Rising global standards of living coupled with the recent agreement to eliminate hydrofluorocarbon refrigerants are creating intense pressure to develop more sustainable climate control systems. In this vein, the use of water as the refrigerant in adsorption heat pumps is highly attractive, but such adsorption systems are constrained to large size and poor efficiency by the characteristics of currently employed water sorbents. Here we demonstrate control of the relative humidity of water uptake by modulating the pore size in a family of isoreticular triazolate metalorganic frameworks. Using this method, we identify a pair of materials with stepped, nonoverlapping water isotherms that can function in tandem to provide continuous cooling with a record ideal coefficient of performance of 1.63. Additionally, when used in a single-stage heat pump, the microporous Ni2Cl2BBTA has the largest working capacity of any material capable of generating a 25 °C difference between ambient and chiller output.
Tunable Metal-Organic Frameworks Enable High Efficiency Cascaded Adsorption Heat Pumps

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Summary
Rising global standards of living coupled with the recent agreement to eliminate hydrofluorocarbon refrigerants are creating intense pressure to develop more sustainable climate control systems. In this vein, the use of water as the refrigerant in adsorption heat pumps is highly attractive, but such adsorption systems are constrained to large size and poor efficiency by the characteristics of currently employed water sorbents. Here we demonstrate control of the relative humidity of water uptake by modulating the pore size in a family of isoreticular triazolate metal-organic frameworks. Using this method, we identify a pair of materials with stepped, non-overlapping water isotherms that can function in tandem to provide continuous cooling with a record ideal coefficient of performance of 1.63. Additionally, when used in a single-stage heat pump, the microporous Ni₂Cl₂BBTA has the largest working capacity of any material capable of generating a 25 °C difference between ambient and chiller output.

Introduction
More than 44% of all primary energy in the U.S. residential and commercial sectors is consumed by climate control systems.[1] Globally, as standards of living in the developing world rise, the resultant rapid growth in demand for air conditioning is a major driver of increased worldwide electricity consumption.[2] Thermally-driven cooling cycles, such as adsorption heat pumps (AHPs), can utilize previously wasted energy resources such as engine exhaust, waste heat from power generation, or chemical process heat, resulting in dramatic reductions in primary energy consumption for heating and cooling.[3–5] Interest in AHPs using water as the refrigerant has spiked in light of the recent agreement to phase out hydrofluorocarbons, the refrigerants most commonly employed in conventional vapor compression heat pumps, due to their very high global warming potential.[6–8] Widespread deployment of AHPs is limited by shortcomings of the currently employed active sorbents, including low water capacity and shallow water uptake step profiles that are not tunable as a function of relative humidity (RH).[3,9–12] These shortcomings result in bulky devices which suffer from low thermal efficiency. In principle, the efficiency of AHP systems, as measured by the coefficient of performance (COP, defined as the cooling energy per unit energy input), could be improved by employing multiple-effect cascaded cycles wherein two or more adsorbent beds are operated in series.[13] Cascaded
temperature stages allow for the reuse of input thermal energy at progressively lower temperatures in each successive bed, resulting in higher overall efficiency. For instance, a three-stage cascaded AHP using silica gel and activated carbon adsorbents can achieve a superb overall COP of 1, but requires an elevated driving temperature of 220 °C.[14] Further improvements require designing pairs of materials with water uptake steps tailored to fully reuse the excess energy from the high temperature stage with the low temperature stage. Such fine tuning in adsorbent design has not been possible so far.

In view of their recently demonstrated utility as sorbents in AHPs,[4,9,10,15,16] metal-organic frameworks (MOFs) are poised to offer thermodynamically matched material pairs that would enable record performance in cascaded systems. In addition to higher overall capacity, MOFs typically exhibit steeper uptake steps, occurring in a narrow RH range, which directly improve efficiency because cycling between the filled state and the empty state requires a minimal temperature swing.[17] Most importantly for cascaded AHPs, MOFs allow for precise control over the position of water uptake as a function of RH. [11,18–25] This is vital because it determines the maximum temperature difference (lift) an AHP system can produce and defines the required driving temperature to regenerate the sorbent.[9] More hydrophilic sorbents will adsorb water at lower RH, and can produce a
higher lift, but will also require a higher temperature heat source for regeneration because water is adsorbed more strongly. Therefore, the ideal sorbent is one whose hydrophilicity provides enough driving force to achieve the target temperature lift, but is not so high as to decrease thermal efficiency.[9,16]

