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Effect of water loading on the stability of pristine and defective UiO-66†

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Materials used for water treatment purposes need to be stable for easy handling and cost-effectiveness. UiO-66 has been identified as a promising option. In this work, we investigate the impact of water loading on the structural and mechanical properties of pristine and defective UiO-66 using classical molecular simulations. We employ and compare two approaches for modeling non-bonded interactions between the framework and water molecules: direct Lorentz–Berthelot (L–B) mixing and hybrid mixing. We conducted molecular dynamics simulations to examine the spatial arrangement of water molecules within the framework, water affinity for specific framework interaction sites, and their impact on the framework's structural parameters under atmospheric conditions, high hydrostatic pressures, and increased water loading. Our results indicate that both methods predict water affinity near zirconium clusters, but differ in identifying principal interaction sites and interaction strength. L–B mixing predicts strong binding to linker oxygen atoms, restricting water movement, while hybrid mixing indicated dynamic water behavior, with site-to-site hopping and pore-to-pore movement observed at moderate and high loadings. Structural analysis at increased water loadings showed adsorption-induced expansion using L–B mixing due to linker–cluster bond stretching, contrasting with slight system contraction predicted by hybrid mixing. High-pressure NPT simulations evidence that water loading reduces amorphization pressure, although values obtained using both approaches differ significantly at moderate and high loadings.

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Introduction

Water pollution is an escalating concern in the world, as diverse sources such as sewage, wastewater, agricultural pesticides, and industrial waste continually introduce harmful substances into natural water bodies.^{1–3} This contamination poses significant threats to ecosystems, the environment, and human health. Recent research has highlighted the potential of metal–organic frameworks (MOFs) in addressing wastewater treatment challenges.^{4–7} MOFs have demonstrated effectiveness in eliminating a variety of pollutants, including phosphates,^{8,9} fluorides,^{10,11} heavy metals,^{12,13} pharmaceuticals,^{14,15} and chemical warfare agents.^{16,17} The limitations of traditional water

harvesting methods, such as their reliance on high humidity and energy inputs, further underscore the need for innovative and efficient approaches like using MOFs.^{18–20}

For industrial applications, it is crucial that materials used in water treatment exhibit structural stability under moist conditions to ensure ease of handling and cost-effectiveness. Despite the development of numerous MOFs, only a select few are chemically stable enough for such applications.^{21–24} Among these, UiO-66, a Zr-based MOF, is particularly noteworthy for its exceptional water stability, which is attributed to the robustness of the Zr–O bond and a unique geometry that minimizes water inclusion and reduces hydrolysis reactions.^{13,21,25–27} UiO-66 features zirconium-based $Zr_6O_4(OH)_4$ clusters interconnected by 1,4-benzenedicarboxylic acid (BDC) ligands. In its pristine, defect-free state, each inorganic cluster is 12-fold coordinated, contributing to its remarkable structural stability under high pressure conditions.^{25,28–32} Functionalized derivatives of UiO-66, developed to tune hydrophobicity, also exhibit impressive stability, further underscoring the potential of Zr-based MOFs in advanced water treatment applications.^{7,17,33–36} Most synthesis procedures for UiO-66 result in a defective structure with missing linkers, typically reducing the coordination number of each zirconium cluster to 11, indicating one missing linker per unit cell on average.^{30,37–40} Defects increase the porosity and surface area, enhancing adsorption behavior,^{31,32,41,42} catalytic

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properties,^{37,43} and proton conductivity.^{44,45} However, defects also reduce structural stability, creating a trade-off between improved adsorption and mechanical integrity.^{38,41,46}

Molecular simulations provide a reliable means to analyze the relationships between the material structure and specific properties, particularly when adsorbates or defects are present. Over the past decade, computational studies have extensively examined the adsorption properties of pristine and defective UiO-66, focusing primarily on water and CO₂ adsorption.^{47–51} These simulations have offered valuable insights for industrial applications such as water harvesting and gas separation, often modeling UiO-66 as a rigid structure interacting with guest molecules solely *via* non-bonded potentials.

To accurately study the structural and mechanical properties of UiO-66, a flexible model of the framework is essential. Flexible models, such as the Rogge *et al.* force fields developed in 2016,⁴⁶ have been extensively validated for predicting the properties of unloaded UiO-66. These models employ *ab initio* derived parameters tailored to each specific structure, constructed through the QuickFF procedure^{52,53} developed by the same group. However, these force fields were primarily validated by reproducing the structural and mechanical properties of evacuated systems, where interactions with adsorbate molecules were not considered. Recent studies^{54,55} have demonstrated that when adsorbates such as acetone and isopropyl alcohol are introduced, the Rogge potential inadequately captures adsorbate–framework interactions, particularly hydrogen bonding with the inorganic clusters. This limitation underscores the need for modifications to the force field to accurately model the interaction between Zr clusters and adsorbates in loaded UiO-66 systems.

