





Supplementary Information for

Breaking the Upper Bound of Siloxane Uptake: Metal-Organic Frameworks as an Adsorbent Platform

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1. Computational details

1.1. Force field parameters for D4. D4 was modelled as a semi-flexible molecule with an all-atom atomistic model. All intramolecular bonds, angles, dihedrals, and cross terms parameters for methyl groups were taken from the consistent-valence force field (CVFF) reminded below.^{S1}

Harmonic Bond

$$U = \frac{1}{2}p_0(r - p_1)^2 \quad [S1]$$

where p_0/κ_B in units K/Å², p_1 in Å.

Table S1: D4 bonding potential parameters.

Pseudo atom	Type of bond	p_0/κ_B (K/Å ²)	p_1 (Å)
Si-O	RIGID_BOND	-	-
Si-C	HARMONIC_BOND	286248.126	1.809
C3-H	HARMONIC_BOND	409668.576	1.105

Harmonic Bend

$$U = \frac{1}{2}p_0(\theta_{ijk} - p_1)^2 \quad [S2]$$

where p_0/κ_B in units K/rad², p_1 in degree.

Table S2: D4 bending potential parameters.

Pseudo atom	Type of angle	p_0/κ_B (K/rad ²)	p_1 (°)
Si-C-H	HARMONIC_BEND	41614.223	112.3
C-Si-C	HARMONIC_BEND	53400.911	113.5
C-Si-O	HARMONIC_BEND	53040.094	117.3
H-C-H	HARMONIC_BEND	47507.567	106.4

CVFF Dihedral

$$U = p_0(1 + \cos(p_1\phi_{ijk} - p_2))^2 \quad [S3]$$

where p_0/κ_B in units K, p_2 in degree.

Table S3: D4 dihedral potential parameters.

Pseudo atom	Type of torsion	p_0/κ_B (K)	p_1 (multiplicity)	p_2 (°)
H-C-Si-C	CVFF_DIHEDRAL	240.545	3	0
H-C-Si-O	CVFF_DIHEDRAL	-60.136	3	0
C-Si-O-Si	CVFF_DIHEDRAL	240.545	3	0

CFF Bond Bond Cross

$$U = p_0(r - p_1)(r' - p_2) \quad [S4]$$

where p_0/κ_B in units K/Å², p_1 and p_2 in Å

Table S4: D4 cross-term bonding potential parameters.

Pseudo atom	Type of bond-bond	p_0/κ_B (K/Å ²)	p_1 (Å)	p_2 (Å)
Si-C-H	CFF_BOND_BOND_CROSS	14312.406	0	0
C-Si-C	CFF_BOND_BOND_CROSS	7336.612	0	0
C-Si-O	CFF_BOND_BOND_CROSS	25257.188	0	0

CFF Bond Bend Cross

$$U = (\theta - p_0)[p_1(r - p_2) + p_3(r' - p_4)] \quad [S5]$$

where p_0 in degrees, p_1 and p_3 in units K/Å/rad, p_2 and p_4 in Å.

Table S5: D4 cross-term bonding-bending potential parameters.

Pseudo atom	Type of bond-angle	p_0 (°)	p_1 (K/Å/rad)	p_2 (Å)	p_3 (K/Å/rad)	p_4 (Å)
Si-C-H	CFF_BOND_BEND_CROSS	0	14733.359	0	9742.06	0
C-Si-C	CFF_BOND_BEND_CROSS	0	781.770	0	0	0
C-Si-O	CFF_BOND_BEND_CROSS	0	11425.871	0	27061.27	0

D4 LJ parameters and charges

The electronic potential (ESP) derived partial charges of D4 were computed by density functional theory (DFT) calculations with PBE (Perdew-Burke-Ernzerhof) functional^{S2} and DNP (double numeric plus polarization) basis set^{S3}, using DMol³^{S4} (Table S6).

Table S6: Charges and LJ parameters for all atoms of D4.

Pseudo atom	Charge (e ⁻)	ϵ/κ_B (K)	σ (Å)
Si	1.321	202.429	3.826
O	-0.763	30.213	3.118
C	-0.889	52.873	3.431
H	0.2032	22.156	2.571

1.2. Screening dataset. Material details.

Table S7: Details of the 29 MOFs added to the COREMOF database.