We recently reported a MOF, Co$_2$Cl$_2$BTDD (BTDD = bis(1H-1,2,3-triazolo[4,5-b],[4′,5′-i])dibenzo[1,4]dioxin) whose pore size allows water sorption at the uptake reversibility limit.[16,26] Co$_2$Cl$_2$BTDD is the best-performing sorbent for single-stage AHPs capable of a 20 °C lift.[9,16] Here we show that a smaller-pore isoreticular MOF made from the shorter bistriazolate linker $1H,5H$-benzo(1,2-d),(4,5-d′)bistriazole (BBTA)[27,28] is significantly more hydrophilic and enables a lift greater than 40 °C, which, in combination with the larger pore system, can enable a record high-efficiency (COP > 1) cascaded double-effect AHP system.

Results

Water vapor adsorption isotherms measured at 298 K for activated samples of M$_2$Cl$_2$BBTA (M = Co, Ni, Cu) are shown in Figure 2. All three materials exhibit type I isotherms with pore filling at low RH, attributed to coordination of water at the metal nodes, followed by further hydrogen bonding-assisted adsorption until pore saturation. The nickel and cobalt materials both adsorb nearly 30 percent water by weight (wt%) below 5% RH. The copper analog exhibits a significantly lower water capacity, as well as an irreversible step at 30% RH. Powder X-ray diffraction (PXRD) analysis and nitrogen adsorption experiments revealed that the copper material loses porosity and undergoes a phase change after water uptake (Figures S1 and S2). Similarly, the cobalt material largely maintains its crystallinity, but loses much of its porosity after reactivation upon water sorption (Figures S3 and S4). By contrast, the nickel MOF retains its crystallinity and porosity (Figures S5 and S6) even after water sorption and desorption. This stability trend is in line with that

![Figure 2: Water vapor isotherms for M$_2$Cl$_2$BBTA materials.](image)

Adsorption (filled symbols) and desorption (open symbols) at 298 K for Co$_2$Cl$_2$BBTA (blue triangles), Ni$_2$Cl$_2$BBTA (green squares, measured at 296 K), and Cu$_2$Cl$_2$BBTA (orange circles).
previously observed for ammonia sorption in this family of materials and is the result of the kinetics of ligand exchange at the metal ion.[29]

Due to its greater stability, Ni$_2$Cl$_2$BBTA would be the more appropriate choice for a cascaded AHP. The enthalpy of water adsorption ($\Delta H_{ads}$) in Ni$_2$Cl$_2$BBTA, determined by variable temperature water vapor sorption isotherms (Figure S7), is approximately 55 kJ mol$^{-1}$ at zero coverage, nearly identical to that found for water binding to the open coordination sites of Co$_2$Cl$_2$BTDD[16] (Figure S8). Importantly, during pore filling of Ni$_2$Cl$_2$BBTA, $\Delta H_{ads}$ remains above 50 kJ mol$^{-1}$, indicating stronger across-pore hydrogen bonding interactions, likely due to confinement effects,[30,31] than those found in the larger pore Co$_2$Cl$_2$BTDD, wherein $\Delta H_{ads}$ during pore-filling is 46 kJ mol$^{-1}$. The variable-temperature water vapor isotherms for Ni$_2$Cl$_2$BBTA were also used to validate a characteristic curve (Figure S9), which converts the two independent variables that govern uptake, vapor pressure and temperature, into a single parameter related to the Gibbs free energy of adsorption (see Supplementary Information)[9,15,16] and allows extrapolation of the water loading at any temperature and pressure (Figure S10).

