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## Selectivity for water isotopologues within metal organic nanotubes†

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Through a combination of many analytical approaches, we show that a metal organic nanotube (UMON) displays selectivity for  $\text{H}_2\text{O}$  over all types of heavy water ( $\text{D}_2\text{O}$ ,  $\text{HDO}$ ,  $\text{HTO}$ ). Water adsorption experiments combined with vibrational and radiochemical analyses reveal significant differences in uptake and suggest that surface adsorption processes may be a key driver in water uptake for this material.

While the mass differences between isotopologues of water ( $\text{H}_2\text{O}$ ,  $\text{H}^2\text{DO}$ ,  $\text{D}_2\text{O}$ ,  $\text{H}^3\text{TO}$ ) are small, the overall isotopic effects can be significant and result in important differences in physical and chemical properties. The natural abundance water consists of mostly  $\text{H}_2\text{O}$  with one deuterium ( ${}^2\text{D}$ ) or tritium ( ${}^3\text{T}$ ) atom for every  $6.42 \times 10^3$  or  $5.41 \times 10^{16}$  H atoms, respectively.<sup>1</sup> Heavy water enriched in  ${}^2\text{D}$  and/or  ${}^3\text{T}$  forms stronger hydrogen bonds, which causes smaller amplitude intermolecular motions and more ordering within the solution phase.<sup>2</sup> This effect imparts differences in the physical properties of light and heavy water, including variations in the viscosity,<sup>3</sup> self-diffusion coefficient,<sup>4</sup> phase behavior,<sup>5</sup> hydration energies,<sup>6</sup> and exchange kinetics<sup>7</sup> that have observable effects on protein stability,<sup>8</sup> biological activity,<sup>7</sup> toxicity,<sup>9</sup> and chemical reactivity.<sup>7</sup> Nuclear quantum effects of water isotopologues are an active area of research and the effects on kinetics and thermodynamics in complex systems are critical if difficult to untangle.<sup>10</sup>

Even with the substantial differences in the properties of light and heavy water, selectively partitioning these isotopologues remains both challenging and important. Evaporation in the environment leads to slight fractional separation in surface waters where variation of  ${}^2\text{D}$  concentrations ranging from 0.0130% in the Arctic ocean to 0.0162% in the Nile River, and these differences can be used for climatic reconstructions.<sup>11</sup> In a more anthropogenic context, isotopic fractionation is also important for heavy water nuclear reactors. Although distillation is the simplest engineering procedure for separation of the isotopologues, a separation factor of 1.015 requires extensive multi-stage systems and large energy costs (10 MW h/kg  $\text{D}_2\text{O}$ ).<sup>11,12</sup> Alternatively, chemical processes for heavy water production, such as the Girdler sulfide and monothermal ammonium hydrogen methodologies, offer a marginal increase in the separation factors

(2.3 to 2.8) with similar energy demands.<sup>13</sup> Clearly, a fractionation process for isotopic separation of water that involves modest energy consumption and high separation factors would offer substantial benefits. So, it is essential to identify materials capable of such large separation factors and to understand the molecular mechanisms by which these processes occur.

In previous studies we showed that a metal organic nanotube (UMON) contains 1.2 nm diameter nanopores composed of six U(vi) nodes connected through iminodiacetate linkers to create macrocyclic units that stack into nanotubular arrays by hydrogen bonding (Fig. 1). The crystalline material is stabilized by charge-balancing piperazinium cations and is durable to 200 °C.<sup>14</sup> Within the nanotubular cavity, water molecules spontaneously arrange in hexagonal arrays resembling crystalline ice that can be removed without collapse or structural degradation of the nanotube. Dehydration of the weakly-bound, confined water within the UMON nanochannels occurs between 40–70 °C and complete removal of the  $\text{H}_2\text{O}$  accounts for 5.87%

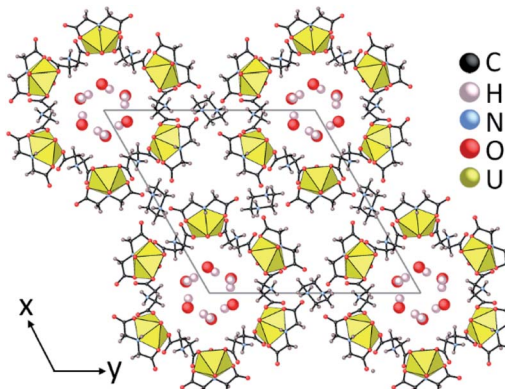


Fig. 1 (a) UMON crystals are composed of oriented metal organic nanotube, with U(vi) (yellow polyhedra) linked by iminodiacetate molecules form the nanotubular arrays and ordered  $\text{H}_2\text{O}$  molecules are located within the nanotubes.

