.³⁴ Besides, other Monte Carlo methods were also attempted; the flat histogram method based on a canonical ensemble (NVT) was illustrated in simulating water adsorption of MOF-806-Zr,³⁴ and Gibbs ensemble Monte Carlo (GEMC) based on Gibbs ensemble (NPT) was adopted to predict water adsorption isotherms of two Al-based MOF-303 and MOF-333.³⁷ However, the consumed time of water adsorption simulation by the abovementioned approaches is not clearly stated; thus their feasibility in large-scale water adsorption simulation cannot be definitively assessed. To the best of our knowledge, a suitable approach to rapidly and accurately simulate water adsorption isotherms of porous adsorbents on a large scale is still in urgent development.

A simulation method based on a hierarchically constructed lattice model (*i.e.*, the discretized free energy model replacing the continuous interaction of host-guest and guest-guest) driven by underlying atomistic potentials, named lattice GCMC method (LGCMC),³⁸ has been developed to quickly estimate gas adsorption in porous materials and practiced in the prediction of CO₂ adsorption isotherms in MOFs.³⁸ In conjunction with appropriately coarse-grained guest-guest interactions, this method can offer agreement as compared to a fully atomistic GCMC simulation. Crucially, it yielded significant two to three orders of magnitude acceleration as compared to conventional GCMC simulations.



In this work, LGCMC attempted to allow the use of a coarse-grained Molinero water (mW) model,³⁹ which may provide the possibility of high-efficiency acquisition of water adsorption isotherms in porous adsorbents. Six typical MOFs were tested to verify the feasibility of LGCMC for efficiently simulating water adsorption isotherms. Eventually, we proved that simulated water adsorption isotherms of MOFs that are consistent with the experimental results can be obtained by LGCMC within several hours. Therefore, accurate and rapid simulation of water adsorption isotherms in porous materials was realized by LGCMC, which opens up the possibility for high-throughput screening of large-scale adsorbents in water adsorption-related applications.

Materials and methods

Lattice model description

Similar to conventional atomistic GCMC, the structure and geometry information of adsorbents were taken into account as input files of LGCMC. A given crystal unit cell is partitioned into an $N_a \times N_b \times N_c$ sub-lattice cell, where N is the number of divisions along the direct lattice vector to control the resolution of the grid. In the lattice model, the original continuous interaction potential energy of guest molecules within the unit cell is replaced with the corresponding discretized free energy at lattice cells, which is the key to determine the guest molecules adsorbed in a given adsorbent. It should be noted that the lattice approach in this work is used where adsorption space is divided into subspaces rather than into specific adsorption sites.⁴⁰

For the occupation configuration of guest molecules in a given crystal unit cell, the total free energy (F_{tot}) is approximated as eqn (1).

$$F_{\text{tot}} = \sum_n F(n) + \sum_{n < m} F_{\text{int}}(n, m) \quad (1)$$

where $F(n)$ is the free energy of guest molecules in lattice cell n . Indices n and m run over all N -occupied lattice cells, and $F_{\text{int}}(n, m)$ represents the effective interaction energy between occupied cells n and m .

Specifically, $F(n)$ replacing the continuous interaction potential energy of guest molecules within adsorbents can be given by

$$F(n) = -kT \ln Q_n \quad (2)$$

where k is the Boltzmann constant, T is temperature and Q_n is the partition function for a guest molecule within the specific lattice cell n , which is related to the potential energy between the host and guest.

$$Q_n = \int_{v_n} dv e^{-\beta E_{\text{host-guest}}} \quad (3)$$

where $E_{\text{host-guest}}$ is the interaction energy between the host and guest, and β is the inverse temperature. The average of the Boltzmann factors (eqn (3)) is evaluated for each lattice cell by

using the specific sampling scheme and the efficient grid-based energy evaluation approach.³⁸ $E_{\text{host-guest}}$ is split into long-range and short-range potentials calculated by the particle mesh Ewald (PME) algorithm.⁴¹ Both long-range and short-range potentials include electrostatic interaction and van der Waals (vdW) interaction, respectively. The long-range electrostatic interaction was calculated only once at the beginning of the LGCMC simulation and stored on a potential grid. The long-range vdW interaction was calculated exactly. For short-range potential, cutoffs of 5.0 Å and 7.5 Å for electrostatic and vdW interactions were utilized, respectively.³⁸

$F_{\text{int}}(n, m)$ is the discrete free energy of guest-guest interaction ($E_{\text{guest-guest}}$), which can be ignored in Henry's region (*i.e.*, very low pressure and uptake), while the approximation of $F_{\text{int}}(n, m)$ determines the accuracy of the total free energy (F_{tot}) when the uptake is high. The continuous interaction potentials of $E_{\text{guest-guest}}$ must be discretized commensurate with the lattice grid to calculate the $F_{\text{int}}(n, m)$. The straightforward discretization has been proven to work well.³⁸

$$F_{\text{int}}(n, m) = E(r_{nm}) \quad (4)$$

where n and m denote the two different lattice cells, and the average is taken over all possible guest orientations with the restriction that their center of mass is confined to their respective cells. E represents the interaction potential, and r_{nm} is the distance between the centers of the respective lattice cells. In the lattice approach, the coarse-grained guest model rather than the atomistic model was adopted to estimate the guest-guest interaction.

LGCMC simulations are executed on the constructed lattice model with a free energy grid including the host-guest and discretized guest-guest interactions. Three types of Monte Carlo moves including particle translations, insertions and deletions are performed in each cycle, in which a random guest is displaced on a random lattice according to the acceptance probability from the standard Metropolis scheme.³⁸ The acceptance probabilities (p) are as follows.

$$p_{\text{tra}} = \min(1, \exp(-\beta(F_{\text{tot}}^{\text{try}} - F_{\text{tot}}))) \quad (5)$$

$$p_{\text{ins}} = \min\left(1, \frac{V}{\Lambda^3} (N+1) \exp(-\beta(F_{\text{tot}}^{\text{try}} - F_{\text{tot}} - \mu))\right) \quad (6)$$

$$p_{\text{del}} = \min\left(1, \frac{\Lambda^3 N}{V} \exp(-\beta(F_{\text{tot}}^{\text{try}} - F_{\text{tot}} + \mu))\right) \quad (7)$$

where Λ is the thermal De Broglie wavelength, V is the total unit cell volume, N is the number of particles, μ is the chemical potential for the guest, and F is the free energy of the system. It should be noted that there are no rotation moves attempted due to the fact that the orientational degrees of freedom of coarse-grained guest molecules used in LGCMC (*i.e.*, single particle) are averaged out during the creation of the lattice potential.



