Selective Decontamination of the Reactive Air Pollutant Nitrous Acid via Node-Linker Cooperativity in a Metal-Organic Framework

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S1
S1 Synthesis

Unless otherwise noted, all reagents were purchased from commercial sources and used without further purification.

Table S1: List of reagents, and their suppliers, used in this work.

| Reagent                                      | Supplier                  | Reagent                           | Supplier                      |
|----------------------------------------------|---------------------------|-----------------------------------|-------------------------------|
| 1,4-diazabicyclo[2.2.2]octane (DABCO)       | Sigma-Aldrich             | 2-aminoterephthalic acid          | Sigma-Aldrich                 |
|                                              |                           | (H2BDC-NH2)                      |                               |
| 2-bromoterephthalic acid (H2BDC-Br)         | Sigma-Aldrich             | Copper (Cu) powder               | Fisher Scientific Company     |
| Dimethylsulfoxide-\(d_6\) (DMSO- \(d_6\))  | Cambridge Isotope Laboratoires, Inc. | Hydrochloric acid (HCl) | Caledon Laboratory Chemicals |
| Methanol (MeOH)                              | ACP Chemicals             | \(N,N\)-dimethylformamide (DMF)  | Fisher Chemical               |
| Sodium acetate (NaOAC)                       | Sigma-Aldrich             | Sodium hydroxide (NaOH)          | ACP Chemicals                 |
| Sodium nitrite (NaNO₂)                       | Sigma-Aldrich             | Sulfuric acid-\(d_2\) (D₂SO₄)    | Sigma-Aldrich                 |
| Terephthalic acid (H₂BDC)                    | Sigma-Aldrich             | Zirconium tetrachloride (ZrCl₄)¹ | Sigma-Aldrich                 |
| Zinc nitrate hexahydrate (Zn(NO₃)₂·6H₂O)    | Sigma-Aldrich             |                                   |                               |

¹ was stored under nitrogen gas prior to use.

S1.1 Synthesis of 2-hydroxyterephthalic acid

The ligand 2-hydroxyterephthalic acid (H₂BDC-OH) was synthesized according to the literature preparation of Zhao et al.¹ A scaled down synthesis was carried out. 1.01 g (5.0 mmol) of H₂BDC-Br and 0.59 g (15.0 mmol) of NaOH were mixed in 20 mL of water.
1.02 g (12.5 mmol) of NaOAc and 0.01 g (0.32 mmol) of Cu powder were added while stirring. The solution was refluxed for 30 hours before a few drops of 1 M NaOH were added to ensure Litmus paper remained blue. The solution was further heated for 6 hours. Subsequently, the terephthalic acid-based compounds were precipitated by addition of concentrated HCl; a white precipitate was filtered and dried. $^1$H NMR (300 MHz, DMSO-$d_6$) $\delta$ 7.90 (dd, $J = 8.1$, 0.6 Hz, 1H), 7.47 (dd, $J = 7.9$, 1.6 Hz, 1H), 7.45 (dd, $J = 1.6$, 0.6 Hz, 1H).

**Fig. S1.** NMR spectrum of synthesized (a) H$_2$BDC-OH and (b) the starting materials, H$_2$BDC-Br.

**S1.2 Synthesis of UiO-66 and UiO-66-NH$_2$:**

UiO-66 and UiO-66-NH$_2$ were synthesized using the same procedure by Katz et al.$^2$ Each MOF was synthesized in 25 mL Duran® glass bottles. Using the preparation of Zwoliński et al.$^3$ the isolated UiO-66-NH$_2$ product was refluxed in 25 mL of MeOH for 24 h to remove any formylated UiO-66-NH$_2$. The purified UiO-66-NH$_2$ was filtered, dried, and stored wrapped in tin-foil in the dark.
S1.3 Synthesis of Zn$_2$(BDC-NH$_2$)$_2$(DABCO)