MOFs	Reference	MOFs	Reference
RAVWAO	S5	DUT-5	S6
RAVWES	S5	DUT-51-Zr	S7
RAVWIW	S5	DUT-67-Zr	S8
RAVWOC	S5	MIL-68(Al)	S9
RAVWUI	S5	Cr-soc-MOF-1	S10
RAVXAP	S5	MIP ^[4] -177	S11
RAVXET	S5	MIP-200	S12
RAVXIX	S5	Zr-IPA ^[5]	S13
MIL-125	S14	Ni-BPM ^[6]	S15
MOF-808-acetate	S14	Ni-BPP ^[7]	S15
MOF-808-formate	S14	Ni-TPM ^[8]	S15
NU ^[1] -1000	S14	Ni-TPP ^[9]	S15
UiO ^[2] -68	S14	Ni-MOF-74	S15
Zr6-AzoBDC ^[3]	S14	PCN ^[10] -224(Ni)	S16

[1]NU: Northwestern University; [2]UiO: University of Oslo;

[3]AzoBDC: azobenzenedicarboxylate;

[4]MIP: Material of the Institute of Porous Materials from Paris;

[5]IPA: isophatale; [6]BPM: biphenyl-meta;

[7]BPP: biphenyl-para; [8]TPM: triphenyl-meta;

[9]TPP: triphenyl-para; [10]PCN: Porous coordination network;

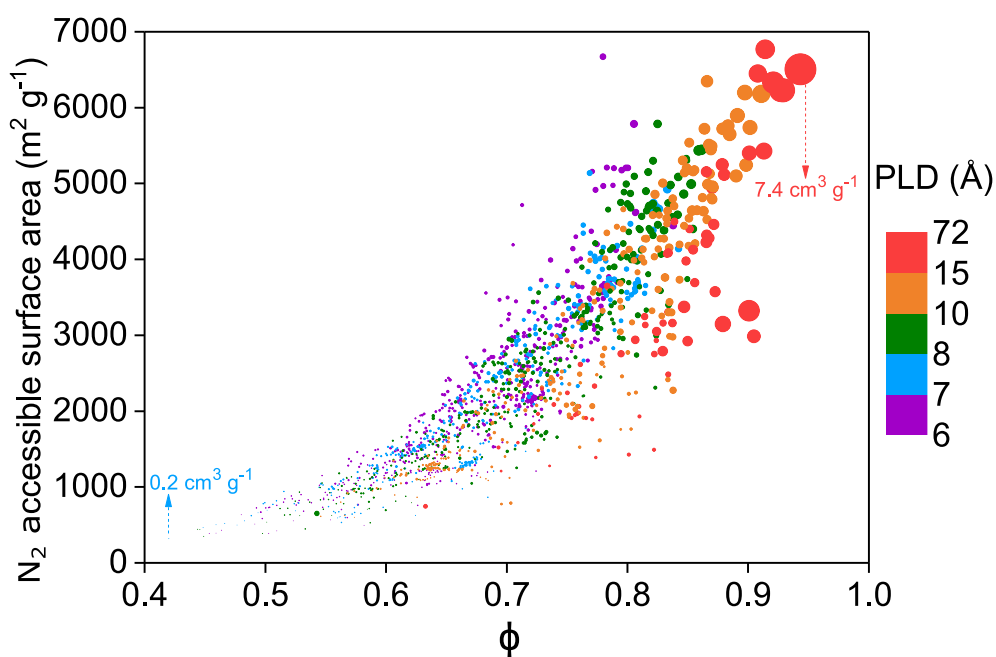


Fig. S1: Overview of the diversity of the MOF database with PLD > 6 Å in terms of void fractions, N_2 accessible surface areas and PLDs. Data points are color coded by PLDs of MOFs. Pore volumes of all structures are represented by size.