Model and force field parameters of adsorbent and guest molecules

In this work, experimental adsorption isotherms of MOFs in our updated experimental water adsorption isotherm database (EWAID)^{13,42} were selected as benchmarks to verify the accuracy of isotherms simulated by LGCMC. The crystal structures of such MOFs were collected from the Cambridge Crystallographic Data Centre (CCDC)⁴³ and RASPA package.⁴⁴ However, the existing difference between the structural properties of experimental samples and perfect crystals can be ascribed to the defects, unremoved solvents, or free ions in experimental samples.⁴⁵ Besides, there are inevitable differences due to different evaluation methods, in which theoretical values are based on geometric methods (e.g., commonly using Zeo++) and experimental ones are usually based on analysis of nitrogen isotherms (e.g., BET method).⁴⁶ The agreement between simulations and experiments is not expected for those adsorbents with inconsistent structural properties.⁴⁵ Thus, to exclude the influence of differences in structural properties on the prediction results to the greatest extent, six MOFs (in Table 1) with consistent theoretical and experimental structural properties including accessible surface area (S_a), available pore volume (V_a) and pore diameter (D_p) were eventually adopted. The process for selecting MOFs is shown in Fig. S1,† and their literature source and structural properties are provided in Table S1.†

For the host–guest interaction ($E_{\text{host-guest}}$ here is $E_{\text{MOF-water}}$), selected MOFs and the Tip4p water model were considered in this work. Their interaction includes both van der Waals (12-6 Lennard-Jones) and coulombic potential as shown in eqn (8).

$$E_{\text{MOF-water}} = \sum_{ij} 4\epsilon_{ij} \left[\left(\frac{\sigma_{ij}}{r_{ij}} \right)^{12} - \left(\frac{\sigma_{ij}}{r_{ij}} \right)^6 \right] + \sum_{ij} \frac{q_i q_j}{4\pi\epsilon_0 r_{ij}} \quad (8)$$

where r_{ij} is the distance between atoms i and j in MOFs and water, LJ parameter ϵ represents the depth of a potential well and σ is the equilibrium position of potential energy. q is the atomic charge, and ϵ_0 is the vacuum permittivity constant. LJ parameters from the universal force field (UFF)⁵² and multilayer connectivity-based atom contribution (mCBAC) charges⁵³ were adopted for all atoms of MOFs. The mCBAC charges were calculated by a trained machine learning model based on the type and connectivity of atoms in the MOF, which can be obtained in seconds and on a par with density-derived electrostatic and chemical charges (DDECs).^{53,54} More calculation details and validation of mCBAC charges can be

found in S2 section (LGCMC and force field parameters) of the ESI.† There is no doubt that the force field of specific MOFs (such as MOF-74-Mg with open metal sites) could also be obtained from first-principles methods,⁵⁵ but the high computation cost restricts their application to a large number of MOFs. The force field parameters of Tip4p water⁵⁶ are provided in Table S2.†

The discretized guest–guest interaction ($E_{\text{guest-guest}}$) in specific adsorbents was calculated. In this work, the guest is coarse-grained mW, which represents each water molecule by a single particle with tetrahedral interactions. The mW model potential^{39,57} uses the short-range interaction form of the Stillinger–Weber potential, consisting of a sum of two-body attraction terms that favor high coordination, and three-body repulsion terms that reinforce tetrahedral hydrogen-bonded configurations (eqn (9)).

$$E_{\text{mW-mW}} = \sum_{i,j>i} A\epsilon \left[B \left(\frac{\sigma}{r_{ij}} \right)^p - \left(\frac{\sigma}{r_{ij}} \right)^q \right] \exp \left(\frac{\sigma}{r_{ij} - a\sigma} \right) + \sum_{i,j\neq i,k>j} \lambda\epsilon [\cos\theta_{ijk} - \cos\theta_0]^2 \exp \left(\frac{\gamma\sigma}{r_{ij} - a\sigma} \right) \exp \left(\frac{\gamma\sigma}{r_{ik} - a\sigma} \right) \quad (9)$$

where r_{ij} is the distance between particles i and j , and θ_{ijk} is the angle defined by particles i, j and k . The parameters A, B, p, q, γ , and a give the form and scale to the potential. σ is the particle diameter, ϵ represents the depth of the two-body interaction potential, and λ scales the repulsive three-body term and determines the strength of the tetrahedral interaction in the model. All parameters are provided in Table S3.† It is worth mentioning that although electrostatic terms or explicit hydrogen atoms are not included in the mW model, it can accurately reproduce the phase behavior for the solid–liquid equilibria in bulk.⁵⁸ Besides, since the decreased number of particles and shorter range of interactions with around 4.3 Å cutoff, the computational cost of simulations using the mW model is only 1% compared with simulations using atomistic water models (such as SPC/E and TIP series models).⁵⁸

LGCMC simulations

In the LGCMC simulation, the adsorption condition was reflected by the chemical potential (μ), and it is used to evaluate the acceptance probabilities of particle translation moves. The chemical potential depends on the adsorption conditions (temperature and pressure) and the properties of the guest molecule.

Table 1 Measured (Mea.) and computed (Com.) structural properties of MOFs used in LGCMC

No.	Adsorbents	Mea. S_a (m ² g ^{−1})	Mea. V_a (cm ³ g ^{−1})	Mea. D_p (Å)	Com. S_a (m ² g ^{−1})	Com. V_a (cm ³ g ^{−1})	Com. LCD (Å)
1	Al(OH)-(1,4-NDC) ⁴⁷	546	0.22	7.70	614	0.19	7.48
2	DUT-67-Zr (ref. 48)	1560	0.60	16.60	1808	0.54	17.05
3	MIL-100-Fe (ref. 49)	1549	0.82	29.00	1931	0.88	26.60
4	MIL-101-Cr (ref. 50)	3124	1.58	34.00	3026	1.61	33.66
5	MOF-74-Mg (ref. 50)	1400	0.65	11.00	1708	0.67	11.87
6	MUF-77-Zn-methyl (ref. 51)	3600	1.85	21.10	4437	1.68	21.78



$$\mu = \mu_{\text{refe}} + RT \ln \left(\frac{P}{P_{\text{refe}}} \right) \quad (10)$$