**Single-Stage Heat Pump Using Ni$_2$Cl$_2$BBTA**

The characteristic curve can be used to calculate water loadings in Ni$_2$Cl$_2$BBTA under conditions relevant for heat pump applications, which are in turn used to calculate the working capacity and COP. Under refrigeration II conditions (i.e. chiller output of 278 K and ambient of 303 K for 25 °C lift),[9] Ni$_2$Cl$_2$BBTA has a maximum COP of 0.70 and a volumetric working capacity of nearly 0.40 mL H$_2$O mL$^{-1}$ MOF using a 400 K desorption temperature (Figure 3A, filled squares). This is the highest reported volumetric working capacity for any material working under Refrigeration II standardized conditions (i.e. capable of a 25 °C lift). Notably, because of the exceptional hydrophilicity of Ni$_2$Cl$_2$BBTA, the temperature lift it could achieve is primarily limited by the regeneration temperature.

**Figure 3: Single-stage heat pump figures of merit.**

* A) Coefficient of Performance (purple symbols, left axis) and volumetric working capacity (aqua symbols, right axis) for Ni$_2$Cl$_2$BBTA with a chiller output temperature of 278 K and an ambient temperature of either 303 K (filled symbols, 25 °C lift) or 313 K (open symbols, 35 °C lift) as a function of desorption temperature. * B) Heat transferred per cycle, determined by varying the evaporator (output) temperature, versus temperature lift for Ni$_2$Cl$_2$BBTA (squares) and Co$_2$Cl$_2$BTDD (blue triangles).
rather than the position of its water uptake step, which is otherwise dominant in other materials. For instance, if the ambient temperature increases to 313 K, the water-Ni$_2$Cl$_2$BBTA working pair can still produce a 278 K chiller output (35 °C lift) with only minimal decreases in the COP and working capacity (Figure 3A, open squares). Further, with an ambient temperature of 30 °C (303 K), Ni$_2$Cl$_2$BBTA can achieve a 30 °C lift (Figure 3B, filled green squares), and with an ambient temperature of 40 °C (313 K) Ni$_2$Cl$_2$BBTA can achieve a 40 °C lift with only a 10 °C increase in desorption temperature (Figure 3B, open pink squares).

In agreement with the characteristic curve calculations, cycling of Ni$_2$Cl$_2$BBTA between 25 °C and 150 °C at a constant vapor pressure of 4.76 mmHg (20% RH at 25 °C, well beyond the saturation RH value at this temperature) revealed an initial slight decline in uptake followed by a plateau at 0.36 g g$^{-1}$ (Figure S11). We note that this is close to the vapor pressure of water ice at 0 °C, 4.585 mmHg (~20% RH at 25 °C). Impressively, nearly all of the capacity of Ni$_2$Cl$_2$BBTA, greater than 0.33 g g$^{-1}$, is reached below this vapor pressure (Figure 2). Although using water as the refrigerant makes it impossible to cool below 0 °C, the unique behaviour of Ni$_2$Cl$_2$BBTA towards water vapour highlights its potential as a desiccant in other applications such as industrial compressed air drying that require sub-zero dewpoints. [32]

**Two-Stage Cascaded Heat Pump Using Ni$_2$Cl$_2$BBTA and Co$_2$Cl$_2$BTDD**

In order to utilize the full potential driving force for heat transfer with Ni$_2$Cl$_2$BBTA, we devised a concept tandem cascaded heat pump cycle (Figure 4A and 4B). Due to the material’s exceptional affinity for water, water vapor can be adsorbed from a low temperature evaporator onto a bed of Ni$_2$Cl$_2$BBTA maintained between 104 °C and 57 °C. When combined in a tandem cycle with a material exhibiting a lower affinity for water.

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**Figure 4:** Two-stage cascaded heat pump cycle diagram and thermodynamics. **A)** Schematic of a cascaded double-effect adsorption heat pump for stationary applications. **B)** Isosteric cycle diagram of a cascaded adsorption heat pump cycle providing a 17 °C lift with evaporator at 283 K, condenser at 300 K, desorption of Co$_2$Cl$_2$BTDD at 330 K and desorption of Ni$_2$Cl$_2$BBTA at 400 K.
vapor, for instance the larger-pore Co$_2$Cl$_2$BTDD, the sensible heat ($Q_1$) as well as the energy from $\Delta H_{ads}(Q_2)$ in the small-pore Ni$_2$Cl$_2$BBTA (Bed 1) at elevated temperature can be used to completely regenerate the larger-pore Co$_2$Cl$_2$BTDD (Bed 2). Both adsorbent beds can be paired with the same evaporator and condenser, at temperatures of 10 °C and 27 °C respectively (17 °C lift). The minimum adsorption temperature of Bed 2 (Co$_2$Cl$_2$BTDD) would be that of the condenser, 27 °C, and it would be regenerated at 57 °C with waste heat from Bed 1 (Ni$_2$Cl$_2$BBTA). In turn, Bed 1 would operate at a minimum adsorption temperature of 57 °C and be regenerated by an external heat source at 127 °C. This configuration is ideal for the application of district air conditioning for buildings (Figure 4A).