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weight loss in thermogravimetric analysis (TGA).<sup>14</sup> Rehydration is spontaneous under humid conditions. Following the loss of water from the nanochannel, an additional 1.2% weight gain can be observed under high humidity conditions due to adsorption of surface water onto the material.<sup>15</sup> The UMON surface contains ample hydrophilic groups (*i.e.* free carboxyl oxygen atoms and protonated amines) that are available for hydrogen bonding with water molecules to facilitate this additional surface adsorption. Removal of the more strongly bound surface water takes place between 80–150 °C. Here we show that, while UMON allows H<sub>2</sub>O to enter its nanochannels, we observe divergent behaviour for water containing <sup>3</sup>T or <sup>2</sup>D. This fractionation process occurs at room-temperature with no additional energy sources or pressurization. We have employed a range of analytical methods to confirm the isotope effects observed for this material including spectroscopic and radiochemical techniques.

To investigate the isotopic fractionation of UMON, we synthesized the material according to the previously reported procedure.<sup>14</sup> We initially examined the UMON behaviour in deuterated systems (D<sub>2</sub>O/HDO) using both flow-through and batch studies. *In situ* flow-through experiments (also see ESI† for details) using a custom-built vapor sorption instrument that was been previously described and validated for reproducibility and reliability. For the batch experiments, dehydrated UMON samples were exposed to D<sub>2</sub>O in a N<sub>2</sub> gas filled glove bag to prevent contamination by H<sub>2</sub>O (described in detail in the ESI†). After 18 hours, the samples were analysed by a TA instruments thermogravimetric analyser (TGA) equipped with a Thermo-Scientific FTIR spectroscopy for the evolved gases to identify the isotopic composition of the adsorbate. A second set of dehydrated UMON were used as controls to confirm that H<sub>2</sub>O was not present in the hydration chamber or introduced as part of the developed procedure. No weight loss or evolved vapor of any kind was observed for the control samples throughout the experiments, which validated the experimental design of the batch experiments as free from protic contamination.

Flow-through experiments demonstrate difference in uptake kinetics and total uptake in the system (Fig. 2). Full uptake (5.8 ± 0.2 wt%) was observed in the H<sub>2</sub>O flow-through experiments (80% RH) whereas only 1.2 ± 0.2 wt% uptake occurs in the presence of

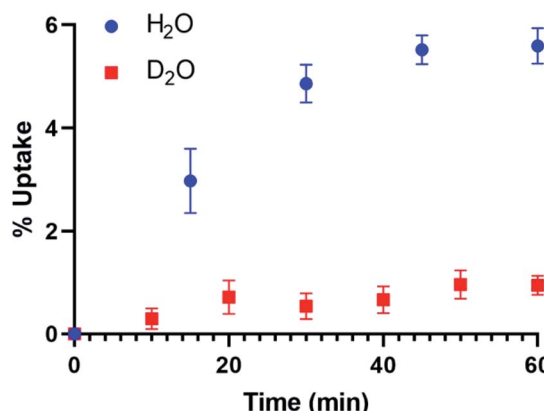


Fig. 2 Uptake of H<sub>2</sub>O and D<sub>2</sub>O vapor into the UMON material using the flow-through apparatus.

D<sub>2</sub>O over the course of the 60 minute experiment. Additional TGA analysis of the UMON sample exposed to H<sub>2</sub>O finds a 5.1 ± 0.5% weight loss between 25 to 100 °C. A similar residual weight is observed for the material exposed to D<sub>2</sub>O, where 0.6 ± 0.5% mass loss occurs between 25 to 100 °C. Result from the flow-through experiments demonstrate differences in uptake between D<sub>2</sub>O and H<sub>2</sub>O and the related weight loss data finds that residual water (0.6–0.7 wt%) remains strongly bound to the material.

