Controlled Gas Uptake in Metal–Organic Frameworks with Record Ammonia Sorption

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Supporting Information

ABSTRACT: Ammonia is a vital commodity in our food supply chain, but its toxicity and corrosiveness require advanced protection and mitigation. These needs are not met efficiently by current materials, which suffer from either low capacity or low affinity for NH3. Here, we report that a series of microporous triazolate metal–organic frameworks containing open metal sites exhibit record static and dynamic ammonia capacities. Under equilibrium conditions at 1 bar, the materials adsorb up to 19.79 mmol NH3 g−1, more than twice the capacity of activated carbon, the industry standard. Under conditions relevant to personal protection equipment, capacities reach 8.56 mmol g−1, 27% greater than the previous best material. Structure–function relationships and kinetic analyses of NH3 uptake in isostuctural micro- and mesoporous materials made from Co, Ni, and Cu reveal stability trends that are in line with the water substitution rates in simple metal–aquo complexes. Altogether, these results provide clear, intuitive descriptors that govern the static and dynamic uptake, kinetics, and stability of MOF sorbents for strongly interacting gases.

INTRODUCTION

As an irreplaceable feedstock for global agriculture and industry,1 ammonia is produced at a rate of 140 million metric tons per year, making it among the largest volume chemicals on the planet.2 Although critical to our food supply and economy, NH3 is a highly toxic gas even in small concentrations. Therefore, sorbents capable of removing trace NH3 are of interest for industrial air remediation, personal protective equipment, capacities reach 8.56 mmol g−1, 27% greater than the previous best material. Further, the interpenetrated nature of the porous polymers leads to a highly tortuous pore network with small windows, which substantially retards gas uptake.9 Indeed, although a high maximum NH3 capacity is desirable, increasing the uptake often comes at the expense of slower sorption kinetics. For instance, nonporous MgCl2 has the greatest capacity for ammonia of any known material, but slow kinetics as well as a large volume expansion upon NH3 uptake limit its utility in many applications.14 Despite its obvious importance, the kinetics of ammonia sorption in porous materials is seldom investigated. Reports of sorbent performance overwhelmingly focus on uptake capacity with no mention of adsorption kinetics or, at best, only indirect investigations through the analysis of gas breakthrough experiments.13,15

Recently, we reported a series of particularly robust MOFs, M2Cl2(BTDD) (M = Mn, Co, Ni, Cu; BTDD = bis(1H-1,2,3-triazolo[4,5-b][4′,5′-i]dibenzo[1,4]dioxin),16,17 which feature large hexagonal mesoporous channels with a diameter of 2.3 nm lined with chains of Lewis acidic metals that exhibit open coordination sites (Figure 1). These materials were the first examples of MOFs that were stable to repeated sorption and their stable all-carbon backbone offering distinct advantages for performance under humid conditions. However, the heterogeneous pore size distribution in these porous organic polymers impedes studies of pore size effects on sorption kinetics.13 Further, the interpenetrated nature of the porous polymers leads to a highly tortuous pore network with small windows, which substantially retards gas uptake.9 Indeed, although a high maximum NH3 capacity is desirable, increasing the uptake often comes at the expense of slower sorption kinetics. For instance, nonporous MgCl2 has the greatest capacity for ammonia of any known material, but slow kinetics as well as a large volume expansion upon NH3 uptake limit its utility in many applications.14 Despite its obvious importance, the kinetics of ammonia sorption in porous materials is seldom investigated. Reports of sorbent performance overwhelmingly focus on uptake capacity with no mention of adsorption kinetics or, at best, only indirect investigations through the analysis of gas breakthrough experiments.13,15