The primary objective of this work is to determine how water loading affects the structural and mechanical properties of flexible UiO-66. Given the established reliability of the Rogge force fields in accurately predicting these properties, we prioritized maintaining their original parameters for simulating water-loaded frameworks. Instead of altering these parameters, we assessed and compared two approaches for modeling the interaction between the framework and water molecules: direct Lorentz–Berthelot (L–B) mixing and hybrid mixing. The hybrid mixing method has been previously employed to study the pore filling process and water diffusion within UiO-66,⁵⁶ using separate water force fields to model water–water interactions (MB-pol⁵⁷) and water–framework interactions (TIP4P2005 (ref. 58)). In our study, the hybrid mixing method utilizes the Rogge force field for interactions between framework atoms and the UFF⁵⁹ force field for interactions between the framework and water molecules. By systematically evaluating these approaches, we aim to provide a deeper understanding of how water impacts the stability and performance of UiO-66, with significant implications for its use in industrial water treatment applications.

Methodology

Along with the pristine structure, we considered the eight defective structures (and corresponding force fields) proposed by Rogge *et al.*,⁴⁶ each containing linker defects: one of them

(type 0) corresponds to a system with one linker vacancy per unit cell, while the rest (types 1–7) represent structures in which an additional, distinct linker defect is introduced to the type 0 parent structure (Fig. S1†). Further details on the force fields and the construction of the structures are provided in the first portion of the ESI.† Before the production simulations, these structures were loaded with different amounts of water molecules, from 10 molecules per unit cell of the framework (corresponding to a relative humidity of 5%) to 120 molecules per unit cell (80% RH), near the water saturation point at room temperature. As was done in recent published work,^{47,56,60} the quantity of water molecules within the MOF structure at various relative humidity levels was determined using the experimental water adsorption isotherm for UiO-66.^{51,61}

Force fields and models

We model the pristine and defective frameworks using the group of force fields proposed by Rogge *et al.*⁴⁶ which have proven to provide an accurate prediction of the structural and mechanical properties of the unloaded systems. The functional form for the interactions between framework atoms, regardless of the inclusion of linker vacancies, is the following:

$$U = U_{\text{bonded}} + U_{\text{non-bonded}} \quad (1)$$

where the covalent interactions between neighboring atoms are mediated through harmonic and Fourier-style potentials

$$U_{\text{bonds}} = \sum_{N=1}^{N_{\text{bonds}}} \frac{1}{2} K_{r,n} (r_n - r_{n,0})^2 \quad (2)$$

$$U_{\text{bends}} = \sum_{N=1}^{N_{\text{bends}}} \frac{1}{2} K_{\theta,n} (\theta_n - \theta_{n,0})^2 \quad (3)$$

$$U_{\text{dihedrals}} = \sum_{N=1}^{N_{\text{dihed}}} \frac{1}{2} K_{\phi,n} (1 - \cos(m\phi - \phi_0)) \quad (4)$$

$$U_{\text{oopd}} = \sum_{N=1}^{N_{\text{oopd}}} \frac{1}{2} K_{d,n} (d_n - d_{n,0})^2 \quad (5)$$

Two terms contribute to the non-bonded portion of the potential energy function. The electrostatics are described by the Coulomb interaction between spherical Gaussian densities with distinct radii d_i and d_j containing charges q_i and q_j , respectively. The potential energy term for particles separated by a distance of r_{ij} is:

$$U_{\text{Electrostatic}} = \frac{1}{2} \sum_{\substack{i,j=1 \\ i \neq j}}^{N_{\text{dihed}}} \frac{q_i q_j}{4\pi\epsilon_0 r_{ij}} \operatorname{erf} \left(\frac{r_{ij}}{\sqrt{d_i^2 + d_j^2}} \right) \quad (6)$$

The second term models the van der Waals interactions between two atoms i and j separated by a distance r_{ij} using the two-parameter molecular mechanics (MM3)^{62,63} Buckingham potential:



$$U_{\text{vdw}} = \varepsilon_{ij} \left[1.84 \times 10^5 \exp\left(-12 \frac{r_{ij}}{\sigma_{ij}}\right) - 2.25 \left(\frac{\sigma_{ij}}{r_{ij}}\right) \right] \quad (7)$$

For bonded atoms, different scaling and exclusion rules are applied for each contribution to the non-bonded part of the potential energy. As defined by the MM3 force field, 1–2 and 1–3 exclusion rules for bonded pairs are applied to VdW interactions, with no scaling applied to atoms separated by two or more atoms (1–4). For framework atoms of different species, the equilibrium distance (σ_{ij}) and well depth (ε_{ij}) parameters are determined *via* the empirical Lorentz–Berthelot mixing rules:

$$\sigma_{ij} = \frac{1}{2}(\sigma_i + \sigma_j), \quad \varepsilon_{ij} = \sqrt{\varepsilon_i \varepsilon_j} \quad (8)$$

The interaction parameters of the energy equation depend on the particular structure to be modelled, that is, each of the 9 structures simulated (pristine, type 0 defective and type 1–7 defective) has distinct associated sets of parameters, and they are provided in the original publication of the force field.⁴⁶

The water molecules were modeled using the extended SPC model,^{64,65} although the Tip4P2005 (ref. 58) force field was initially implemented and considered as well. The SPC/E model was selected primarily due to its lower computational cost, as it involves taking into account fewer interactions, and because early results showed that both models influenced the framework similarly. The non-bonded parameters for the description of water–water interactions were derived using Lorentz–Berthelot mixing rules as well.