Synthesis of Zn$_2$(BDC-NH$_2$)$_2$(DABCO) was carried out using the literature preparation from McGrath et al.$^4$ In a 25 mL Duran® glass bottle 0.0947 g (0.844 mmol) DABCO was sonicated to dissolution in 7 mL DMF. To this, 0.5010 g (1.684 mmol) of Zn(NO$_3$)$_2$·6H$_2$O was added along with another 7 mL aliquot of DMF and the resultant solution was sonicated to dissolution. Finally, 0.2675 g (1.526 mmol) of H$_2$BDC-NH$_2$ was added with an additional 7 mL aliquot of DMF and the final solution was sonicated to dissolution. Subsequently, the sample was placed into a 120 °C oven for 48 h. After 48 h, the sample was removed from the oven and the cooled crystals were transferred to a 50 mL centrifuge tube and centrifuged for 5 min at 5000 rpm. The mother liquor was then decanted off and the sample was washed with three 10 mL aliquots of DMF followed by three 10 mL aliquots of MeOH under the same centrifugation conditions mentioned above. The crystals were characterized by NMR.

S2 Characterization

S2.1 Nuclear Magnetic Resonance (NMR)

$^1$H-NMR analyses were measured using a Bruker AVANCE III 300 MHz NMR. The solution phase NMR of all MOF samples were prepared using ca. 5 mg of MOF first dissolved in 2-3 drops of D$_2$SO$_4$. Once dissolved, approximately 1 mL of DMSO-$d_6$ was added as the NMR lock solvent. It should be noted that small chemical shift differences can occur between samples due to the change in dielectric constant caused by the slight variance in pH between samples.

S2.2 Infrared (IR) Spectroscopy

A Bruker Alpha FTIR spectrometer equipped with an ATR crystal was used for IR characterizations. The scanning range was 500 cm$^{-1}$ – 4000 cm$^{-1}$. 
S2.3 Surface Area (SA) Analysis

All samples were thermally activated using a Micromeritics Smart VacPrep gas adsorption sample preparation instrument. The samples were initially heated to 90 °C while a vacuum level below 1.00 mmHg was reached at 5.00 mmHg s\(^{-1}\). Subsequently, the samples were heated to 150 °C for 500 min at a ramp rate of 5 °C min\(^{-1}\). The nitrogen-accessible gas adsorption isotherms (77 K) were measured using a 3Flex Surface Characterization instrument, with the accompanying MicroActive software suites. The accessible surface area was calculated based on Brunauer–Emmett–Teller (BET) theory using the 4-point criterion.\(^5\)

S2.4 Powder X-Ray Diffraction (PXRD)

Powder X-ray diffractograms were measured on a Rigaku Ultima IV X-ray diffractometer with a Cu X-ray source operating at 40 kV x 44 mA (1.76 kW) and a scintillation counter detector. A continuous scan mode was applied between \(2\theta = 5^\circ - 30^\circ\) with a sampling width of 0.020\(^\circ\) and a scan rate of 1.000 \(^\circ/\)min.
S3 Nitrous Acid Measurements

**Fig. S2.** Schematic of the nitrous acid gas-phase source connected to the MOF sample holder and the downstream air generator and detector.

**S3.1 Custom Instrumentation for Nitrous Acid Generation**

The generation of gas-phase nitrous acid was based on an acid-displacement reaction (Fig. S2). To accomplish this, an HCl permeation tube is necessary. A 5 cm long PFA tube (ID = 3 mm, OD = 5 mm) is capped at one end with a 0.5 cm long 1/8" polytetrafluoroethylene (PTFE) rod. The tube is then filled with 2.5 M HCl and the open end is sealed with an identical PTFE rod. A heat gun is used to soften the PFA tube at both ends for insertion of the PTFE caps in order to seal the permeation tube. The PTFE material is porous to HCl gas.

To make a stable/tunable nitrous acid source, a 120 sccm stream of nitrogen gas is divided. Each 60 sccm stream flows through PFA tubing (ID = 1 mm, OD = 3 mm), with one stream
bubbled through water to produce a 100% relative humidity (RH) nitrogen stream. The other stream flows through a heated channel in an aluminum block containing the porous HCl permeation tube. The two streams are recombined to yield 50% RH gaseous HCl. The humid HCl flows over solid NaNO₂ residing in the second channel of the heating block to produce gas-phase nitrous acid. This nitrous acid gas is subsequently used for MOF breakthrough experiments. Higher Al block temperatures result in higher mass-emission rates of HCl and increased nitrous acid production. The output concentration of the nitrous acid source is measured using an Ecotech Serinus 40 Oxides of Nitrogen Analyzer (American Ecotech, Warren, RI).