To address this need, recent efforts have focused on developing sorbents with Lewis or Bronsted acidic active sites that impart higher affinity for the basic NH3 molecule.8,9 Metal–organic frameworks (MOFs) with coordinatively unsaturated, Lewis acidic open metal sites such as Cu2(trimesate)2− (HKUST-1), Mg2(2,5-dihydroxybenzenedicarboxylate) (Mg-MOF-74), and Co(isonicotinic acid)2 have emerged as superlative dry NH3 adsorbents. However, their performance almost invariably suffers upon repeated exposure to NH3 and, more critically, they often degrade upon exposure to water vapor.10–12 Porous polymers with a high density of Bronsted acid sites also demonstrate superior NH3 sorption,5,13

Received: January 9, 2018
Published: February 9, 2018
desorption of ammonia. Their extreme stability to corrosive
gases was further highlighted by repeated exposure to chlorine
and bromine gas, which can also be stored reversibly in these
materials over multiple cycles. Here, we utilize these
materials and their smaller-pore isoreticular analogues,
M2Cl2BBTA (M = Co, Ni, Cu; BBTA = 1H,5H-benzo(1,2-
d),(4,5-d′)bistriazole), as a ready-made platform for
interrogating the effects of modifying the pore size and active
site density on the total uptake and the kinetics of ammonia
sorption. We show that the higher capacity for ammonia in
the smaller-pore materials cannot be accounted for simply by
the increase in the density of open metal sites. Instead, cooperative
proximity effects likely lead to the anomalous observed increase
in ammonia saturation uptake for M2Cl2BBTA. Using Infra-
SORP, a recently developed calorimetric technique, we
further draw correlations between pore size and sorption kinetics. Finally, we show that MOF stability to ammonia or
water, an oft-empirical parameter of fundamental and practical
importance, correlates directly with the water exchange rates in
homoleptic metal−aquo complexes, a fundamental property
established decades ago that will serve as an easy guide for
new MOF design in the future.

■ RESULTS AND DISCUSSION
Ammonia sorption experiments for activated samples of
M2Cl2BBTA (M = Co, Ni, Cu) at 298 K revealed type I
isotherms, with steep uptakes at low absolute pressure (Figure
2). The total NH3 uptakes at 1 bar were 17.95, 14.68, and 19.79
mmol NH3 g−1 for the Co, Ni, and Cu analogues, respectively.
As compared to the larger pore BTDD materials, the uptakes
observed for BBTA materials are considerably higher, positioning Cu2Cl2BBTA as the material with the highest
NH3 uptake among all MOFs. Intriguingly, the increase in
gravimetric capacity in going from M2Cl2BTDD to the smaller
pore M2Cl2BBTA is greater than what would be expected from
simply increasing the density of active open metal sites. For
instance, the equilibrium uptake of 2.72 molecules of NH3 per
Co atom at 1 bar and 298 K in mesoporous Co2Cl2BTDD increases to 3.20 molecules of NH3 per Co atom in
microporous Co2Cl2BBTA. We attribute this anomalous increase in ammonia density to the shorter distance between
neighboring chains of metal-bound NH3 molecules in
Co2Cl2BBTA. This may allow for increased hydrogen-bonding
interactions between the first, metal-bound NH3 molecules and
additional NH3 adsorbed in the pore to an extent that is not
available in the larger-pore M2Cl2BTDD series. In support of
this hypothesis, we note that a similar confinement effect is
operative for NH3 uptake in microporous zeolites.