Two distinct methods for modeling the interaction between water molecules and the framework atoms were employed; in both cases the VdW interactions are modelled using the Lennard-Jones potential and the electrostatics are modelled using the canonical Coulomb potential for point charges. In the first method, the Lennard-Jones parameters for the description of the water–framework interaction were derived by applying Lorentz–Berthelot (L–B) mixing rules directly. Here, it should be noted that although the empirical combination of L–J parameters from popular rigid water models and generic force fields have been used extensively in published work to study this and similar systems, this is not the case for the combination of SPC/E and MM3 parameters. This first mixing methodology is thus implemented as an intuitive point of reference, reflective of the usual default choice in the simulation community. In the second method, the Lennard-Jones contribution of the water–framework interaction was modified by employing universal force field (UFF)⁵⁹ ε and σ coefficients for the framework atoms instead of the MM3 force field (MM3)^{62,63} parameters used for intra-framework non-bonded interactions, therefore making use of two framework force fields in a hybrid manner.

Simulation details

The pressure-volume behavior of UiO-66 structures was determined using molecular dynamics (MD) simulations in the flexible NPT ensemble ($N, P, \sigma_\alpha = 0, T$). This ensemble allows

dynamic changes in both simulation cell volume and shape, enabling the estimation of the amorphization pressure (P_{am}) of the framework. For each system, flexible NPT simulations were conducted starting from a high pressure. The unloaded structures were first energy minimized and equilibrated under atmospheric conditions to record the reference volume (V_r). After loading with water molecules, the structures were equilibrated at high pressure, monitoring the instantaneous volume (V_i). Amorphization was deemed to occur if V_i dropped below a predefined threshold (V_t), proportional to V_r . The pressure was adjusted in subsequent simulations based on whether amorphization occurred, using a step size (ΔP) of 0.005 GPa. The final P_{am} was determined as the highest pressure at which the structure remained stable. Large simulation cells were used to mitigate fluctuations in instantaneous pressure, which can induce phase transitions at artificially low pressures, particularly in flexible MOFs.^{66,67}

All simulations were performed using the LAMMPS package.⁶⁸ The Verlet integrator with a timestep of 1 fs ensured energy conservation, while independent thermostats optimized equilibration times by grouping atoms of the framework and water. A Nose–Hoover thermostat (300 K, 100 fs relaxation time) and an MTTK barostat (1000 fs relaxation time) controlled temperature and pressure, respectively. Radial distribution functions (RDFs) and 3D water density distribution maps⁶⁹ were averaged from 500 ps NVT ensemble runs after equilibration. Simulations used a $2 \times 2 \times 2$ supercell with non-bonded interactions truncated at 14.0 Å and long-range electrostatics calculated *via* Ewald summation. Periodic boundary conditions were applied in all dimensions, with specific scaling of 1–2, 1–3, and 1–4 non-bonded interactions as per the force field requirements.

Results and discussion

Effect of water loading on the pristine structures

In order to evaluate the performance of each parameter mixing method, we start by analyzing the spatial arrangement of water molecules inside the undefective framework at different water loadings. The Radial Distribution Function (RDF) describes how the average density varies as a function of distance from a chosen reference particle. By selecting framework atoms as references, they can be used to identify preferential adsorption sites within its surface and characterize the interaction environment in the vicinity of the metal clusters and organic ligands.

Regarding the initial interaction sites, at low loadings both models predict that water molecules tend to agglomerate around the metal clusters while keeping away from the central portion of the organic ligands, in line with previously published research.^{47,56,70} However, each model suggests different species near the metal cluster as the primary interaction site for water. L–B mixing predicts that the oxygen atoms connecting the Zr atoms with the linkers are the main interaction site, whereas hybrid mixing points to the hydroxyl groups within the inorganic cluster.



For the L-B mixing simulations at low loadings, the RDF describes the spatial correlation between O_{H_2O} and O_L peaks at a radial distance of 1.9 Å, contrasting with a clear separation from O_{Ox} and O_{Oh} sites (Fig. S2a†). This behavior is not consistent with previous computational studies indicating that water preferentially binds to the hydroxyl hydrogens in the zirconium clusters. Recently published computational studies of water diffusion and pore filling processes in UiO-66 predict an interaction site affinity order of $O_{Oh} > O_L > O_{Ox}$.⁵⁶ Furthermore, the peak distance indicates heavy perturbation of the Zr- O_L bond electronic environment. While the interaction site water affinity order predicted by hybrid mixing (Fig. 1) is qualitatively congruent with said findings and the radial distance to O_{Oh} is the same (≈ 2.8 Å) the density of water molecules near the preferential site at low loadings is underestimated by 36%. The RDFs between water oxygen atoms (O_{H_2O}) and potential bonding sites were analyzed at higher loadings of 40 and 100 water molecules per unit cell using both mixing methods (Fig. S2†). Hybrid mixing predicts a broadening of RDF peaks with increasing water content, indicating a more dispersed distribution of water molecules. As loading increases, more water molecules shift away from the zirconium clusters toward the pore centers, forming one-dimensional water chains and clusters (Fig. S3†). In contrast, L-B mixing shows a slight decrease in water presence around zirconium clusters, with less pronounced occupancy within the pore volume.