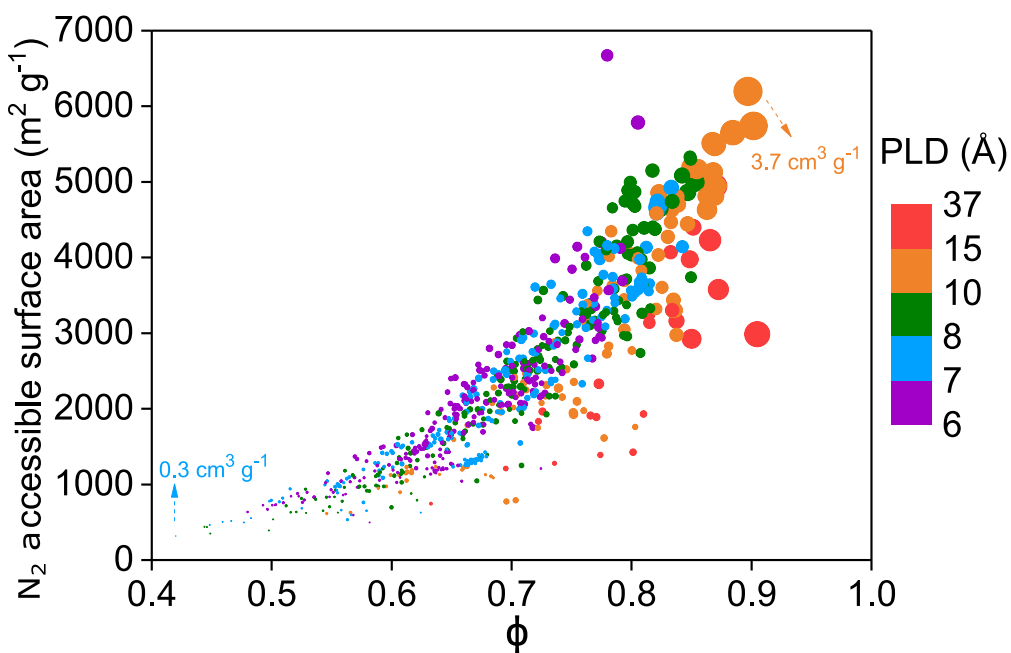


Fig. S2: Overview of the diversity of the hydrophobic MOFs database in terms of void fraction and N_2 accessible surface areas. Data points are color coded by PLDs of MOFs. Pore volumes of all structures are represented by size.

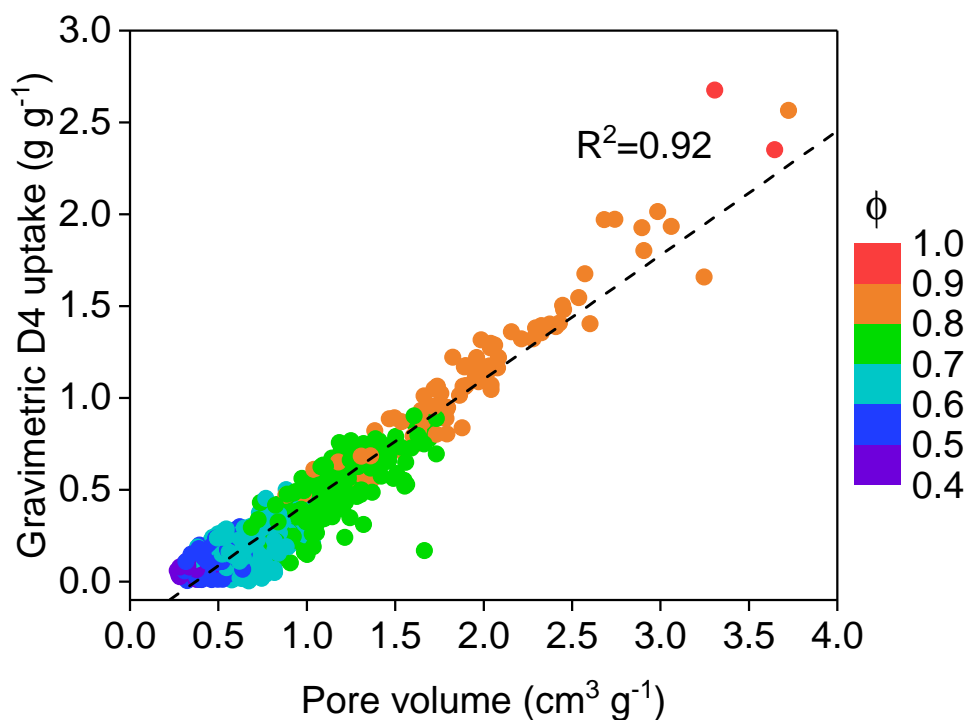


Fig. S3: The relation between the gravimetric D4 uptake of the 811 hydrophobic MOFs (g g^{-1}) and their pore volumes ($\text{cm}^3 \text{g}^{-1}$), color coded by void fraction of the MOFs.