After exposure to NH3 at 1 bar and 298 K, Ni2Cl2BBTA(retains its crystallinity and porosity (Figures S4.1 and S5.2), whereas Cu2Cl2BBTA becomes amorphous and nonporous
(Figures S4.2 and S5.3), both in line with the behavior observed for the respective larger pore analogues (for
Cu2Cl2BTDD, Figures S3.2, S4.3, and S5.4). By contrast, powder X-ray diffraction (PXRD) measurements indicate that
Co2Cl2BBTA, unlike its larger pore analogue, loses much of its
crystallinity upon exposure to 1 bar ammonia (Figure S4.4). This also leads to increased hysteresis during desorption
(Figure 2) and substantially reduced Brunauer−Emmett−
Teller (BET) surface area (Figure S5.5 and Table S5.1).
Although Co$_2$Cl$_2$BBTA collapses upon exposure to 1 bar NH$_3$, it remains stable and retains its porosity at 1 mbar NH$_3$, conditions that are more relevant for personal protection equipment (Figures S4.4 and S5.5). Furthermore, NH$_3$ cycling experiments revealed that, despite its collapse at higher ammonia pressure, Co$_2$Cl$_2$BBTA retains a substantial uptake upon cycling with reactivation at 200 °C. These results mirror those observed for mesoporous Mn$_2$Cl$_2$BTDD (Figure S3.1) and are in line with other studies that have shown that a decrease in surface area does not predict a decline in the ammonia uptake; rather, the interaction strength between binding sites and ammonia is a more important predictor of ammonia uptake. Although the total surface area decreases, the open metal sites that strongly bind ammonia evidently remain accessible.

The different stability observed for Co$_2$Cl$_2$BBTA and Co$_2$Cl$_2$BTDD under identical conditions was intriguing because in the vast majority of MOFs where the primary coordination sphere of the metal is conserved, as here, it is the larger pore material that undergoes pore collapse more readily. To investigate this anomalous behavior, we measured a series of variable temperature ammonia isotherms for the larger pore Co$_2$Cl$_2$BTDD. More pronounced hystereses became evident as the isotherm temperature dropped below 298 K, and at 263 K we observed a hysteresis in Co$_2$Cl$_2$BTDD similar to that in Co$_2$Cl$_2$BBTA (Figure 3). As with the small pore analogue, this lower temperature NH$_3$ isotherm also led to partial structural collapse and decreased crystallinity (Figure S4.5). These experiments showed that pore collapse and hysteresis are closely correlated with the local concentration of ammonia in the pore. The cobalt frameworks are stable when ammonia uptake is limited at and around the open metal sites in a layer/cluster mechanism, but if the pores of these materials start filling with ammonia, which begins to occur at room temperature for the small-pore analogue and at lower temperature for the large-pore material, the crystallinity suffers. With Ni$_2$Cl$_2$BTDD, ammonia sorption remains quasi-reversible even at 263 K, although the isotherm profile changes relative to the 298 K isotherm, with the 298 K capacity reached at only 0.4 bar, followed by a pronounced step increase and saturation at 35 mmol g$^{-1}$ at 1 bar (Figure 3). The increase in uptake followed by saturation is characteristic of pore-filling behavior, observed previously in these systems for water adsorption. Importantly, Ni$_2$Cl$_2$BTDD retains its crystallinity and high surface area even after pore filling (Figures S4.6 and S5.6) and represents, to our knowledge, the first example of reversible ammonia pore-filling behavior in a MOF, of potential interest for adsorption heat pumps.