A series of 3 ns simulations in the NVT ensemble were conducted at increasing loadings to monitor water molecule positions and analyze their trajectories. Fig. S4† compares the mean square displacement (MSD) of water molecules for the two interaction methods at a moderate loading of 40 water molecules per unit cell. As shown in Fig. 2, L-B mixing results in water molecules remaining highly localized near the linker oxygen, with minimal dimer formation. In contrast, hybrid mixing simulations exhibit diffusive behavior around the predicted adsorption sites, with occasional formation of water dimers.

To assess how differences in predicted adsorption sites affect the framework structure, NPT simulations were performed under atmospheric conditions with increased water loadings. System volume and bond distances between inner-cluster atoms were monitored and averaged. As shown in Fig. 3, L-B mixing predicts a significant increase in the unit cell lattice parameter with loading, reaching a 7% volume increase at 80 molecules per unit cell compared to the unloaded structure. In contrast, hybrid mixing simulations indicate a slight decrease in lattice parameters, suggesting water-induced framework contraction.

The expansion correlates with stretching of Zr- O_L bonds. In L-B mixing, increased water presence near O_L results in noticeable elongation of Zr- O_L bonds, while hybrid simulations show a more pronounced effect on O_{Oh} -H bonds. Non-preferential site bonds, such as Zr- O_x and Zr- O_h , experience minimal changes in L-B mixing (Fig. S5†). The observed weakening of framework bonds, particularly Zr- O_L , suggests a reduction in cluster-linker coordination and a decrease in mechanical stability upon water loading.

The UiO-66 crystal structure is known for its stability under high pressure, remaining intact up to approximately 1.4 GPa.³³ To assess the impact of water loading on structural stability, NPT simulations were conducted under varying hydrostatic pressures and water loadings. Fig. 4 shows that the amorphization pressure (P_{am}) decreases with increasing water loading. At low loadings (10–20 H_2O molecules per unit cell), P_{am} values are similar for both models. However, at intermediate loadings (30–40 H_2O molecules), significant differences emerge, due to the hybrid model predicting water dimer and cluster formation at lower water concentrations than the L-B model. The water groupings tend to occupy space closer to the middle of the pores and away from the metal clusters, thus contributing less to cluster-linker weakening. L-B mixing likely predicts unrealistically low P_{am} values, and the use of hybrid mixing is suggested at moderate and low loadings.

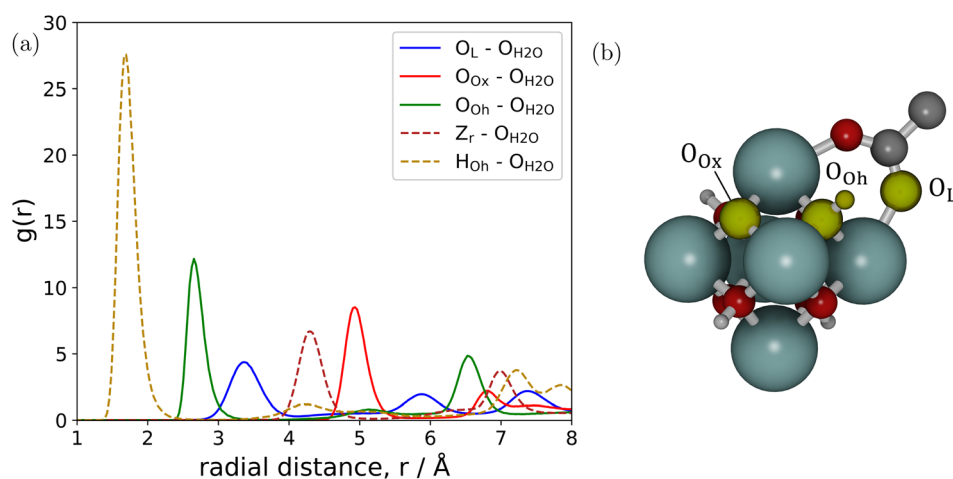


Fig. 1 (a) RDF calculated using hybrid mixing between water molecules and framework interaction sites near the zirconium (Zr) cluster. Obtained at a temperature and pressure of 300 K and 1 atm, respectively, with a water loading of 10 molecules per unit cell. (b) The identified interaction sites in the inorganic cluster are the linker oxygen O_L , the hydroxyl group O_{Oh} and the dehydroxylated group O_{Ox} within the cluster.