The chemical potentials (μ_{exp}) for experimental water can be calculated by combining the Peng–Robinson–Stryjek–Vera (PRSV) state equation of water.⁵⁹ μ_{exp} corresponding to various relative pressures (P/P_0) at 298 K are provided in Table S4†. For the mW model, it is noted that the predicted saturation pressure of the coarse-grained mW model ($P_{0,\text{mW}}$, 68 Pa at 298 K (ref. 60)) is lower than the experimentally measured result ($P_{0,\text{exp}}$, 3158 Pa at 298 K (ref. 60)). Such a discrepancy is essentially determined by the monatomic nature of the mW model, of which the hydrogen-bonded water is mimicked through the introduction of a non-bond angular-dependent term that encourages tetrahedral configurations.³⁹ A loss of rotational entropy in mW particles that is significant in the gas phase entails abnormal vapor pressure. As a consequence, it causes a shift in chemical potential that refers to the excess entropy at the condensed phase between the experimental saturation pressure ($P_{0,\text{exp}}$) and the theoretical one ($P_{0,\text{mW}}$). Thus, the chemical potential for mW (μ_{mW}) is calculated and expressed as follows according to previous studies.^{58,60}

$$\mu_{\text{mW}} = \mu_{\text{exp}} + RT \ln \left(\frac{P_{0,\text{exp}}}{P_{0,\text{mW}}} \right) \quad (11)$$

where R is the ideal gas constant and T is the reference temperature. $P_{0,\text{exp}}$ and $P_{0,\text{mW}}$ are the water saturation pressure from the experiment and the mW model at T , respectively. $P_{0,\text{exp}}$ and $P_{0,\text{mW}}$ at varying temperatures can be obtained by the Clausius–Clapeyron equation according to the experimental enthalpy of vaporization of 44.52 kJ mol^{−1} and 44.18 kJ mol^{−1} of the mW model, respectively.⁶⁰ Combined with the μ_{exp} at different temperatures and pressures, the μ_{mW} of the water model can be calculated (Table S4†).

In this work, a total of 2×10^7 cycles including 1×10^7 cycles for equilibration and 1×10^7 cycles for production were implemented. The manual and original simulation code of LGCMC can be found on the websites of Schmidt's group.⁶¹ Output results including water uptake, the heat of adsorption and water distribution in adsorbents can be obtained from LGCMC. In our pre-test, the LGCMC simulations are extremely rapid and typically finish within a few minutes using a single CPU core, which benefits from the exclusive use of the pre-tabulated discretized free energy model. As such, the majority of the computational cost of simulation is contributed by the generation of a free energy model.

Results and discussion

Grid size optimization

For a given adsorbent, the grid size determining the resolution of the lattice cell is one of the most important factors affecting the computational time of the free energy model. In the construction of the lattice model, the periodic boundary conditions were used, and the unit cell

size of MOFs was set to larger than twice the cutoff (*i.e.*, larger than 15 Å). For specific MOFs, the number of gridded sub-cells depends on the size of their unit cell (Table S5† shows the size of the crystal unit cell for six MOFs used in this work) and grid size. Owing to the trade-off between the high precision and the long computational time of the free energy model using a small grid size, grid size optimization was performed first. The representative MOFs, MOF-74-Mg, DUT-67-Zr and MIL-101-Cr with small-size (<30 Å), medium-size (30–50 Å) and large-size (>50 Å) unit cells, respectively, were chosen for LGCMC under varying grid sizes (0.5 Å, 1 Å and 2 Å). It is indicated that the equilibrium water uptakes obtained using smaller grid sizes (0.5 Å and 1 Å) are slightly higher than that using a large grid size of 2 Å (Fig. 1), which may be due to the high-precision free energy model increasing the insertion sites of the water molecule. The water uptake values obtained using grid sizes of 0.5 Å and 1 Å are almost exactly the same, but there is a huge difference in computational cost. Taking MIL-101-Cr with a large-size unit cell as an example, a computation time of about 24 h is required to produce the free energy grid with a grid size of 1 Å, while around 200 h is needed for a grid size of 0.5 Å. Therefore, taking both the computational precision and time into account, a grid size of 1 Å was adopted for the computation of free energy. The free energy maps of most adsorbents can be obtained within a few hours in our tests.

Water adsorption isotherms by LGCMC

The water adsorption isotherms of six MOFs were simulated using LGCMC at 298 K and compared with reported experimental results (Fig. 2). Two important indexes including water uptake (W) and step position of isotherms (α) were focused on. In this work, α was defined as the relative pressure when half of the saturation water uptake was reached. In experiments, MOF-74-Mg shows a type I (*i.e.*, inverted L shape) water adsorption isotherm; the MOFs Al(OH)-(1,4-NDC) and MUF-77-Zn-methyl exhibit type V (*i.e.*, S shape) isotherms, and the rest of the MOFs MIL-100-Fe, MIL-101-Cr and DUT-67-Zr with hierarchical pores show type VI isotherms with multiple adsorption steps. LGCMC simulations successfully predicted similar isotherms for MOF-74-Mg and Al(OH)-(1,4-NDC) with slightly higher saturation water uptake and larger step position of the isotherm (*i.e.*, $\alpha_{\text{LGCMC}} > \alpha_{\text{exp}}$). However, almost no water uptake throughout the pressure range ($P/P_{0,\text{exp}} = 0-1$) was observed in the predicted isotherms of the other four MOFs. A rapid increase in water uptake in the LGCMC isotherm was observed at $P/P_{0,\text{exp}} > 1$ (Fig. 2 and S3†). For example, a rapid increase in water uptake was observed at $P/P_{0,\text{exp}} = 2-3$ (*i.e.*, α_{LGCMC}) for MIL-100-Fe and MIL-101-Cr, $P/P_{0,\text{exp}} = 3-4$ for DUT-67-Zr and $P/P_{0,\text{exp}} = 6-7$ for MUF-77-Zn-methyl; all of them are larger than α_{exp} in experimental water adsorption isotherms. It also indicates that the saturation vapor pressure