**S3.2 Quantification of Nitrous Acid in MOF Breakthrough Experiments**

To measure the time-resolved downstream nitrous acid concentration through various MOFs, a thermally activated MOF (activated under vacuum at 100 °C) sample (ca. 10 mg) was packed into a 4 cm PFA tube between two plugs of glass wool (ca. 13 mg each). Downstream nitrous acid measurements made through only glass wool (i.e., in the absence of MOF) resulted in negligible nitrous acid uptake compared to that seen in the presence of MOF. The tubing containing the MOF was connected to the nitrous acid source via push-to-connect fittings. The downstream concentration of nitrous acid through the MOF was measured on the Serinus 40. Pre- and post-MOF gas flow was measured consistently during breakthrough experiments with no observed flow/pressure reduction.

Nitrous acid desorption experiments were done by replacing the nitrous acid source with a 50% RH nitrogen gas stream at 120 sccm. The downstream concentration was monitored with the Serinus 40.
S3.3 Oxides of Nitrogen Analyzer

To measure nitrous acid concentrations, the Serinus 40 was used. The instrument is designed to measure NOx (NO₂ + NO) in the range of 0-20 ppm with a sensitivity of 0.4 ppb. The instrument operates at 0.1 Hz and averages to 1 min measurements. A minimum flow rate of 1500 sccm is needed for the instrument to operate correctly. To that end, a 1500 sccm flow of ultrapure air (GasCal 1100 + 8301 Compressed Air source from American Ecotech, Warren, RI) is combined with the experimental flow at the instrument inlet (downstream of the MOF) through a backpressure exhaust. This ensures nitrous acid concentrations are within the calibrated and linear response range of the instrument and in excess of its intake flow rate; the data accounts for the dilution factor from this.

The general detection principle of the Serinus 40 involves reacting nitric oxide with ozone (O₃) to generate excited state NO₂*. During luminescent relaxation, a photon is released and detected by a photomultiplier. Using this technique, nitric oxide can be measured easily in the O₃-reaction channel. Other nitrogen oxides must first be converted to nitric oxide before they can be measured. The mechanisms for nitrogen dioxide and nitrous acid analysis are synonymous and involve converting either species into nitric oxide using the Mo catalyst heated to 325 °C in the NOx channel. Thus, nitrogen dioxide is determined by difference between the NOx and nitric oxide channels. If only nitrous acid is introduced to the NOx channel, then the instrument can also be exploited to measure nitrous acid, which is well documented. To ensure that we are producing nitrous acid and not nitrogen dioxide, a control experiment is run periodically in which a sodium carbonate (Na₂CO₃)-coated annular denuder (URG Corp., Chapel Hill, NC) is used to remove nitrous acid from our gas stream. Under these conditions, the analyzer is only measuring nitrogen
dioxide and nitric oxide. With the denuder in line, no downstream nitrogen dioxide, or nitric oxide were observed. Thus, our nitrous acid source was producing nitrous acid and no NOx.

The mass-emission rate of nitrous acid was determined by bubbling the output flow through 1 mM KOH solution. Under this condition, nitrous acid is converted back to the nitrite anion. The nitrite concentration was subsequently measured by anion ion chromatography with conductivity detection (IC). The separation was performed using a ThermoScientific ICS-2100 fitted with AS11-HC guard and analytical columns, a KOH eluent generator system (EGC-III), running a gradient elution program at 30 °C and a mobile phase flow of 1.5 mL/min, followed by suppressed conductivity detection.13

S4 MOF Characterization

**Table S2:** Breakthrough times for the first sorption cycle featured in the main text represented in different units (column 2, 3, and 4 correspond to the first sorption curve in Fig. 2a, 2b, and 2c respectively, while column 5 corresponds to the first sorption curve in Fig. 4b).