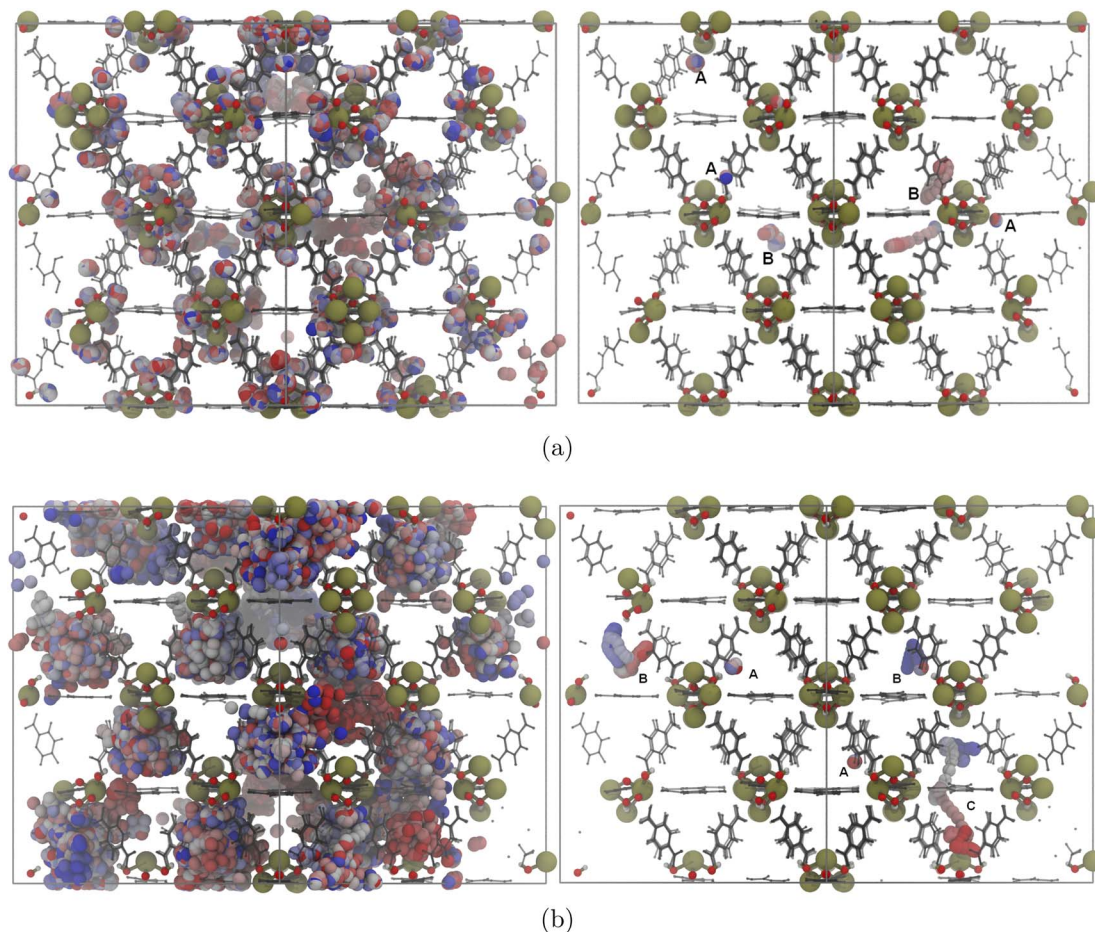


Fig. 2 Trajectories described by water molecules inside the framework over a simulation period of 500 picoseconds at moderate loading (40 molecules per uc) using (a) L–B and (b) hybrid mixing under atmospheric conditions. On the left side, the cumulative trajectories of all O(H₂O) atoms are pictured. On the right side a subset of trajectories is presented, representative of 3 types of observed behavior: (A) molecules bound to an interaction site, (B) molecules orbiting already occupied sites, and (C) molecules of the previous type hopping from one interaction site to another within the same pore. Trajectories are colored based on a gradient from red (start of the trajectory) to blue (end of the trajectory).

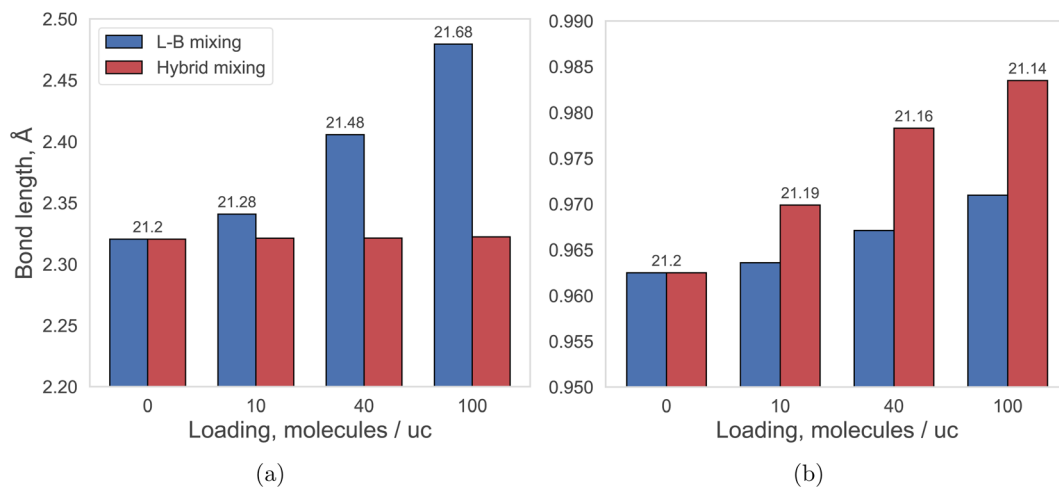


Fig. 3 Average bond lengths between (a) Zr and O_L and (b) O_{OH} and its bonded hydrogen. The unit cell lattice parameter is indicated above each bar. The data for (a) and (b) at a given loading are from the same four simulation sets.