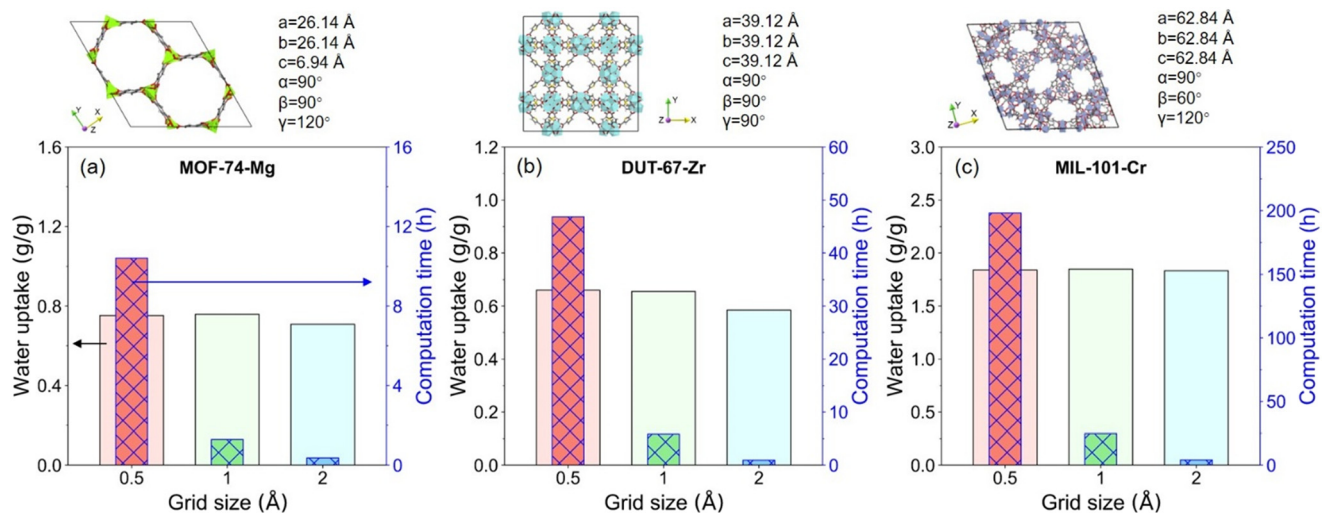


Fig. 1 Simulated water uptake (at 298 K, $P/P_{0,\text{exp}} = 7$) and calculation time of representative MOFs in LGCMC with different grid sizes of 0.5 Å, 1 Å and 2 Å. (a) MOF-74-Mg, (b) DUT-67-Zr and (c) MIL-101-Cr with different sizes of unit cells.

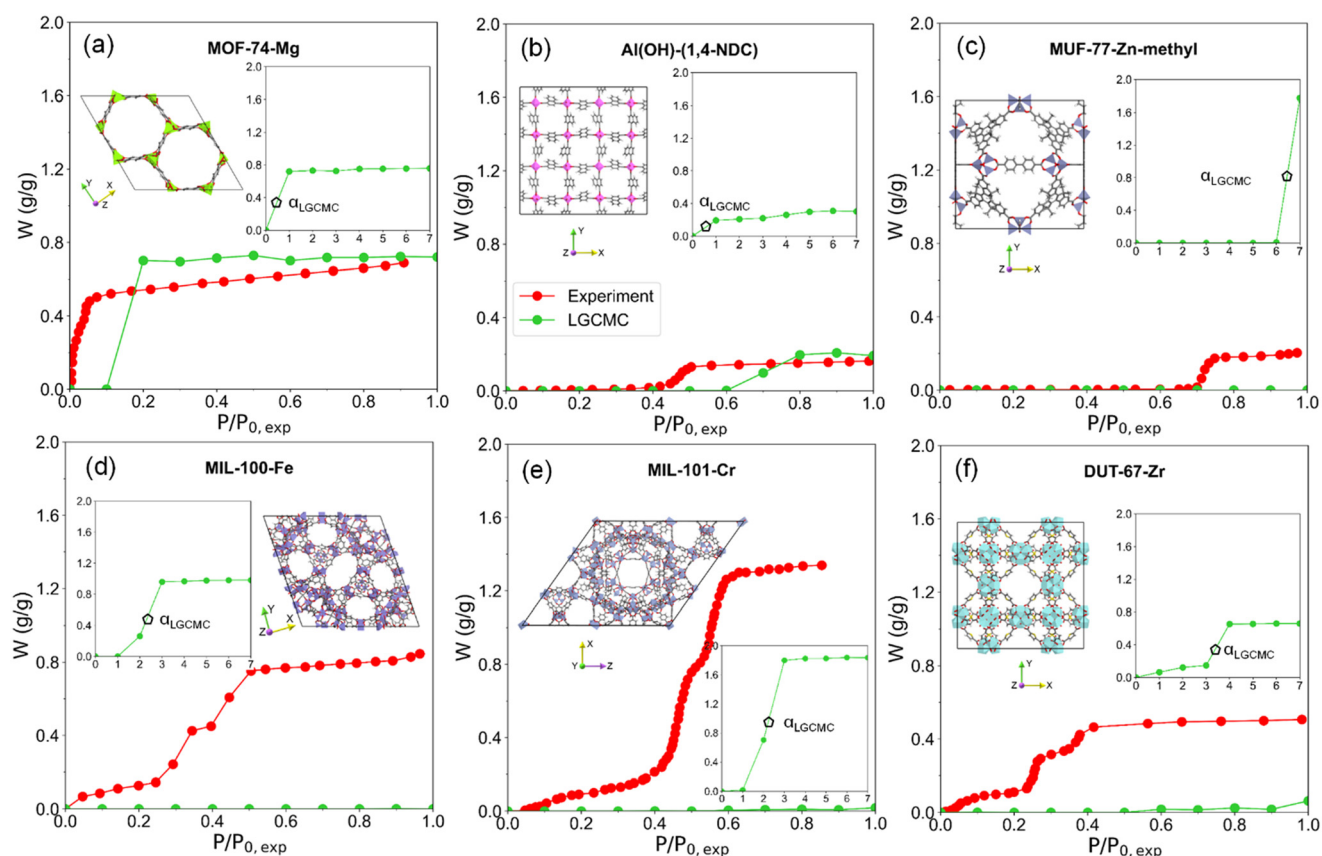


Fig. 2 Experimental water adsorption isotherms collected from the literature (sources provided in Table 1) and simulated water adsorption isotherms of MOFs by LGCMC. The inset figures show the LGCMC simulation at $P/P_{0,\text{exp}} > 1$. (a) MOF-74-Mg with a type I isotherm (i.e., inverted L shape), (b) Al(OH)-(1,4-NDC) and (c) MUF-77-Zn-methyl with type V (i.e., S shape) isotherms, (d) MIL-100-Fe, (e) MIL-101-Cr and (f) DUT-67-Zr with multiple-step isotherms. α_{LGCMC} is the step position of LGCMC isotherms.

of mW in LGCMC simulation isotherms is larger in the theoretical than the experimental measured result.