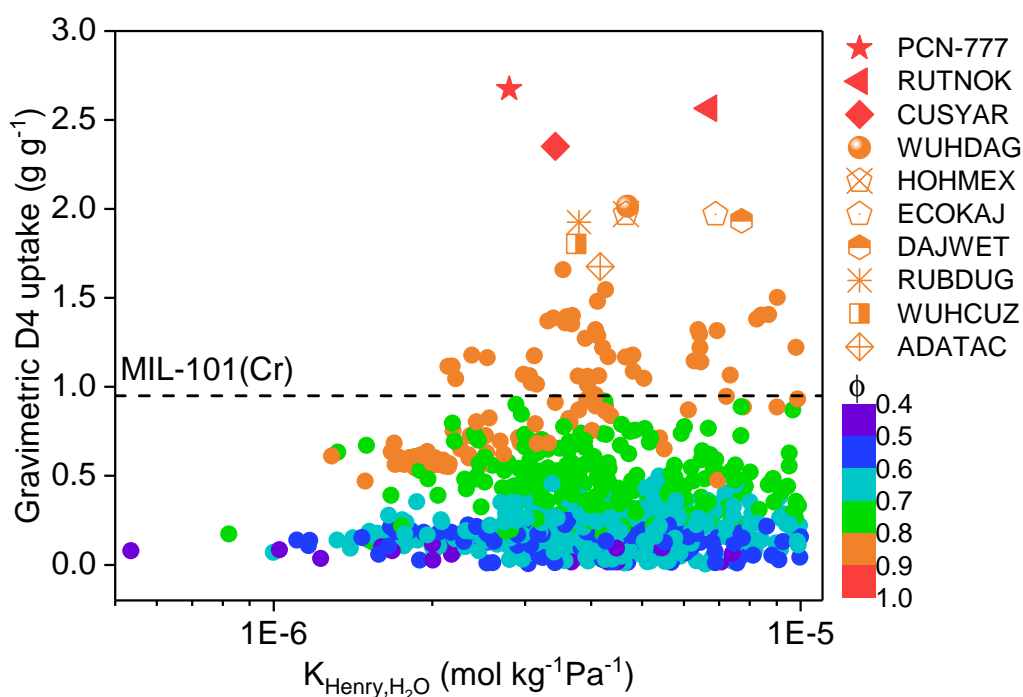
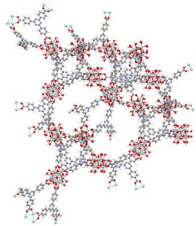
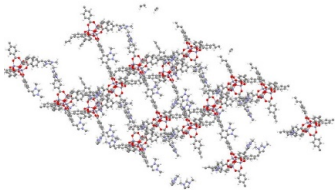
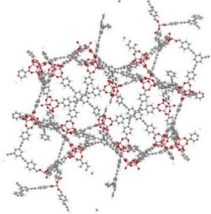
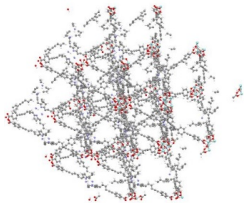
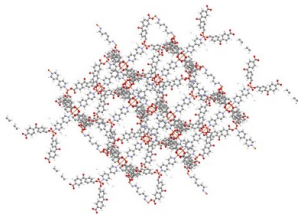
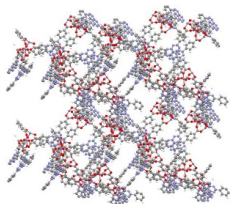
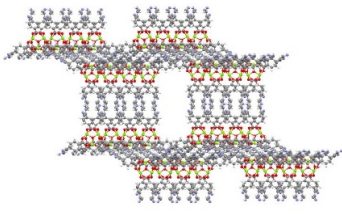
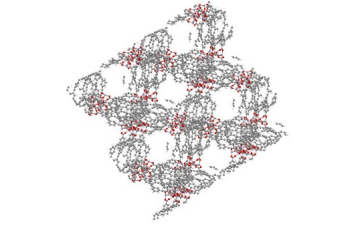
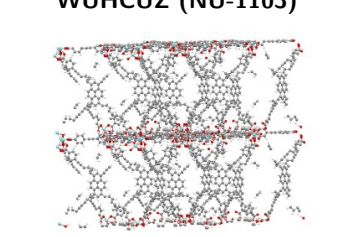
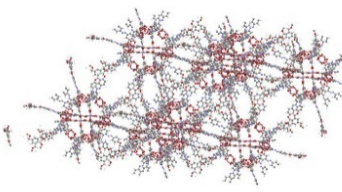