Altogether, Ni$_2$Cl$_2$BBTA and Co$_2$Cl$_2$BTDD combine for a cascaded double-effect AHP cycle in which the input thermal energy for regeneration is effectively used twice, once at 127 °C to regenerate the small pore MOF, and again at 57 °C to regenerate the large-pore MOF. In this tandem configuration, Co$_2$Cl$_2$BTDD has a working capacity of 0.52 mL mL$^{-1}$ and a COP of 0.91 when regenerated at 57 °C, whereas Ni$_2$Cl$_2$BBTA has a working capacity of 0.31 mL mL$^{-1}$ and a COP of 0.72 when regenerated at 127 °C (Figure 5B). Because the COP of the cascading pump is the sum of the COPs for the constituent cycles, the ideal system COP for the combination presented here is 1.63.

The thermodynamic cycle is diagrammed in process flow in Figure 4A and in a tandem isosteric cycle in Figure 4B. Beginning with a saturated bed of Ni$_2$Cl$_2$BBTA (Bed 1, point A1) and an empty bed of Co$_2$Cl$_2$BTDD (Bed 2, point C2), Bed 1 undergoes isosteric heating to point B1 and its vapor pressure increases from 9.5 mmHg (the pressure of the evaporator) to 26.6 mmHg (the pressure of the condenser). At the same time, Bed 2 undergoes isosteric cooling from point C2 to point D2. Subsequently, Bed 1 is further heated while connected to the condenser for isobaric desorption to point C1, while Bed 2 is connected to the evaporator and undergoes isobaric adsorption to point A2. At this point the roles of the beds
reverse in the second half-cycle (Figure 4A). Bed 1 is cooled isostERICally to point DI, connected to the evaporator, and undergoes isobaric adsorption back to A1 while the sensible heat $Q_1$ and the heat of adsorption $Q_2$ from this process are used to regenerate Bed 2, moving from A2 to B2 and finally C2 (Figure 4B).

Discussion

The foregoing results are the first example of materials with thermodynamically-tuned, steep, and non-overlapping water isotherms (Figure 5A) employed in a cascaded adsorption cycle, which enables significantly greater thermal efficiency by reducing the required temperature swings and minimizing the dead thermal mass. For example, using conventional sorbents, which, due to overlapping isotherms necessitate employing different refrigerants for each stage, an AHP with a cooling COP of 1 requires a three-stage cascade with a source temperature of 220 °C.[14] Additionally, previous screening of sorbent-sorbate pairs for tandem heat pumps found theoretical maxima for COP of 1.15 for a tandem activated carbon-methanol and zeolite-water cycle.[13] By contrast, using MOFs with stepped water isotherms tuned by pore size, we demonstrate a record ideal COP of 1.63, over 150% higher than the previous best, notably decreasing complexity by using water as the sole refrigerant, and employing a much lower source temperature of 127 °C.

Triazolate MOFs are uniquely suited to water vapor isotherm engineering by virtue of their exceptional stability as well as their hydrophilic chains of metals exhibiting open coordination sites. The microporous BBTA MOFs adsorb water vapor near 0% RH. This unusually high hydrophilicity engenders a high driving force for heat transfer. Compared to other materials capable of generating a 25 °C lift, Ni$_2$Cl$_2$BBTA has the greatest uptake capacity and positions it as a key material for the hot adsorption bed of a cascaded adsorption heat pump. When used in tandem, Ni$_2$Cl$_2$BBTA and Co$_2$Cl$_2$BTDD combine for an unprecedented COP of 1.63, which can be achieved using an accessible low driving temperature of only 127 °C. The ability to provide continuous cooling with simultaneously high COP and low source temperatures is a significant advance over the state-of-the-art. This work moves sustainable adsorption heat pumps using water as a refrigerant closer to widespread adoption.