**Simple Kinetic Descriptors Define MOF Stability toward Polar Coordinating Molecules.** Calorimetric measurements and theoretical calculations have shown that porous phases of MOFs are almost universally metastable, that is, uphill in energy with respect to their dense phases. Finding the more stable structures in this class has largely been empirical, but, as with any material, stabilization of a metastable porous MOF phase relative to the dense phase can only be kinetic. It must involve raising the reaction coordinate energy barrier between the two phases, either by decreasing the energy of the porous phase or by increasing the transition state energy. The former is more intuitive: extensive previous work has shown that increasing the metal–ligand bond strength, generally the weakest link in MOF structures, results in enhanced stability toward polar analytes, for example, water and ammonia, a result of lowering the energy of the porous frameworks. Similarly, the triazolate frameworks investigated here already exhibit enhanced stability over their carboxylate containing counterparts such as MOF-74. Here, we further posit that the stability trend within this family of triazolate MOFs follows the kinetic metal–aquo substitution rate, an effect of the transition state energy. The Ni$^{2+}$ materials are most stable toward ammonia: the small-pore Ni$_2$Cl$_2$BBTA framework is the only one to retain its surface area after exposure to 1 bar NH$_3$, and the large-pore Ni$_2$Cl$_2$BTDD withstands pore filling with NH$_3$. The Co$^{2+}$ materials both withstand layer-cluster ammonia binding, but not pore filling, whereas the Cu$^{2+}$ and Mn$^{2+}$ materials degrade even at low concentrations of ammonia. This stability trend is in line with the experimentally observed kinetic metal–aquo substitution rate. Ni$^{2+}$ is 2 orders of magnitude more kinetically inert than Co$^{2+}$, which, in turn, is approximately 2 orders of magnitude more kinetically inert than either Mn$^{2+}$ or Cu$^{2+}$. We note that kinetic stability has also been invoked to explain the exceptional robustness of the carboxylate-based Cr$^{3+}$-containing MIL-101. Notably, in the MIL-53 and 47 family of isostructural frameworks, cation inertness is a more relevant predictor of stability: chemical robustness decreases in the order Cr$^{3+}$ > Al$^{3+}$ > V$^{3+}$, in line with the water substitution rates in the respective aquo complexes, but diverging from the thermodynamic trend in metal–oxygen bond strength. Further, even partial replacement of Mg$^{2+}$ in Mg-MOF-74 with more inert Ni$^{2+}$ ions significantly enhances the stability of the Mg material. Although incorporation of kinetically inert metals into MOFs can be challenging synthetically, future efforts in this direction can provide a route toward enhancing chemical robustness.

**Pore Size Effects on Breakthrough Performance and Kinetics.** The small- and large-pore cobalt frameworks contain identical, stable, strongly adsorptive active sites, which are regularly spaced through one-dimensional pores of varying diameters, providing an ideal platform to investigate the effects of pore size on NH$_3$ breakthrough performance as well as kinetics. Dynamic breakthrough measurements at 1000 ppm of NH$_3$ revealed that Co$_2$Cl$_2$BBTA has a record dry capacity (uptake until 1000 ppm is reached) of 8.56 mmol g$^{-1}$, 27% higher than the state-of-the-art HKUST-1, while the larger-pore Co$_2$Cl$_2$BTDD has a capacity of 4.78 mmol g$^{-1}$ (Figure 4A, Table 1). These breakthrough capacities are equal to 1.48
molecules of NH$_3$ per Co atom for the small-pore material and 1.08 molecules per open Co site for the large-pore material. Interestingly, the breakthrough criterion for ammonia (5 ppm) is reached after the small-pore material has adsorbed 6.24 mmol g$^{-1}$ NH$_3$ and the large-pore MOF has adsorbed 4.42 mmol g$^{-1}$ NH$_3$, corresponding to 1.08 and 1.00 molecules of ammonia per open metal site, respectively. These results demonstrate that increasing the density of open metal sites results in a linear increase in the breakthrough performance, while the saturation capacity is further enhanced in the small pore material, in line with the proximity effects observed in the static isotherms discussed above. In the dry breakthrough curves, the NH$_3$ concentration remains at the baseline until all metal sites have been filled, and then the downstream concentration begins to rise (Figure 4A). The steep increase in downstream NH$_3$ concentration for the large-pore material further evidences weaker secondary interactions as compared to the shallow sigmoidal rise in concentration for the small-pore material. Expectedly, NH$_3$ breakthrough measurements performed at 80% relative humidity (RH) revealed that both small-pore Co$_2$Cl$_2$BBTA and large-pore Co$_2$Cl$_2$BTDD are affected by moisture (Figure 4B, Table 1). Co$_2$Cl$_2$BBTA retains a saturation capacity of 4.36 mol kg$^{-1}$, whereas Co$_2$Cl$_2$BTDD has a humid saturation capacity of 3.38 mol kg$^{-1}$. These values correspond to 0.76 and 0.77 NH$_3$ molecules per open metal site, respectively. These humid breakthrough capacities remain superior to conventional porous materials such as zeolites and activated carbons, and show promise for air purification applications.