Fig. S4: Predicted D4 uptake performance at 298 K for the hydrophobic MOFs database plotted as a function of their computed Henry constant of water, color coded by void fraction, ϕ . Top performing 10 candidates are represented by different symbols in the legend to the right.

Table S8: Structural details of the top promising 10 hydrophobic materials which exhibit the highest D4 uptake.

MOF	Details
FOTNIN (PCN-777) 	Organic ligand: 4,4',4'-s-triazine-2,4,6-triyl-tribenzoic acid Metal site: Zr PLD: 28.36 Å SA: 2990 m ² g ⁻¹ Density: 0.27 g cm ⁻³ PV: 3.31 cm ³ g ⁻¹ ϕ : 0.90 Gravimetric D4 uptake: 2.68 g g ⁻¹ Volumetric D4 uptake: 0.72 g cm ⁻³
RUTNOK (IRMOF-76) 	Organic ligand: 4,7-bis(4-carboxylphenyl)-1,3-dimethylbenzimidazolium-tetrafluoroborate Metal site: Zn PLD: 14.65 Å SA: 6200 m ² g ⁻¹ Density: 0.24 g cm ⁻³ PV: 3.72 cm ³ g ⁻¹ ϕ : 0.90 Gravimetric D4 uptake: 2.57 g g ⁻¹ Volumetric D4 uptake: 0.62 g cm ⁻³
CUSYAR (MOF-210) 	Organic ligand: biphenyl-4,4'-dicarboxylate Metal site: Zn PLD: 12.18 Å SA: 5700 m ² g ⁻¹ Density: 0.25 g cm ⁻³ PV: 3.65 cm ³ g ⁻¹ ϕ : 0.90 Gravimetric D4 uptake: 2.35 g g ⁻¹ Volumetric D4 uptake: 0.59 g cm ⁻³
WUHDAG (NU-1104) 	Organic ligand: meso-tetrakis-(4-((phenyl)ethynyl)benzoate) porphyrin Metal site: Zr PLD: 10.50 Å SA: 5500 m ² g ⁻¹ Density: 0.29 g cm ⁻³ PV: 2.99 cm ³ g ⁻¹ ϕ : 0.87 Gravimetric D4 uptake: 2.01 g g ⁻¹ Volumetric D4 uptake: 0.58 g cm ⁻³
HOHMEX 	Organic ligand: 4,4'-carbonyldibenzoato - (μ 2-4,4'-bipyridine) Metal site: Cu PLD: 14.89 Å SA: 5000 m ² g ⁻¹ Density: 0.32 g cm ⁻³ PV: 2.74 cm ³ g ⁻¹ ϕ : 0.87 Gravimetric D4 uptake: 1.97 g g ⁻¹ Volumetric D4 uptake: 0.63 g cm ⁻³
ECOKAJ 	Organic ligand: s-heptazine tribenzoate Metal site: Zn PLD: 17.58 Å SA: 3600 m ² g ⁻¹ Density: 0.33 g cm ⁻³ PV: 2.68 cm ³ g ⁻¹ ϕ : 0.87 Gravimetric D4 uptake: 1.97 g g ⁻¹ Volumetric D4 uptake: 0.65 g cm ⁻³