TGA of the dehydrated metal organic nanotubes exposed to D<sub>2</sub>O vapor (99.96% purity) in the batch experiments indicates a 6.1 ± 0.5% weight loss occurring between 25 and 100 °C, but the FTIR spectrum of the evolved gases only provides evidence for the presence of H<sub>2</sub>O in the channels (Fig. 3A). The observed

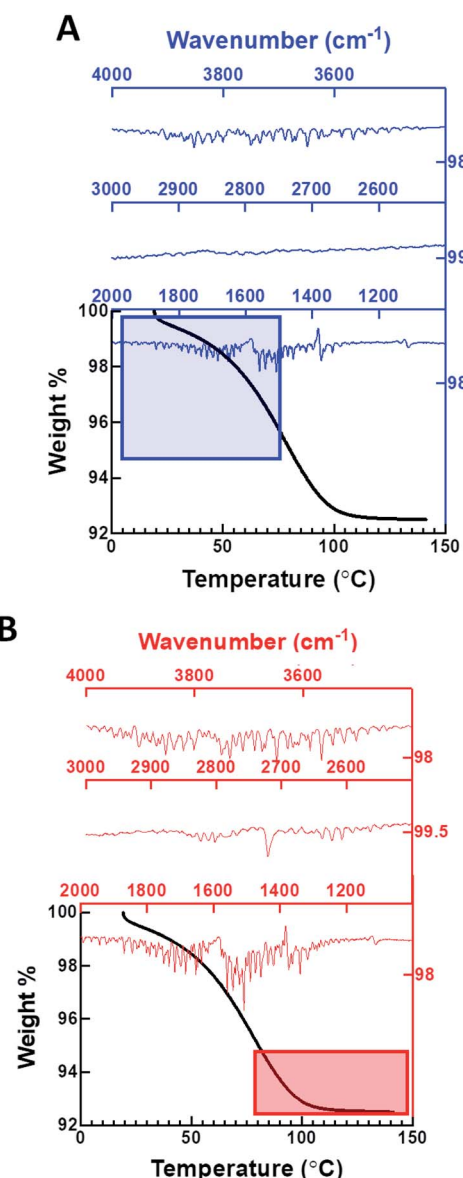


Fig. 3 Thermogravimetric analysis combined with FTIR spectroscopy of the evolved gases determine the presence of water associated with the UMON sample exposed to pure D<sub>2</sub>O vapor. (A) Initially gases evolved during the heating process are confirmed as H<sub>2</sub>O. (B) Evolved gases observed at higher temperatures are identified as HDO by the appearance of a band at 2721 cm<sup>-1</sup>.



weight loss is slightly higher than expected for water confined only within the nanotubes (5.8%); thus, we assume that the additional water must correspond to surface adsorbed species. Evolved water vapor released during the heating process is evaluated using a coupled FTIR spectrometer to determine the isotopic composition. Initially, only O-H stretching ( $\nu_1$  and  $\nu_3$ ; 3400–4000 cm<sup>-1</sup>) and bending modes ( $\nu_2$ ; 1250–1900 cm<sup>-1</sup>) are present in the spectra for the first 5.8% of the total weight loss, indicating that the vapor removed during the initial heating process is H<sub>2</sub>O. The presence of D<sub>2</sub>O or HDO vapor would exhibit characteristic signals at 2671 or 2723 cm<sup>-1</sup>, respectively. These bands are noticeably absent in the initial heating stage, but a weak signal at 2721 cm<sup>-1</sup> does arise during the final 5.8–6.1% portion of the total weight loss (>80 °C) (Fig. 3B). We also conducted an extended vapor sorption experiment (6 hours) and found identical uptake (6.1%), mass loss (5.8%), and no evidence of D<sub>2</sub>O in the FTIR of the evolved gas (Fig. S17†). These data suggest that H<sub>2</sub>O is present inside the nanochannel and additional heating results in the removal of surface adsorbed D<sub>2</sub>O/HDO species.