The breakthrough experiments demonstrate that the kinetics of NH$_3$ uptake, obviously vital for both personal protection and gas separations applications, are not easily predicted by static isotherm measurements, although many studies of NH$_3$ adsorbents focus solely on capacity. We interrogated the kinetics of NH$_3$ uptake in Co$_2$Cl$_2$BBTA and Co$_2$Cl$_2$BTDD using InfraSORP, an optical calorimetric method. Measurements revealed that the large-pore material has significantly more rapid ammonia uptake (Figure 5). Fitting of the initial 500 s of calorimetric data to two exponentials according to the equation:

$$T = T_0 - k_1(1 - e^{-kt}) - k_2(1 - e^{-kt})$$

showed first-order rate constants, $k_1$, for NH$_3$ uptake of 0.42 and 0.28 s$^{-1}$ for the large-pore and the small-pore Co frameworks, respectively (Figures S7.1 and S7.2). This is not surprising: NH$_3$ molecules are able to more swiftly reach all open Co sites in the larger pores of Co$_2$Cl$_2$BTDD.

The optical calorimetric method assumes that the two materials have equivalent heats of adsorption for ammonia and similar heat capacities. To test these assumptions, the areas under the kinetics curves were integrated and multiplied by the sample masses, providing an estimate for the energy absorbed into the materials (Figures S7.3–S7.6), which should correlate with the amount of ammonia adsorbed, as confirmed previously in other systems. The ratio of heat transferred to the small pore versus the large pore as determined by InfraSORP was 1.49, in excellent agreement with the ratio of 1.48 for equilibrium NH$_3$ capacities at 1 mbar determined from low pressure NH$_3$ isotherms (Figure S3.4), validating the applicability of the InfraSORP method for ammonia.

The kinetic data provide predictive power when combined with relevant models for gas diffusion, such as the Knudsen

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**Table 1. Saturation NH$_3$ Breakthrough Capacities at 1000 ppm in mmol g$^{-1}$ for Co$_2$Cl$_2$BTDD, Co$_2$Cl$_2$BBTA, and Cu$_2$Cl$_2$BBTA (Figure S3.3)**

<table>
<thead>
<tr>
<th>Material</th>
<th>dry (0% RH)</th>
<th>wet (80% RH)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Co$_2$Cl$_2$BTDD</td>
<td>4.78</td>
<td>3.38</td>
</tr>
<tr>
<td>Co$_2$Cl$_2$BBTA</td>
<td>8.56</td>
<td>4.36</td>
</tr>
<tr>
<td>Cu$_2$Cl$_2$BBTA</td>
<td>7.52</td>
<td>5.73</td>
</tr>
</tbody>
</table>

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**Figure 4.** Ammonia breakthrough curves for large-pore Co$_2$Cl$_2$BTDD (green) and small-pore Co$_2$Cl$_2$BBTA (blue) under (A) dry (0% RH) and (B) wet (80% RH) conditions. Dashed lines in (A) indicate 1 molecule of NH$_3$ adsorbed per open coordination site. The challenge concentration $C_o$ was 1000 ppm with a flow rate of 140 mL min$^{-1}$.

**Figure 5.** "InfraSORP" heat flux upon introduction of 1000 ppm ammonia in nitrogen at a flow rate of 140 mL min$^{-1}$ to samples of large-pore Co$_2$Cl$_2$BTDD (green) and small-pore Co$_2$Cl$_2$BBTA (blue).
model. The latter applies when the mean free path of the analyte molecule is larger than the pore diameter of the sorbent. Here, the mean free path for NH₃ with a kinetic diameter of 0.326 nm at 25 °C and 1 bar is 871 Å, indeed much greater than either of the pore diameters of our materials, 13 and 23 Å. In these scenarios, the flux from gas-phase ammonia to an intrapore adsorption site can be approximated as that for a circular tube with rigid, nonadsorbing walls, and is given by