<p>DAJWET</p> 	<p>Organic ligand: tetrakis (4-carboxylatophenyl) porphyrin Metal site: Mg PLD: 26.59 Å SA: 5000 m² g⁻¹ Density: 0.28 g cm⁻³ PV: 3.06 cm³ g⁻¹ ϕ: 0.87 Gravimetric D4 uptake: 1.93 g g⁻¹ Volumetric D4 uptake: 0.54 g cm⁻³</p>
<p>RUBDUP</p> 	<p>Organic ligand: phenylene ethynylene macrocycle Metal site: Zn PLD: 19.25 Å SA: 4200 m² g⁻¹ Density: 0.30 g cm⁻³ PV: 2.90 cm³ g⁻¹ ϕ: 0.87 Gravimetric D4 uptake: 1.93 g g⁻¹ Volumetric D4 uptake: 0.58 g cm⁻³</p>
<p>WUHCUZ (NU-1103)</p> 	<p>Organic ligand: 4,4',4'',4'''-((pyrene-1,3,6,8 tetrayltetrakis(benzene-4,1-diyl)) tetrakis(ethyne-2,1 diyl))tetrabenzoate Metal site: Zr PLD: 12.21 Å SA: 5500 m² g⁻¹ Density: 0.30 g cm⁻³ PV: 2.91 cm³ g⁻¹ ϕ: 0.87 Gravimetric D4 uptake: 1.80 g g⁻¹ Volumetric D4 uptake: 0.54 g cm⁻³</p>
<p>ADATAC</p> 	<p>Organic ligand: 5,5',5''-(4,4',4'''-[1,3,5-phenyltris(methoxy)] tris-phenylazo) tris-isophthalic acid Metal site: Zn PLD: 10.28 Å SA: 5130 m² g⁻¹ Density: 0.34 g cm⁻³ PV: 2.57 cm³ g⁻¹ ϕ: 0.87 Gravimetric D4 uptake: 1.68 g g⁻¹ Volumetric D4 uptake: 0.57 g cm⁻³</p>

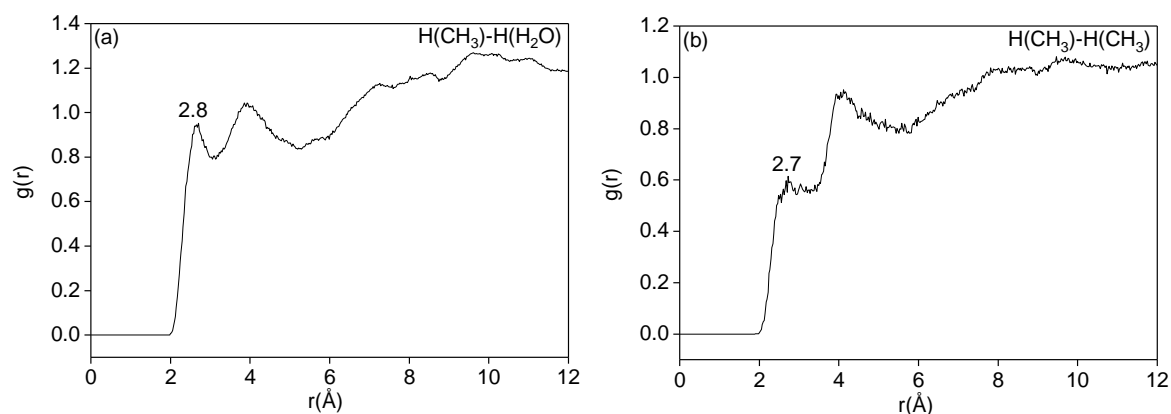


Fig. S5: All-atom averaged radial distribution functions between (a) H atom from CH₃ group of D4 molecules and H atom from coordinated water of the framework at 10% total loading and (b) H atom from CH₃ groups of D4 at 100% loading.

1.3. Radial distribution functions for D4/PCN-777.

2. MOF samples

2.1. MIL-101(Cr). The benchmark MIL-101(Cr) sample was taken from a previous work^{S17}, with all textural characteristics as stated in reference.