Experimental Procedures

Water vapor adsorption isotherms were measured by a volumetric method using a Micromeritics ASAP 2020 gas sorption analyzer with a vapor dose option and a heated manifold. A typical sample of ca. 40 mg of metal-organic framework, pre-activated at 100°C to remove all residual solvent, was transferred in an Ar filled glovebox to a pre-weighed analysis tube. The tube with sample inside was weighed again to determine the mass of the sample. The tube was capped with a Micromeritics TranSeal, brought out of the glovebox, and transferred to the analysis port of the gas sorption analyzer. Free space correction measurements were performed using ultra-high purity He gas (UHP grade 5, 99.999% pure). Water vapor adsorption isotherms were measured using Milli-Q water. The water was degassed on the ASAP 2020 manifold prior to measurement. All analyses were performed using a water bath held at constant temperature with a recirculating chiller. The manifold was held 15 °C above the temperature of the sample water bath, and the vapor dosing tube was held 10 °C above the temperature of the sample water bath. Oil-free vacuum pumps were used to prevent contamination of sample or feed gases.
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Author Contributions
AJR and MD conceived of experiments. AJR and AMW synthesized materials and performed experiments. AJR, SR, HK, and ENW designed and calculated parameters for the cascaded thermal cycle. ADL performed cycling experiments. AJR wrote the initial manuscript. AJR, AMW, SR, HK, ADL, ENW, and MD revised the manuscript.

Declaration of Interests
AJR and MD are inventors on a patent pertaining to the materials discussed herein.

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Supplemental Information

Tunable Metal-Orgnic Frameworks Enable High Efficiency Cascaded Adsorption Heat Pumps

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Materials and Methods

CuCl$_2$•4H$_2$O (98%, Strem Chemicals), CoCl$_2$•6H$_2$O (99.9%, Alfa Aesar), NiCl$_2$•6H$_2$O (Strem Chemicals), HCl (32-35%, BDH – VWR Analytic) methanol (99.9%, VWR), N,N-dimethylformamide (99.8%, Millipore), ethanol (ACS grade, Mallinckrodt) were used as received.

Powder X-ray diffraction (PXRD) patterns were recorded with a Bruker Advance II diffractometer equipped with a $\theta/2\theta$ Bragg-Brentano geometry and Ni-filtered CuK$_\alpha$ radiation ($K\alpha_1=1.5406$ Å, $K\alpha_2=1.5444$ Å, $K\alpha_1/K\alpha_2=0.5$). The tube voltage and current were 40 kV and 40 mA, respectively. Samples for PXRD were prepared by placing a thin layer of the appropriate material on a zero-background silicon crystal plate.

Nitrogen adsorption isotherms were measured by a volumetric method using a Micromeritics ASAP 2020 gas sorption analyzer. A typical sample of ca. 40 mg of metal-organic framework, pre-activated at 100°C to remove all residual solvent, was transferred in an Ar filled glovebox to a pre-weighed analysis tube. The tube with sample inside was weighed again to determine the mass of the sample. The tube was capped with a Micromeritics TranSeal, brought out of the glovebox, and transferred to the analysis port of the gas sorption analyzer. Free space correction measurements were performed using ultra-high purity He gas (UHP grade 5, 99.999% pure). Nitrogen isotherms were measured using UHP grade nitrogen. All nitrogen analyses were performed using a liquid nitrogen bath at 77 K. Oil-free vacuum pumps were used to prevent contamination of sample or feed gases.

Water vapor adsorption cycling experiments were performed using a Surface Measurement Systems DVS Adventure dynamic gravimetric water sorption analyzer. A typical sample of ca. 5 mg of MOF, pre-activated at 150°C, but later exposed to air, was loaded into the microbalance. The instrument was set to deliver a constant vapor pressure of 4.76 mmHg. The baseline mass of the sample was measured over 2 hours at the desorption temperature. The sample was then cycled between the adsorption temperature, and the desorption temperature 10 times. The sample mass was measured once per minute. Heating and cooling ramp rates were 1°C per minute. The instrument switched to the next cycle when the change in mass per minute was less than 0.02%.