| Exposure ppbv<sup>a</sup> | BDC | BDC-NH<sub>2</sub> | BDC-(NH<sub>2</sub>)<sub>1/6</sub> | Zn<sub>2</sub>-(BDC-NH<sub>2</sub>)<sub>2</sub>(DABCO) |
|--------------------------|-----|----------------|----------------|----------------------------------|
| 75                      | 35  | 70            | 105          |
| 20                      | 1500| 1500          | 100          |
| 35                      | 17  | 17            | 3.5          |
| 0.011                   | 0.37| 0.74          | 0.074        |
| 0.018 (0.016)<sup>b</sup> | 0.65 (0.57)<sup>b</sup> | 1.2 (1.1)<sup>b</sup> | 0.045        |
| 0.0029 (0.0039)<sup>b</sup> | 0.11 (0.14)<sup>b</sup> | 0.21<sup>c</sup> (0.28)<sup>b</sup> | 0.022        |
| 0.0029 (0.0026)<sup>b</sup> | 0.11 (0.09)<sup>b</sup> | 0.21<sup>c</sup> (0.18)<sup>b</sup> | 0.022        |
| 0.0044 (0.0034)<sup>b</sup> | 0.16 (0.12)<sup>b</sup> | 0.31 (0.24)<sup>b</sup> | N/A          |

<sup>a</sup> Average value of nitrous acid produced prior to the start of breakthrough measurements (i.e., C<sub>0</sub>).

<sup>b</sup> Values in brackets calculated based on 2 missing linkers and the addition of two charge compensating hydroxides and two coordination number compensating water units.

<sup>c</sup> Only 1 in 6 linkers in UiO-66-(NH<sub>2</sub>)<sub>1/6</sub> are BDC-NH<sub>2</sub>. The maximum uptake would equate to 0.17 moles HONO/moles linker (6 linkers per node); the measured value is within error of the expected value.

<sup>d</sup> The acidic proton comes from the Zr<sub>6</sub>O<sub>4</sub>(OH)<sub>4</sub><sup>12+</sup> node. The µ<sub>3</sub>-OH units contain acidic protons.
**S4.1 Triplicate Nitrous Acid Breakthroughs of UiO-66**

![Graph showing nitrous acid breakthrough curves for three samples of UiO-66.](image)

**Fig. S3.** Nitrous acid breakthrough curves for three samples of UiO-66. Each sample was run through 3 sorption and desorption cycles under 50% RH. Blue lines separate sorption and desorption curves, while black lines separate different cycles on the same MOF sample.
S4.2 Nitrous Acid Breakthroughs of UiO-66 with 0% RH Desorption

Fig. S4. Nitrous acid breakthrough curves for one sample of UiO-66. The sample was run through three sorption and desorption cycles. Each desorption cycle flowed a dry (0% RH) N₂ gas stream through the sample. Blue lines separate sorption and desorption curves, while black lines separate different cycles on the same MOF sample. Note the minimal desorption using 0% RH, and the spike in concentration when the MOF is exposed in subsequent (cycles 1 and 2) sorption cycles due to the presence of 50% RH in the subsequent HONO gas stream sorption cycle. The present work illustrates that, unlike the 50% RH desorption in Fig. 2 and Fig. S3, no nitrous acid is desorbed under 0% RH desorption gas. This demonstrates that nitrous acid is chemically trapped as the nitrosonium cation within UiO-66.
S4.3 Triplicate Nitrous Acid Breakthroughs of UiO-66-(NH₂)₁/₆

Fig. S5. Nitrous acid breakthrough curves for three samples of UiO-66-(NH₂)₁/₆. Each sample was run through 3 sorption and desorption cycles under 50% RH. Blue lines separate sorption and desorption curves, while black lines separate different cycles on the same MOF sample.
S4.4 Triplicate Nitrous Acid Breakthroughs of Zn$_2$(BDC-NH$_2$)$_2$(DABCO)

Fig. S6. Nitrous acid breakthrough curves for three samples of Zn$_2$(BDC-NH$_2$)$_2$(DABCO). Each sample was run through 3 sorption and desorption cycles under 50% RH. Blue lines separate sorption and desorption curves, while black lines separate different cycles on the same MOF sample.
S4.5 NMR Data for Pre/Post-Nitrous Acid Exposure