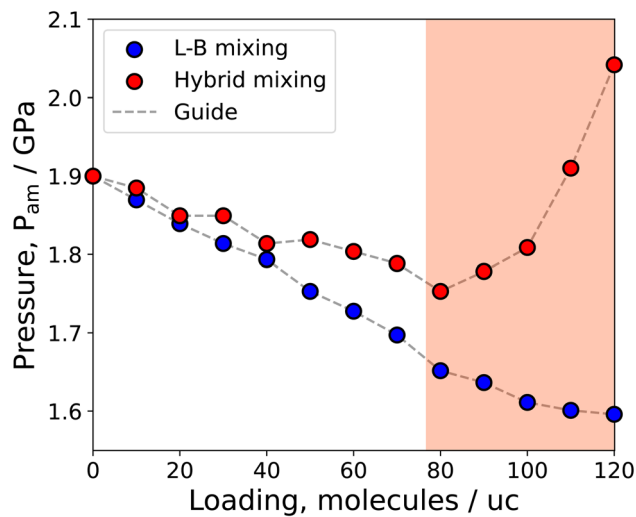


Fig. 4 Amorphization pressure (P_{am}) with increased water loading at 300 K. Both models show similar P_{am} values at low loadings but diverge at higher loadings. Hybrid mixing predicts a higher P_{am} at high loadings, reflecting methodological limitations.

At high loadings (>80 molecules per uc), simulations using hybrid mixing exhibit a noticeable increase in P_{am} values, a behavior absent in L–B mixing simulations. This difference stems from the water distribution within the crystal and highlights a limitation in our framework collapse detection methodology, in which a framework collapse is defined by a system volume drop below a threshold during equilibration. At high particle loadings, the collapse may be impeded or masked, preventing detection using our criteria.

Our findings indicate that in a system as complex as UiO-66 and water, the selection of guest–host interaction parameters significantly influences the distribution of the former within the available pore volume. When comparing the employed

methodologies, hybrid mixing predicts water molecule behavior more in line with previously published simulation studies on the diffusion and pore-filling process of different adsorbates in UiO-66 and analogous structures.^{47,54–56} The results obtained when employing direct Lorentz–Berthelot mixing rules predict an unusually strong coordination between the water molecules and the framework oxygen atoms connecting the organic linkers with the zirconium cluster, heavily restricting the movement of individual water molecules and inhibiting the formation of water clusters at moderate loadings. Moreover, in both cases, the results indicate that the particular spatial arrangement of the water molecules at different loadings influences both inner-cluster and framework structural parameters. This is more evident when employing L–B mixing, as water coordination causes a stretching of the linker–cluster bonds and a subsequent increase in system volume. This mechanical response to loading also impacts loss of crystallinity pressures and is uncharacteristic of a MOF as rigid as UiO-66, and it might be evidence of poor cross-interaction parameter selection.

Effect of water loading on defective structures

Previous simulation studies show that although the inclusion of defects corresponds to a negligible decrease in equilibrium volume at a particular temperature, they introduce substantial decreases in both amorphization pressure and bulk moduli.^{46,71} The effect of water loading at high pressure was investigated for structures containing one and two linker defects per unit cell. NPT simulations were conducted at 300 K, systematically varying the water loading from 10 molecules per unit cell to 80. The resulting P_{am} data obtained using both methods are visually presented in Fig. 5.

A decrease in amorphization pressure is predicted upon increasing water inclusion for all defective structures, using

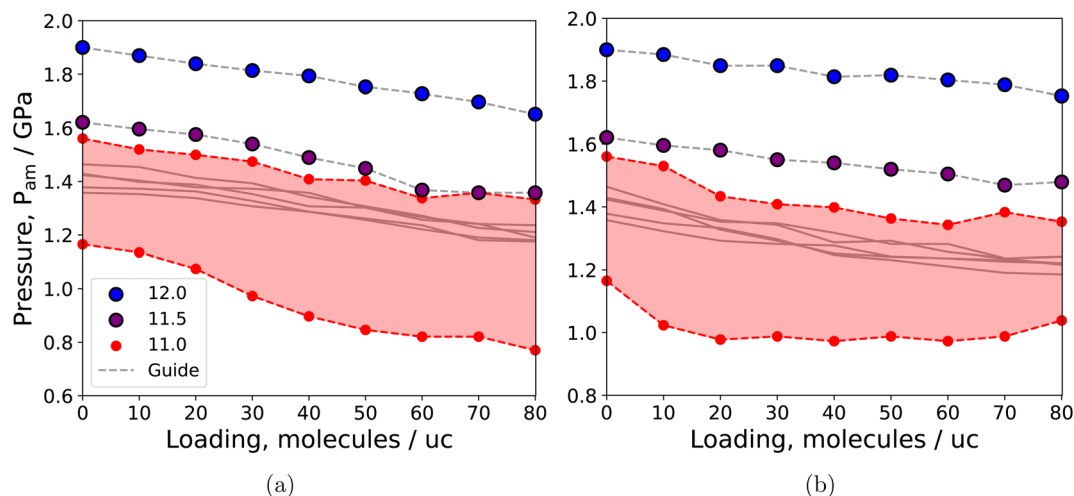


Fig. 5 Amorphization pressure predicted with increased water loading for structures containing linker defects using (a) L–B mixing and (b) hybrid mixing. The color of the curve indicates the average linker–cluster coordination per unit cell. The area in red indicates the P_{am} range of values obtained for structures with an average cluster coordination of 11.0, where the limiting upper and lower red curves indicate the most stable (type 3) and least stable (type 5) structures. The results for the pristine structures (blue) are included for comparison.