\[ F = \frac{1}{3} \sqrt{\frac{8RT}{\pi M L}} \frac{d \varepsilon}{L_{\tau}} \]

where \( M \) is the molecular weight of NH₃, \( R \) is the gas constant, \( T \) is the temperature, \( d \) is the pore diameter, \( \varepsilon \) is the void fraction, \( L \) is the length of the path from the gas phase to the inner adsorption site, and \( \tau \) is the tortuosity. Because \( M, R, T, \) and \( \tau \) are identical in both MOFs, the ratio \( F_L/F_S \) is thus expected to be \( L_S/L_L \). The ratio \( L_S/L_L \) is difficult to quantify because the average diffusion distance to each active site depends on particle length and aspect ratio, as well as differences in active site density. As such, \( L_S/L_L \) was estimated by scanning electron microscopy (Figures S6.1 and S6.2), which showed that crystallites of the small pore material were on average one-half the size of those observed for the large pore MOF, allowing us to approximate \( L_S/L_L \) as \( \sim 0.5 \). Overall, the expected flux ratio \( F_L/F_S \) was thus expected to be approximately 1.25, in good agreement with the experimentally determined ratio of rate constants of 1.5.

Altogether, the kinetic data show that the Knudsen model serves as a reasonable baseline for understanding and predicting the effects of modifying the pore size and the crystallite size on the speed of gas uptake in MOFs, even when the analytes are strongly interacting gases, and that decreasing particle size can mitigate the slower uptake kinetics resulting from reducing the pore size. Future research aimed at uncovering the kinetics of gas sorption in MOFs in detail will further aid prediction of material suitability for varied applications including personal protective equipment, separations, and heat pumps.

### CONCLUSIONS

Stable triazolate frameworks containing open metal sites provide a unique platform for interrogating coordinating and corrosive gas sorption. Because of their higher density of open metal sites, the smaller-pore BBTA materials have significantly greater capacities than their larger-pore counterparts, with Cu₂Cl₂BBTA exhibiting the greatest static uptake at 1 bar and 298 K of any MOF or porous organic polymer. Additionally, increasing the density of open metal sites results in a linear increase in breakthrough performance, as the breakthrough criterion for NH₃, 5 ppm, is reached after one molecule of ammonia is adsorbed per open metal site in both small-pore Cu₂Cl₂BBTA and large-pore Cu₂Cl₂BTDD. Increasing the density of open metal sites results in a record dry dynamic ammonia capacity for Cu₂Cl₂BBTA at 1000 ppm of 8.56 mmol g⁻¹, exceeding that of the state-of-the-art HKUST-1 by 27%. Decreasing the pore size from 23 to 13 Å results in slower sorption kinetics, although this effect can be mitigated by synthesizing smaller MOF particles. Kinetic analyses correlate with the Knudsen flux model, demonstrating that this theoretical framework serves as a good baseline to predict the effects of modifying the pore size as well as the particle size on sorbent kinetic performance. Further, the stability trend of the isoreticular frameworks substituted with various metal ions follows the metal–aquo substitution rate, with the large-pore framework formed with the most inert metal, Ni₂Cl₂BTDD, able to withstand pore filling with NH₃ at low temperature. This work provides a clear path toward designer sorbents by controlling the uptake, kinetics, and stability of the active materials.

### ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/jacs.8b00313.

Synthetic details, physical methods, InfraSORP curve fitting methods, and Figures S3.1–S7.6 (PDF)

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Notes

The authors declare the following competing financial interest(s): AJR and MD are listed as inventors on a patent pertaining to the materials used in this work.

### ACKNOWLEDGMENTS

This work was supported by the Tata Center for Technology and Design. Fundamental studies of metal-small molecule interactions in MOFs were supported by a National Science Foundation CAREER Award to M.D. (DMR-1452612). We thank Dr. R. Day for assistance with acquiring the SEM images.

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