M$_2$Cl$_2$BBTA MOFs were synthesized and activated according to previously published procedures.[1–3]
Calculation of the Characteristic Curve for Ni$_2$Cl$_2$BBTA.

In order to make the loading dependent on only one free variable (A, energy) rather than two (P,T), we constructed a characteristic curve.[4–6] The energy parameter used is the adsorption potential (A), which is the Gibbs free energy of adsorption with inverse sign:

$$ A = RT \ln \left( \frac{p_o(T)}{p} \right) $$

$p_o$ is the saturation vapor pressure of the working fluid at the temperature of analysis, $p$ is the pressure at each loading value, R is the ideal gas constant, and T is the analysis temperature.

Isotherms measured at different temperatures should collapse onto a single characteristic curve. Water isotherms for Ni$_2$Cl$_2$BBTA were measured at 283 K, 288 K, and 296 K (Figure S7). After calculating the characteristic curve using each of these isotherms, it is evident that they all collapse into a single characteristic curve (Figure S9) confirming the validity of the characteristic curve model.

The characteristic curve for the highest temperature 296 K isotherm, that closest to the temperatures typically found in AHP systems, was then fitted with a simple cubic equation in order to easily extrapolate the water loading for any A value (Figure S10).

Calculation of the Heat Transferred per cycle as a Function of Temperature Lift for the Water-Ni$_2$Cl$_2$BBTA working pair (Figure 3B).

To determine temperature lift, the evaporator temperature, which determines the vapor pressure of the water being adsorbed by the MOF, was varied while the condenser at ambient temperature (set equal to the minimum temperature of the MOF bed) was held constant at 303 K or 313 K and the desorption temperature was held constant at 405 K or 415 K. The adsorption potential was then calculated for the empty state, using the vapor pressure and temperature at desorption conditions, that is, the vapor pressure of the condenser, and the desorption temperature. The adsorption potential was also calculated for the full state, using a bed temperature of 303 K or 313 K and with the water vapor pressure allowed to vary with the condenser temperature from 273 K to 303 K or 313 K (30° or 40° lift). The adsorption potential values were then converted to loadings of water in Ni$_2$Cl$_2$BBTA by using the previously determined cubic equation fitted to the characteristic curve. The desorption loading was then subtracted from the adsorption loading to determine the net working capacity. Loading values of water per gram MOF were then converted into energy values using the enthalpy of evaporation of water (44 J g$^{-1}$). Gravimetric values were converted into volumetric values using the crystal density of the MOF (1.1 g cm$^{-3}$).

Calculation of the Coefficient of Performance for the Water- Ni$_2$Cl$_2$BBTA working pair.

The COP for cooling with a 25 °C temperature lift (Refrigeration II conditions as defined by De Lange, et. al)[6] was calculated as described in the literature.[4,5] Briefly, the COP for cooling is defined as:

$$ \frac{Q_{ev}}{Q_{rejen}} $$
The COP for single-stage cooling may vary between 0 and 1. \( Q_{ev} \) is the heat energy removed from the evaporator through evaporation of the working fluid, and \( Q_{regen} \) is the heat energy required to regenerate the adsorbent, including isosteric heating, of both the adsorbent and the working fluid, and isobaric desorption steps. The enthalpy of adsorption of water in Ni\(_2\)Cl\(_2\)BBTA was assumed to be 53 \( \text{kJ mol}^{-1} \) throughout the working range. This is the enthalpy of adsorption at the pore filling step (Figure S8). The heat capacity of the sorbent was assumed constant at 1 \( \text{J g}^{-1} \text{K}^{-1} \), in accordance with the literature.[6] The relevant temperatures for the calculation for refrigeration II[6] conditions are \( T_{\text{cond}} = 303 \text{K} \), \( T_{ev} = 278 \text{K} \). The desorption temperature was varied. The COP for cooling applications with a 25°C or 35°C temperature lift for water in Ni\(_2\)Cl\(_2\)BBTA at varied desorption temperature is shown in Figure 3A.