both parameter mixing methods. In terms of discerning the relative stability of 11-fold coordination structures under increasing water loading, both methodologies predict that inclusion of type 3 vacancies yields the most stable configuration, while type 5 defects exhibit the least stability to high hydrostatic pressure. A close inspection of the results obtained for the structure with 1 missing linker per unit cell (Fig. S6a†) reveals a similar comparative behavior to the pristine structures, where the amorphization values predicted at low loadings using both models are almost identical, followed by deviations at middle and high loadings. Interestingly, a different trend is observed for the 11-fold coordinated structures (Fig. S6b–d†), where the hybrid mixing simulations predict a sharper decrease in P_{am} values at low loadings when compared to the L–B mixing results. Furthermore, unlike with the less-defective structures, both methods predict comparable high-loading values for all structures with two linker vacancies per unit cell.

Fig. 6 depicts the RDF of water molecules with respect to the metal sites in a structure containing type 3 defects at low loading. For this structure as well as all others containing two linker vacancies per unit cell, when hybrid mixing is employed, water molecules preferentially interact with open Zr atoms and thus tend to agglomerate around them, evidencing strong $\text{O}_{\text{H}_2\text{O}}-\text{Zr}$ interactions between them. Furthermore, water molecules bonded to the open metal sites exhibit reduced translational and rotational movement, similarly to water molecules near the O_{L} atoms when using L–B mixing in the pristine structures. Water molecules near defective inorganic clusters, but not directly bonded to Zr, frequently change hydrogen-bond partners and exhibit higher orientational mobility. As loading increases water begins to occupy the pore regions, with molecules near the center displaying less restricted movement compared to those near interaction sites.

As was done for the pristine framework, the structural properties of the defective systems under atmospheric conditions, using both mixing methods, were also analyzed. Fig. 7a illustrates the change in unit cell volume for pristine, type 0 defective, and type 3 defective structures when using hybrid mixing. Although all three systems generally contract upon loading, the defective structures show a slight expansion at low loadings. This predicted volume expansion is more noticeable in structures with more linker vacancies, as seen when comparing the results for type 0 and type 3. The type 3 defective structure, representative of other structures with two linker defects per unit cell, exhibits a linear increase in system volume followed by contraction once a loading of 20 molecules per unit cell is reached. Comparing the trends for the type 0 and type 3 structures with that for the pristine one, the initial increase in system volume is likely related to the number of open metal sites available after linker vacancies are introduced. When employing hybrid mixing, water molecules bound to open metal sites contribute to the stretching of linker–cluster bonds, and once these sites are occupied, the general trend of volume contraction resumes.

For all three structures and over the loading range, the simulations employing hybrid mixing predict an overall gradual reduction in volume, eventually plateauing for loading values close to saturation. As was the case for the pristine structure, direct Lorentz–Berthelot mixing predicts a significant volume expansion (a 7% to 8% increase over the loading range) of framework volume upon water loading for defective systems (Fig. 7b). Although the results obtained using hybrid mixing are more in line with the expected behavior of UiO-66 and its employment over direct L–B mixing is recommended (comparisons in Fig. S13–S15†), further force field refinement might be necessary to more accurately capture the interactions between water molecules and open metal sites in the defective structures.

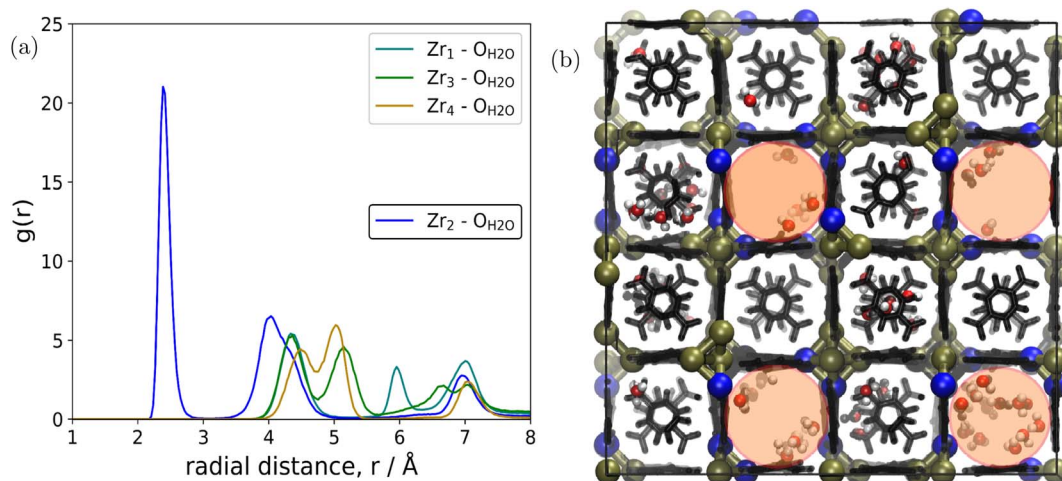


Fig. 6 (a) RDF between water molecules and the zirconium atoms for the structure containing type 3 defects. The blue curve depicts the RDF between H_2O and open Zr_2 sites, where all others show the average density of water with respect to fully coordinated Zr sites. (b) Representative snapshot along the XY plane of the structure containing type 3 defects. Open metal sites are pictured in blue. 1D channels created by linker vacancies are highlighted in orange. Similar water molecule behavior observed in simulations with other defective structures at low loading, using hybrid mixing (Fig. S7–S12†).