In order to eliminate the discrepancy of the saturation vapor pressure between mW and experimental water in the

bulk phase, $P_{0,\text{exp}}$ was used instead of $P_{0,\text{mW}}$ (eqn (11)). However, the phase behaviors of adsorbed water molecules confined in nanoporous materials are still significantly different from that of bulk water.⁶² The monatomic mW

model with a loss of rotational entropy makes it hard to accurately reproduce the phase behaviors of adsorbed water under nanopore confinement effects, which leads to the difference between the saturation pressure of mW and that of experimental water in nanoporous adsorbents. Therefore, the difference in saturation pressures between mW and experimental water due to the nanoconfinement effects needs to be further corrected.

Isotherm correction

The quantification of the differences in saturation pressures of mW and experimental water is crucial for accurate correction, but the saturation pressure of mW in nanopores is difficult to predict, which complicates this process. In nanopores, the pressure of water in the nanopore beginning or undergoing phase transition can be described by the condensation pressure (P_c , vapor to liquid phase during adsorption). Assuming that the nanopore confinement effects on the phase behavior of mW and experimental water are equivalent under varying pressures, then the difference in condensation pressures between mW and experimental water can also reflect differences in saturation pressures. In previous studies on water adsorption, the condensation pressure represents the water uptake that begins to rapidly increase in water adsorption isotherms,^{58,63,64} thus, the condensation pressure can be estimated according to the adsorption isotherms.

As for the mW in adsorbents, the condensation pressure ($P_{c,mW}$) can be identified based on the LGCMC simulated isotherms. Besides, the condensation relative pressure ($P_{c,mW}/P_{0,exp}$) coincides with the step position of isotherms (α_{LGCMC}) in this work. As for the experimental water in adsorbents, there is a problem in obtaining the condensation pressure of experimental water in adsorbents. First of all, direct experimental measurement of the phase transition pressure of water in nanoconfinement is very difficult. Secondly, the determination of condensation pressure based on the experimental water adsorption isotherm of the adsorbent is contrary to the original intention of simulations. Fortunately, the water condensation phenomena in nanoconfinement can be described by the century-old Kelvin equation (eqn (12)). The condensation pressure of confined water ($P_{c,Kelvin}$) in nanopores can be estimated using the Kelvin equation. In particular, the Kelvin equation has been proven applicable to atomic-scale confinement with a pore size of around 4 Å,⁶⁵ including MOFs.^{63,64}

The computation details are described as follows.

$$\ln\left(\frac{P_c}{P_0}\right) = \frac{2V_m\gamma \cos\theta}{r_p RT} \quad (12)$$

The Kelvin equation relates the condensation relative pressure (P_c/P_0) of confined water to the physiochemical properties of bulk water (the molar volume of liquid water, $V_m = 18 \text{ cm}^3 \text{ mol}^{-1}$, and the experimental liquid/vapor surface tension, $\gamma = 73 \text{ mJ m}^{-2}$ at 298 K) and the structural properties of adsorbents, *i.e.*, pore radius (r_p) is defined as one-half of

the computed pore diameters (LCD) in this work, and the pore surface hydrophilicity of adsorbents is described by the apparent contact angle (θ). The apparent contact angle of adsorbents with various hydrophilicity was theoretically estimated based on Factorovich *et al.*'s work.⁵⁸ They predicted the adsorption isotherms and the contact angles of mW in a series of cylindrical nanopores with varying hydrophilicities by grand canonical molecular dynamics (GCMC), and the correlation between the contact angle (θ) and the condensation relative pressure ($P_{c,mW}/P_{0,exp}$) of isotherms was extracted as shown in Fig. 3. It should be noted that the contact angle of water was estimated from simulations of droplets on flat surfaces with varying hydrophilicities employing molecular dynamics (MD) simulation procedures proposed in Giovambattista's work.⁶⁶ Thus, the approximate contact angle of each adsorbent can be identified based on the predicted $P_{c,mW}/P_{0,exp}$ of water isotherms by LGCMC in this work. In such a case, the condensation pressure of confined water in each adsorbent ($P_{c,Kelvin}$) can be estimated using the Kelvin equation.

Compared with the condensation pressure ($P_{c,mW}$) from isotherms by LGCMC, the difference between the condensation pressures from the Kelvin equation ($P_{c,Kelvin}$) and $P_{c,mW}$ can be converted to the difference in chemical potential by $\Delta\mu$ (schematic diagram in Fig. 4; Table S6† provides the $\Delta\mu$ of six MOFs in this work). Assuming that the chemical potential difference ($\Delta\mu$) of each adsorbent is identical throughout the pressure range, thus, the corrected chemical potential (μ_{cor}) at varying pressures for each adsorbent in LGCMC simulations was obtained by eqn (13).

$$\mu_{cor} = \mu_{mW} + \Delta\mu = \mu_{mW} + RT \ln\left(\frac{P_{c,mW}}{P_{c,Kelvin}}\right) \quad (13)$$

Based on the μ_{cor} , the corrected water isotherms predicted by LGCMC exhibit improved agreement with experimental results as displayed in Fig. 5. As for MOF-74-Mg with a small step position of $\alpha_{LGCMC} = 0\text{--}1$ or type I water adsorption isotherm (Fig. 5a), the corrected LGCMC isotherm is more consistent with experimental results, in which the step position of the

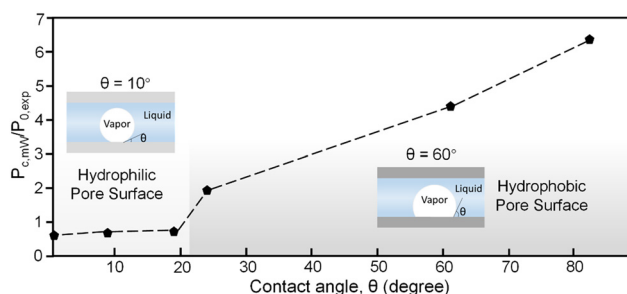


Fig. 3 The relationship between the contact angle (θ) of water in nanopores and the condensation relative pressure ($P_{c,mW}/P_{0,exp}$) of water adsorption isotherms. The contact angles of mW were adapted based on a series of cylindrical nanopores with different hydrophilicity, and the condensation relative pressures were extracted from their predicted water adsorption isotherms by GCMC simulations from Factorovich *et al.*'s work.⁵⁸