Fig. S7: High concentration (> 1 ppmv) nitrous acid experiment analyzed via solution-phase NMR. (a) UiO-66-NH₂ post nitrous acid exposure showing the formation of only UiO-66-(N₂⁺⁻⁻) (b) UiO-66-NH₂ prior to nitrous acid exposure. (c) UiO-66-(N₂⁺⁻⁻) after air exposure for 3 weeks illustrating that the diazonium salt has been converted to new products (likely UiO-66-OH as well as UiO-66-Cl). (d) H₂BDC-OH.
**Fig. S8:** Solution phase NMR measured in DMSO-$d_6$ (lock solvent) with 2 drops of D$_2$SO$_4$ of (a) UiO-66 pre-nitrous acid exposure and (b) UiO-66 post-nitrous acid exposure. Note that there are no additional peaks in the NMR to suggest a chemical transformation of the BDC linker.
Fig. S9: Solution phase NMR measured in DMSO- $d_{6}$ (lock solvent) with 2 drops of D$_2$SO$_4$ of (a) UiO-66-NH$_2$ (same as Fig. 2a in the main text) for comparison. (b)/(c) Zn$_2$(BDC-NH$_2$)$_2$DABCO measured after different concentration of nitrous acid exposure. (d) Zn$_2$(BDC-NH$_2$)$_2$DABCO measured before nitrous acid exposure. The change in chemical shift between Zn$_2$(BDC-NH$_2$)$_2$DABCO samples is due to changes in the solvent due to the effects of D$_2$SO$_4$ concentration on the linker; DABCO and BDC-NH$_2$ are confirmed via the addition of DABCO and H$_2$BDC-NH$_2$. Note that for the post-exposed Zn$_2$(BDC-NH$_2$)$_2$DABCO samples, there is no evidence of BDC-OH suggesting that BDC-NH$_2$ did not react with HONO in Zn$_2$(BDC-NH$_2$)$_2$DABCO. Furthermore, the small amount of BDC-N$_2^+$ that may be present, is likely due to homogeneous reaction between BDC-NH$_2$ and adsorbed nitrous acid during D$_2$SO$_4$ dissolution; this also confirms that the chemistry observed in the main text is heterogeneous in origin.
S4.6 IR Data for Pre/Post-Nitrous Acid Exposure

Fig. S10: (a) IR of UiO-66-NH₂. (b) IR of UiO-66-NH₂ post-moderate concentration (ca. 10x larger than low concentration exposures; ranging between 650 and 850 ppbv) nitrous acid exposures. (c) IR of UiO-66-NH₂ post-high concentration (> 1 ppmv) nitrous acid exposures. (d) IR of UiO-66 post-low (ranging from 35 to 85 ppbv) concentration nitrous acid exposures. (e) IR of UiO-66-OH. Note the presence of new vibration at 2283 cm⁻¹ in both blue and green traces that is consistent with the formation of a diazonium salt.
S4.7 Gas Adsorption Data for Pre/Post-Nitrous Acid Exposure

**Fig. S11**: N$_2$ gas adsorption isotherms measured at 77 K for (a) UiO-66 pre-HONO exposure (red circles), UiO-66 post-moderate (650 and 850 ppbv) HONO exposure (red triangles), (b) UiO-66-($\text{NH}_2$)$_{1/6}$ pre-HONO exposure (green circles), UiO-66-($\text{NH}_2$)$_{1/6}$ post-moderate HONO exposure (green triangles), (c) UiO-66-NH$_2$ pre-HONO exposure (blue circles), and UiO-66-NH$_2$ post-high (> 1 ppmv) HONO exposure (blue triangles).
S4.8 PXRD Data Pre/Post-Nitrous Acid Exposure

Fig. S12: (a) PXRD of UiO-66 post-high concentration (> 1 ppmv) HONO exposure. (b) PXRD of UiO-66-NH₂ post-high concentration HONO exposure. (c) PXRD for UiO-66.

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