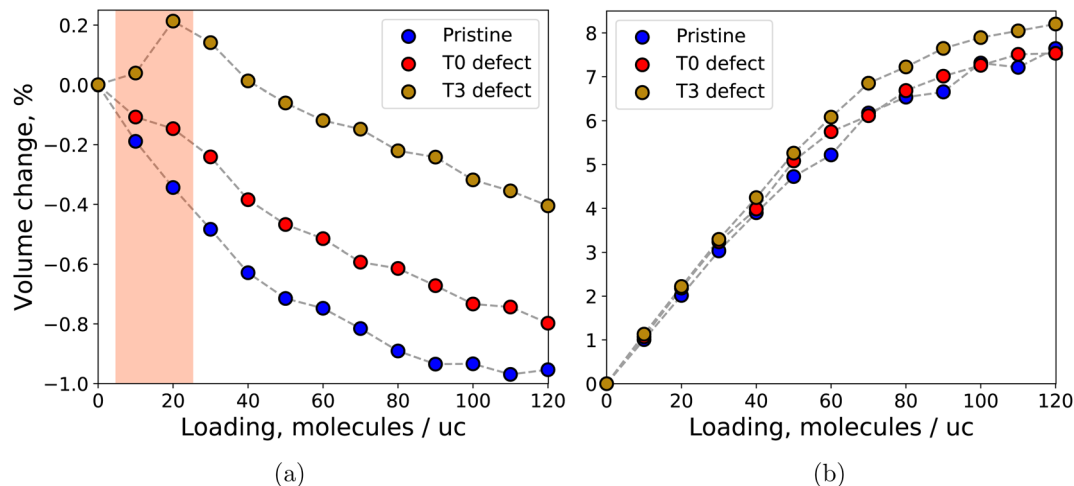


Fig. 7 System volume change upon water loading under atmospheric conditions, using (a) hybrid mixing and (b) L–B mixing. The results for the pristine, type 0 (1 defect per uc) and type 3 (2 defects per uc) structures are presented. Unlike pristine behavior upon water loading, a slight increase in framework volume is observed at low loadings for defective structures when using hybrid mixing (highlighted in orange).

Conclusions

This work investigated the effect of water loading on the stability of pristine and defective UiO-66 *via* molecular simulations. We used two methodologies for modeling non-bonded interactions: direct Lorentz–Berthelot (L–B) mixing and hybrid mixing. L–B mixing combines Lennard-Jones parameters for framework–framework and water–water interactions, while hybrid mixing employs Universal Force Field (UFF) parameters for the framework. We examined water distribution within the framework under atmospheric conditions. Both methodologies predicted water clustering around zirconium clusters, but RDF analysis showed differences in interaction sites and strengths. L–B mixing suggests strong binding of water to linker oxygen atoms (O_L), restricting movement and diffusion. Hybrid mixing, however, shows interactions with the hydroxyl groups of the inorganic cluster (O_{OH}), resulting in dimer formation, site-to-site hopping at low loadings, and pore-to-pore diffusion at higher loadings, which aligns with recent studies on water diffusion in UiO-66.

NPT simulations under atmospheric conditions demonstrated that interaction strength significantly influences structural parameters. At high water loadings, hybrid mixing predicted slight contraction of the system, whereas L–B mixing indicated over a 7 percent expansion due to stretching of linker–cluster bonds. This suggests that L–B mixing inadequately models the rigidity of UiO-66. High-pressure NPT simulations showed that water loading decreases amorphization pressure for both methodologies compared to the unloaded structure. L–B mixing consistently predicted lower amorphization pressures, with notable deviations at middle and high loadings, consistent with previous studies where water agglomeration reduces linker–cluster bond weakening. Similar trends were observed for structures with one missing linker per unit cell. For 11-fold coordinated structures, hybrid mixing predicted a sharper decrease in amorphization

pressure at low loadings compared to L–B mixing, though both methods converged at high loadings for structures with two linker vacancies per unit cell.

For structures with two linker vacancies, hybrid mixing simulations showed water molecules preferentially interacting with open Zr atoms, leading to agglomeration and strong O_{H_2O} –Zr interactions. These water molecules exhibited reduced movement, similar to those near O_L atoms in L–B mixing. Initial volume expansion was observed at low loadings due to interactions with open metal sites, which then resumed contraction once these sites were filled. This indicates a need for refining force field parameters to better model interactions with open Zr sites in defective UiO-66. Overall, our findings emphasize the significant impact of water loading on UiO-66's structural stability and the importance of selecting appropriate interaction parameters for accurate simulations. The differences between L–B and hybrid mixing highlight the need for careful parameter selection and further development of force fields for defective frameworks.

Data availability

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

Conflicts of interest

There are no conflicts to declare.

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