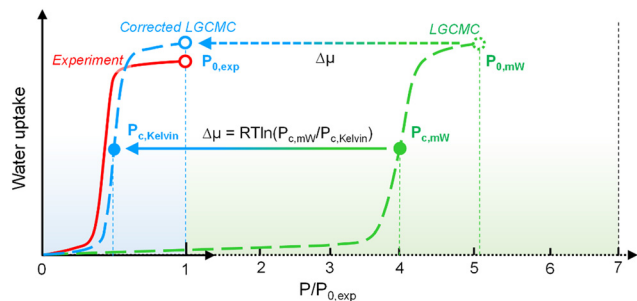


Fig. 4 Schematic diagram of water adsorption isotherms of LGCMC and corrected LGCMC. The chemical potential difference ($\Delta\mu$) was converted from the condensation relative pressure difference between LGCMC isotherms ($P_{c,mw}$) and the Kelvin equation ($P_{c,Kelvin}$).

isotherm locates at a low-pressure region (<0.05). For $\text{Al}(\text{OH})$ -(1,4-NDC) with $\alpha_{\text{LGCMC}} = 0$ –1 in Fig. 5b, the step position of the corrected isotherm is even smaller than the experimental result ($\alpha_{\text{exp}} = 0.4$ –0.5), indicating the underestimation of step position. However, subsequent comparison with the results from atomistic GCMC demonstrates that such a difference also exists in atomistic GCMC, which does not result from the mW model of LGCMC. For adsorbents MUF-77-Zn-methyl, MIL-100-Fe, MIL-101-Cr and DUT-67-Zr (Fig. 5c–f) with large step positions greater than 1, corrected LGCMC gives rise to

improved consistency with experimental isotherms. It is also found that the corrected LGCMC isotherms reproduce the experimental results of MIL-100-Fe, MIL-101-Cr and DUT-67-Zr, specifically. As for the difference in saturation uptakes between simulation and experiment, the simulated value is usually higher than experimental ones, which may be attributed to the higher porosity of the perfect crystal in the simulation than experimental samples, and the inevitable decrease in the adsorption capacity due to defects and residual solvents in the experimental samples.⁴⁵ It should be noted that the great deviation in the saturation uptake of MUF-77-Zn-methyl may be ascribed to its chemical instability during water adsorption measurement.⁵¹

Besides, water adsorption isotherms obtained by LGCMC were also compared with the results from conventional GCMC based on the atomistic water model by other researchers (Fig. 6). The saturation water uptake obtained by atomistic GCMC is generally consistent with LGCMC, which indicates that it is able to provide a quantitatively accurate prediction while using the same perfect crystal structure in simulation. The step positions of isotherms from atomistic GCMC is inevitably slightly overestimated (e.g., MOF-74-Mg (ref. 26)) or underestimated (e.g., $\text{Al}(\text{OH})$ -(1,4-NDC)³⁵ and MIL-101-Cr (ref. 30)) compared with experiments. It should be noted that the atomistic GCMC isotherm of $\text{Al}(\text{OH})$ -(1,4-NDC) showing type V in Fig. 6b was

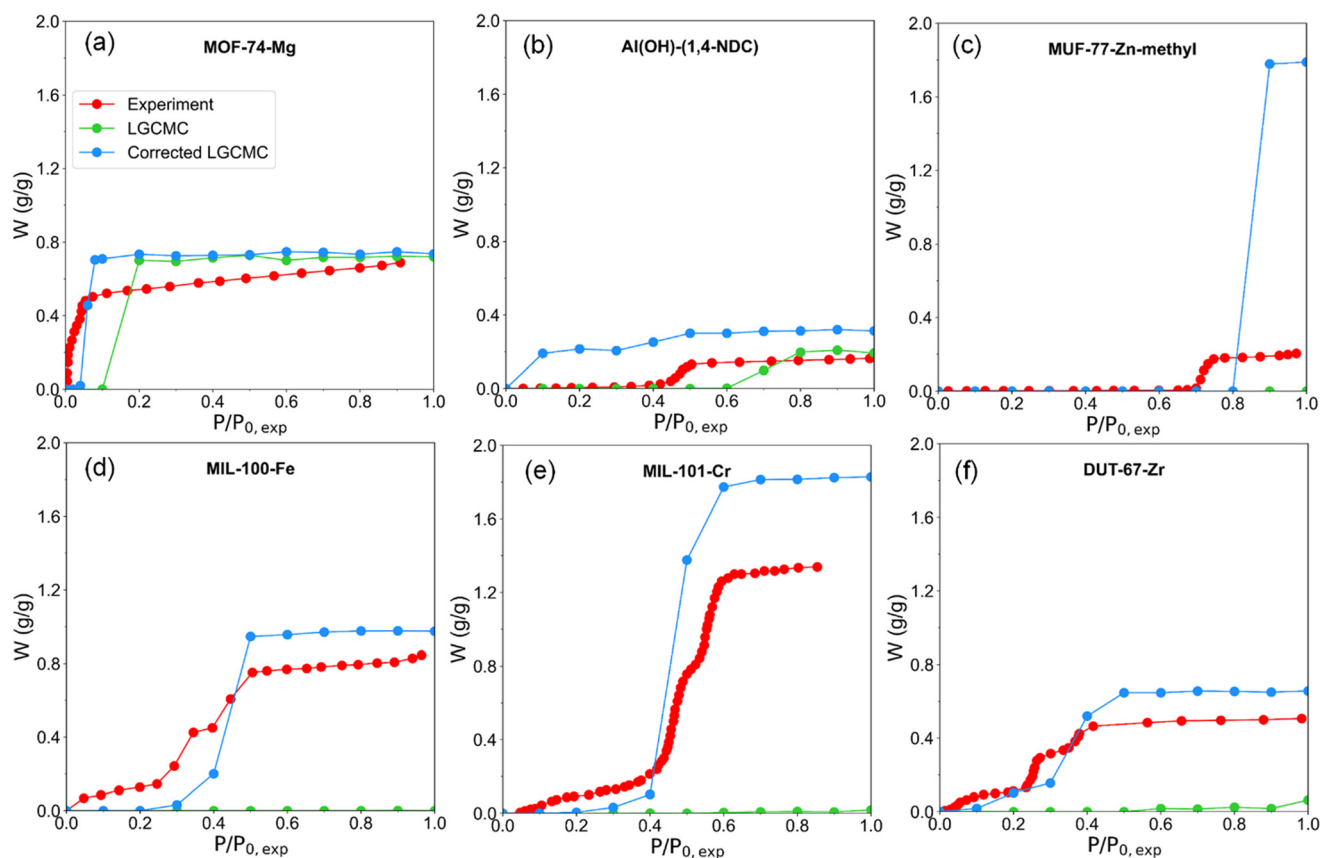


Fig. 5 Experimental water adsorption isotherms from the literature and simulated water adsorption isotherms of MOFs by LGCMC. The sky-blue curve is the corrected isotherms using the corrected chemical potential (μ_{cor}) in LGCMC. (a) MOF-74-Mg, (b) $\text{Al}(\text{OH})$ -(1,4-NDC), (c) MUF-77-Zn-methyl, (d) MIL-100-Fe, (e) MIL-101-Cr and (f) DUT-67-Zr.