Our previous calorimetric study supports the idea that the water in the channels desorbs before the surface waters because the interaction between the confined water and the interior walls of the nanotube is weak ( $-7.8 \pm 0.9$  kJ mol<sup>-1</sup>) whereas multiple, strong H-bonding interactions on the surface increases the binding energy, and therefore the temperature required to remove the water.<sup>17</sup> In addition, water adsorption calorimetry experience noted two different types of water environments and related adsorption energies. Initially, we described this as two different water sites within the channel, but an alternative explanation is that the first value ( $-59.8 \pm 1.1$  kJ mol<sup>-1</sup>) corresponds to interactions with the UMON surface. After initial strong adsorption to the surface (0.25 mol H<sub>2</sub>O = 0.7 wt%), filling begins, and the differential enthalpy increases sharply to 50 kJ mol<sup>-1</sup> and remains at this value until complete hydration.<sup>17</sup>

One puzzling aspect of the batch experiments is the full uptake of H<sub>2</sub>O in the presence of saturated D<sub>2</sub>O environment. We further characterized the dehydrated UMON samples exposed in the batch reactor (99.96% D<sub>2</sub>O vapor) by FTIR analysis and solid-state 2D NMR measurements. The infrared spectrum of dehydrated UMON exposed to D<sub>2</sub>O (Fig. S18†) shows the broad O-H stretching band ( $\nu_1$  and  $\nu_3$ , 3400–4000 cm<sup>-1</sup>) and an additional band at 2280 cm<sup>-1</sup> that does not correspond to any HDO or D<sub>2</sub>O spectral modes. Instead, we propose that this band is due to hydrogen/deuterium exchange with  $-\text{NH}_2$  functional groups within iminodiacetate or piperazinium cations. This spectral interpretation agrees with a study by Heacock and Marion showing that deuteration of secondary amines results in bands between 2000–2400 cm<sup>-1</sup>.<sup>18</sup> The solid-state 2D-NMR confirms H/D exchange on the UMON surface (Fig. S21 and S22†). UMON hydrated with H<sub>2</sub>O serves as the control sample and does not produce a detectable 2D-NMR resonance, as expected. In contrast, the spectrum collected from the dehydrated UMON sample exposed to D<sub>2</sub>O vapor indicates the presence of a relatively immobile D species. The fit to the data shows that the major resonance (>99%) has a chemical shift of 3.8 ppm with respect to TMS. This shift is

lower in frequency than bulk water (4.8 ppm) and is consistent with a change in the chemical environment of the D species that was detected in the UMON material. This data also provides evidence of H/D exchange, likely on the UMON surface, that produces H<sub>2</sub>O that subsequently enters the channels even in the presence of 99.96% pure D<sub>2</sub>O. We hypothesize D<sub>2</sub>O encounters the UMON surface and adsorbs through H-bonding interactions with iminodiacetate and piperazinium molecules (Fig. S23†). The selectivity of the channels prevents a majority of the surface-bound D<sub>2</sub>O from entering the channel, although it can undergo proton exchange with the  $-\text{NH}_2$  groups associated with the organic molecules on the surface. Ultimately, surface exchange with <sup>1</sup>H atoms results in the formation of light water molecule and this molecule can enter the channel. This hypothesis explains the uptake of H<sub>2</sub>O even in the presence of 99.96% pure D<sub>2</sub>O (Fig. 3A) and the fraction of D<sub>2</sub>O/HDO molecules that thermally desorb from the surface at high temperatures in the TGA experiments (Fig. 3B). In the flow-through experiments, surface adsorption can readily occur, but the exchange reaction requires longer time periods and can only be observed when the experiment is extended to 6 hours. The mass difference between the uptake (6.1%) and release (5.8%) for the flow-through experiments suggests that in this particular experiment, exchangeable <sup>2</sup>D is still present on the surface.