**Calculation of the Coefficient of Performance for the Tandem Cycle working pair.**

For the tandem cycle, similar calculations as above were employed for each bed separately. The total COP for the combined tandem cycle is then the sum of the COP for each component bed.

In order to match the thermal energy available from bed 1 of Ni\(_2\)Cl\(_2\)BBTA with the required regeneration energy for bed 2, the bed sizes can be adjusted based on the relative thermal energy required for each material. For instance, moving from \( C1 \) to \( A1 \) (Figure 4B), Ni\(_2\)Cl\(_2\)BBTA dissipates 958 J g\(^{-1}\) of thermal energy and for regeneration from \( A2 \) to \( C2 \), Co\(_2\)Cl\(_2\)BTDD requires 1997 J g\(^{-1}\), meaning that 2.08 times as much Ni\(_2\)Cl\(_2\)BBTA than Co\(_2\)Cl\(_2\)BTDD by mass should be employed. Thus, for a 10 kg (14.4 L) bed of Co\(_2\)Cl\(_2\)BTDD, a 20.8 kg (18.9 L) bed of Ni\(_2\)Cl\(_2\)BBTA should be used, which will provide continuous cooling with a 17°C lift and an ideal COP of 1.63. Per cycle, 18300 kJ of thermal energy will be transferred for Co\(_2\)Cl\(_2\)BTDD and 14400 kJ for Ni\(_2\)Cl\(_2\)BBTA.
**Figure S1.** Powder X-ray diffraction patterns of activated Cu$_2$Cl$_2$BBTA (orange), Cu$_2$Cl$_2$BBTA after a single water isotherm at 298 K (purple), and a simulated pattern for the structure (black).
Figure S2. N\textsubscript{2} adsorption isotherms of activated Cu\textsubscript{2}Cl\textsubscript{2}BBTA (orange), and Cu\textsubscript{2}Cl\textsubscript{2}BBTA reactivated at 150°C after a single water isotherm at 298 K (purple).
Figure S3. Powder X-ray diffraction patterns of activated Co$_2$Cl$_2$BBTA (blue), Co$_2$Cl$_2$BBTA after a single water isotherm at 298 K (light blue), and a simulated pattern for the structure (black).
Figure S4. \( \text{N}_2 \) adsorption isotherms of activated \( \text{Co}_2\text{Cl}_2\text{BBTA} \) (blue), and \( \text{Co}_2\text{Cl}_2\text{BBTA} \) reactivated at 150°C after a single water isotherm at 298 K (light blue).
Figure S5. Powder X-ray diffraction patterns of activated Ni$_2$Cl$_2$BBTA (dark green), Ni$_2$Cl$_2$BBTA after a single water isotherm at 296 K (light green), and a simulated pattern for the structure (black).
**Figure S6.** \( \text{N}_2 \) adsorption isotherms of activated Ni\(_2\)Cl\(_2\)BBTA (dark green), and Ni\(_2\)Cl\(_2\)BBTA reactivated at 150°C after a single water isotherm at 296 K (light green).
Figure S7. Variable temperature water adsorption isotherms for Ni$_2$Cl$_2$BBTA at 296 K (purple squares), 288 K (orange circles), and 283 K (blue triangles).
Figure S8. The enthalpy of adsorption for water in Ni$_2$Cl$_2$BBTA versus loading.
Figure S9. Characteristic curves for water vapor adsorption in Ni$_2$Cl$_2$BBTA calculated using the isotherms at 296 K (purple squares), 288 K (orange circles), and 283 K (blue triangles). The characteristic curve is temperature invariant, justifying its usage to calculate isotherms at other temperatures relevant for AHPs.
Figure S10. Cubic least-squares fit to the characteristic curve measured at 296 K. The black line is according to the equation $W = -0.1869A^3 + 1.332A^2 - 18.23A + 481.2$. The $R^2$ value for the fit was 0.988. The fitted equation was used to calculate loading at temperatures and pressures relevant for AHPs.
Figure S11. Temperature-swing water cycling of Ni$_2$Cl$_2$BBTA at a constant water vapor pressure of 4.76 mmHg.
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