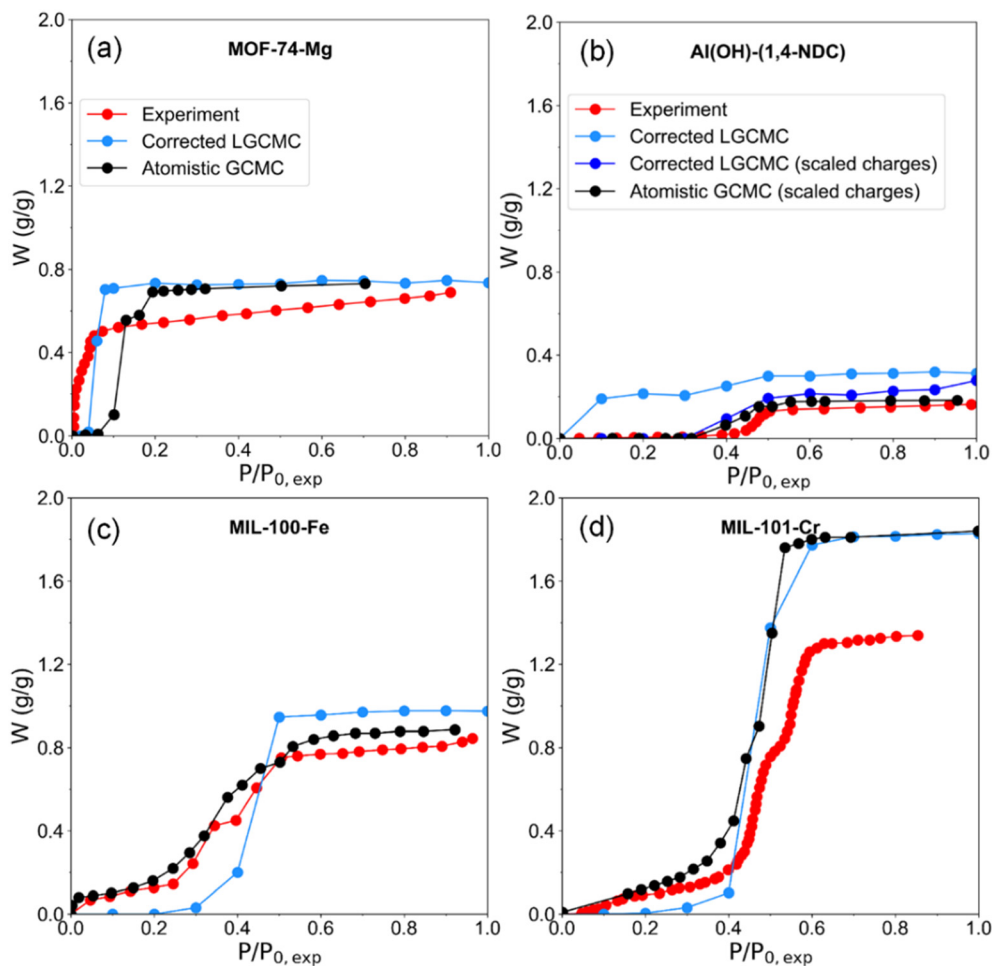


Fig. 6 Experimental and GCMC simulated water adsorption isotherms of MOFs, in which the isotherm by GCMC simulation is extracted from other studies. (a) MOF-74-Mg,²⁶ (b) Al(OH)-(1,4-NDC),³⁵ (c) MIL-100-Fe,⁶⁷ and (d) MIL-101-Cr.³⁰ It should be noted that the isotherms of “corrected LGCMC (scaled charges)” and “atomistic GCMC (scaled charges)”³⁵ of Al(OH)-(1,4-NDC) were obtained using the same force field.

obtained by scaling all partial charges of framework atoms down to about 30% for better reproducing the experimental adsorption isotherm, which in fact weakens the interactions between water and adsorbents.³⁵ Without scaling the atomic charges of Al(OH)-(1,4-NDC) by LGCMC in previously reported work and this work, the water adsorption takes place at low pressure with a small step position owing to the strong interaction with water molecules.³⁵ When using the same force field parameters of Al(OH)-(1,4-NDC) as in the literature,³⁵ LGCMC obtains a consistent adsorption isotherm with atomistic GCMC and experiment (Fig. 6b and S4†). In contrast with GCMC, it has to be emphasized that the computation cost of LGCMC is within a few hours for most MOFs, which is the most attractive aspect of using LGCMC. Overall, the correction strategy of LGCMC simulation is effective in achieving consistent isotherms with the experiment and atomistic GCMC simulation.

Water adsorption behaviors and heat of adsorption

Based on the water adsorption isotherms obtained by LGCMC, the performance of applications focusing on the

water delivery capacity such as water harvesting and desalination can be easily and directly evaluated according to the water uptake of adsorbents under operation conditions. Not only the water uptake but also the corresponding heat of adsorption (H_{ads}) can be obtained from LGCMC simulations (examples in Fig. 7). Thus, the performance of water adsorption-related applications with an energy conversion process, such as adsorption chillers or heat pumps, including cooling/heat capacity and energy conversion efficiency can be predicted.