The one caveat of these results is the relatively modest limits of detection for both the FTIR of the evolved gases and condense phased characterization. To enhance the limit of detection and more rigorously assess the isotope selectivity, we expose dehydrated UMON crystallites to HTO vapor in a closed hydration chamber to assess the amount of <sup>3</sup>T adsorption on the hydrophilic surfaces or within the channels of the UMON material. Natural abundance hydrated samples are used as controls because the protons on the surface in the hydrated samples will exhibit exchange with <sup>3</sup>T but the water in the channels of the fully hydrated samples will not exchange because the confined water does not exchange with the bulk on this time scale. We have previously shown that there is minimal exchange of the confined water inside the channels on the timescale of these experiments, making the hydrated UMON material an ideal control.<sup>16</sup> Liquid scintillation counting quantifies the activities of both the experimental and control samples, and the experimental samples show activities that are, within error, identical to those of the control samples (Fig. S24 and S25†). Using the measured specific activity of the tritiated water and subtracting the background (surface adsorption) allows us to calculate the average amount of HTO present at  $80 \pm 40$  nmol HTO per mg UMON (Fig. S26†). If the tritiated water were to occupy both the nanochannels and the surface for the initially dehydrated samples, then we would expect to observe 3222 nmol HTO per mg UMON. The amount we observe represents  $2.48 \pm 1.24$  wt%, which falls within the range observed for surface bound water (0.6–3%). Due to the radioactivity of the sample, we could not further evaluate the material using TGA analysis, but our results also suggest selectivity between isotopologues.

Although we have considerable experimental evidence to support both the conclusion that the UMON material selectively



prefers light water over heavy forms and a proposed model for surface exchange and uptake of light water even in an overwhelming isotopic abundance of heavy water, the mechanism of the isotopic selectivity of the channel remains unclear. In prior studies on hydration of UMON, we showed that the water molecules in the fully hydrated UMON nanochannel adopt an ice-like structure that is well ordered based on high resolution X-ray crystallographic analysis. There are multiple mechanisms for isotope effects within materials and both kinetic and chemical affinity quantum sieving have been explored within metal organic frameworks.<sup>19–26</sup> Kinetic quantum sieving can occur when the difference between the pore size and the molecular hard core of the guest molecule becomes comparable to the de Broglie wavelength of the guest molecule.<sup>19</sup> Work by Heine and Hirscher and others indicate that materials with pore sizes of  $\sim 2.5\text{--}9.1\text{ \AA}$  can selectively bind heavier  $\text{D}_2$  isotopes over lighter  $\text{H}_2$  molecules at separation factors of 11–12 at temperatures  $<100\text{ K}$ .<sup>20–24</sup> Nevertheless, this level of selectivity would not be enough to explain our observations and is only achievable in the previously studied systems at low temperatures. Chemical affinity quantum sieving relies on a difference in the adsorption enthalpy within the channels as a result of the mass effect on the zero-point energy differences for the bound isotopologues, but is challenging to precisely design the optimal system for this process to be used effectively and also yields modest separation factors and only at relatively low temperatures.<sup>25,26</sup> Thus, it seems unlikely that either of these quantum sieving mechanisms could be responsible for the observed selectivity of the UMON nanopores for light water over all forms of heavy water.

A recent study of hydrophobic nanoconfinement of light and heavy water at interfaces shows significant nuclear quantum effects on the stability and ordering of the hydrogen bond network at hydrophobic interfaces that arise as a purely nanoscale phenomenon that does not significantly affect the bulk properties of the liquid–solid interface.<sup>27</sup> Interestingly, this study shows greater stabilization of  $\text{D}_2\text{O}$  relative to  $\text{H}_2\text{O}$  under nanoconfinement, which is in contrast to the observations reported here. Still, the observation of distinctive nuclear quantum effects as a result of nanoscopic confinement illustrates the potential for these kinds of effects to ultimately lead to the level of selectivity that we report in our experiments. Although the precise molecular mechanisms that lead to the high level of isotope fractionation that we observe in the UMON material remain uncertain, it seems likely that nuclear quantum effects on the structure and stability of hydrogen bond networks under confinement in the UMON pores is central to the reported results. Future studies will focus on untangling the contributions to this effect by probing the structure–function relationships that determine the isotope effect observed within the UMON material.

## Author contributions

T. Z. F., M. K. P., A. L. J. and P. S. designed the experiments; M. K. P., A. L. J., P. S., L. C. A., A. G., G. C. performed the experiments; T. Z. F., M. K. P., P. S., G. C., C. M. C. analysed data; T. Z.

F. and M. K. P. wrote the manuscript. All authors edited and revised the manuscript.

## Conflicts of interest

There are no conflicts to declare.

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