Besides, users can explore the water adsorption mechanism by LGCMC based on the trajectory of mW molecules in adsorbents. Taking three MOFs with different isotherm types as examples (Fig. 7), the snapshots of water molecule distribution during the adsorption process in MOFs were displayed. As for MOF-74-Mg with a type I isotherm and MUF-77-Zn-methyl with a type V isotherm, water molecules fill their accessible pores instantaneously. The lower pressure of MOF-74-Mg than MUF-77-Zn-methyl resulting in different shapes of adsorption isotherms is mainly attributed to its stronger hydrophilicity or stronger interaction between MOFs



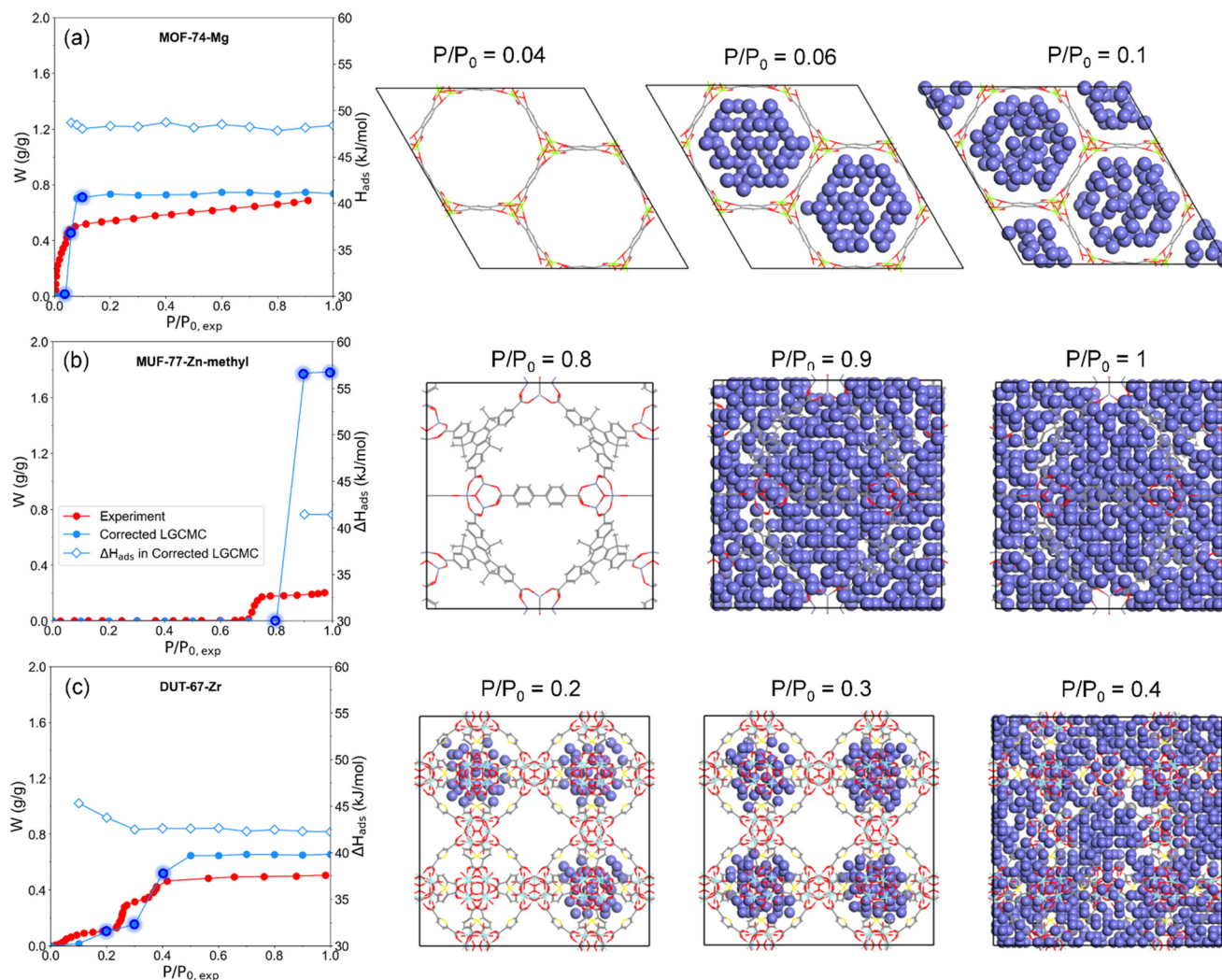


Fig. 7 Water uptake (W), heat of adsorption (H_{ads}) and snapshots of water adsorption in typical MOFs with different-type isotherms obtained from corrected LGCMC. (a) MOF-74-Mg with a type I isotherm (i.e., inverted L shape), (b) MUF-77-Zn-methyl with a type V (i.e., S shape) isotherm, and (c) DUT-67-Zr with multiple adsorption step isotherm.

and water. In DUT-67-Zr with two pore sizes, water molecules preferentially adsorb in the small pores in the form of water clusters at low pressure, and then fill large pores at high pressure. Thus, not only the isotherm with the characteristics of two adsorption steps is reproduced but also the corresponding water adsorption behaviors can be explored.

Conclusions

In this work, we demonstrated lattice GCMC (LGCMC) simulation using a coarse-grained mW model to accurately and rapidly predict water adsorption isotherms of porous adsorbents within a few hours. Water adsorption isotherms of adsorbents obtained by primitive LGCMC under the chemical potential for the mW model (μ_{mW}) cannot accurately predict the step position of isotherms compared to experimental ones. Thus, we proposed an isotherm correction strategy to improve the consistency by correcting the difference in saturation pressure between mW from

LGCMC and experimental water in nanoconfinement, in which the condensation pressure of confined water in each adsorbent was obtained from the Kelvin equation. The difference in saturation pressure was then converted to the chemical potential (μ_{cor}) for the corrected LGCMC simulation. Based on the correction strategy, the water adsorption isotherms obtained by LGCMC exhibit improved consistency with experimental results. It should be noted that the complicated channel and surface heterogeneity of MOFs including various pore sizes, shapes and connectivity add up to the difficulty in accurate prediction of their water adsorption performance. The simplification of the physical properties inside the MOFs during estimation of the contact angle and Kelvin condensation pressure in our correction strategy may cause over- or underestimation of the step positions of water isotherms. For MOFs with open metal sites such as MOF-74-Mg, MIL-100-Fe and MIL-101-Cr, using a more accurate force field from first principles methods may further improve the prediction accuracy. However, there is no



doubt that it is an effective strategy for the prediction of water adsorption isotherms of various adsorbents.

Besides, not only the water uptake but also the corresponding heat of adsorption and the distribution of water molecules in adsorbents can be obtained from LGCMC, which permits the performance evaluation of water adsorption-driven systems, including water delivery capacity and energy conversion efficiency, and enables the in-depth understanding of the microscopic mechanism of water adsorption behaviors in adsorbents. It has to be emphasized that the total computation time of LGCMC can be accomplished within a few hours for most adsorbents, which yields a significant acceleration of two to three orders of magnitude compared to conventional atomistic GCMC simulations. The LGCMC approach reported in this work opens up the possibility for high-throughput computational screening of an enormous number of adsorbents as well as greatly speeds up the discovery of potential adsorbent candidates in water adsorption-related applications.

Data availability

The data supporting this article have been included as part of the ESI.†

Conflicts of interest

There are no conflicts of interest to declare.

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