2.2. DUT-4. DUT-4 was purchased from Materials Center (TU Dresden, Germany).

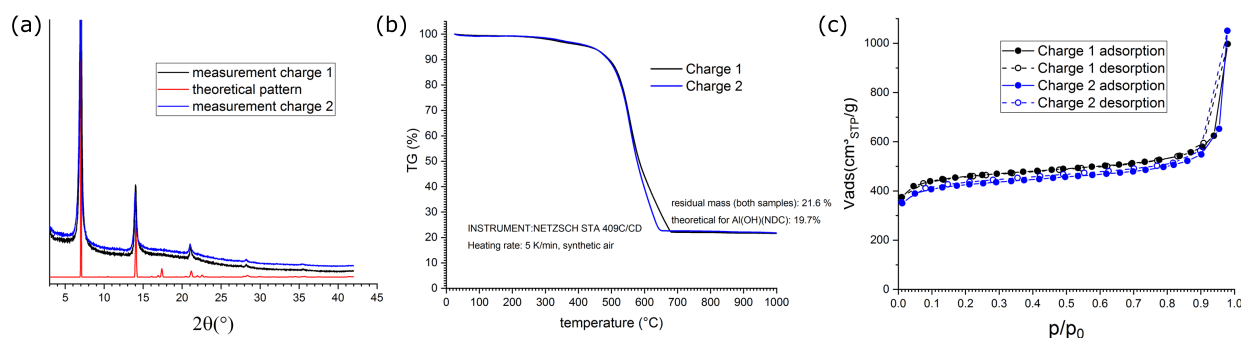


Fig. S6: Characterization of the DUT-4 sample, in duplicates as black and blue: (a) PXRD, alongside simulated pattern in red (b) TGA curves and (c) N_2 physisorption isotherms at 77 K.

2.3. PCN-777. Synthesis

To synthesize the PCN-777, $\text{ZrOCl}_2 \cdot 8\text{H}_2\text{O}$ (1.08 g, 3.351 mmol) and 4,4',4''-s-Triazine-2,4,6-triyl-tribenzoic acid (0.270 g, 0.612 mmol) were put into 36 ml N,N-Diethylformamide (DEF) in a 100 ml Teflon-lined autoclave reactor, alongside an amount of trifluoroacetic acid (1.8 ml) to form a reaction solution. After sonicating the reaction solution at room temperature for 10 min, the reactor was transferred to a convection oven followed by heating at 423 K for 12 h. The PCN-777 crystalline solid was recovered by filtration after purification with 100 ml N,N-Dimethylformamide (DMF) and acetone for 3 h at room temperature. The collected crystalline solid was dried at 393 K for 12 h.

Characterisation

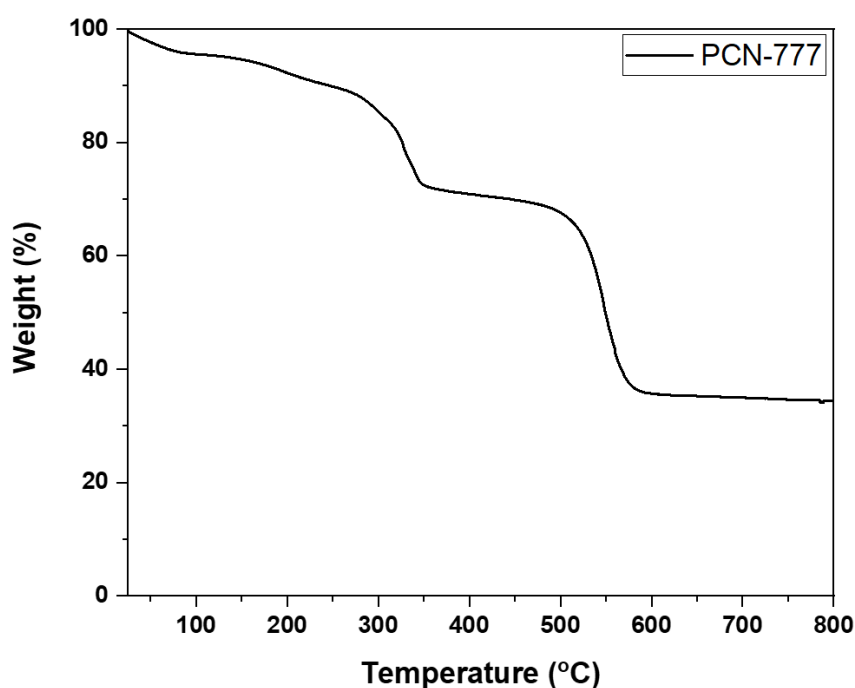


Fig. S7: Thermogravimetric curve recorded on as-synthesised PCN-777.

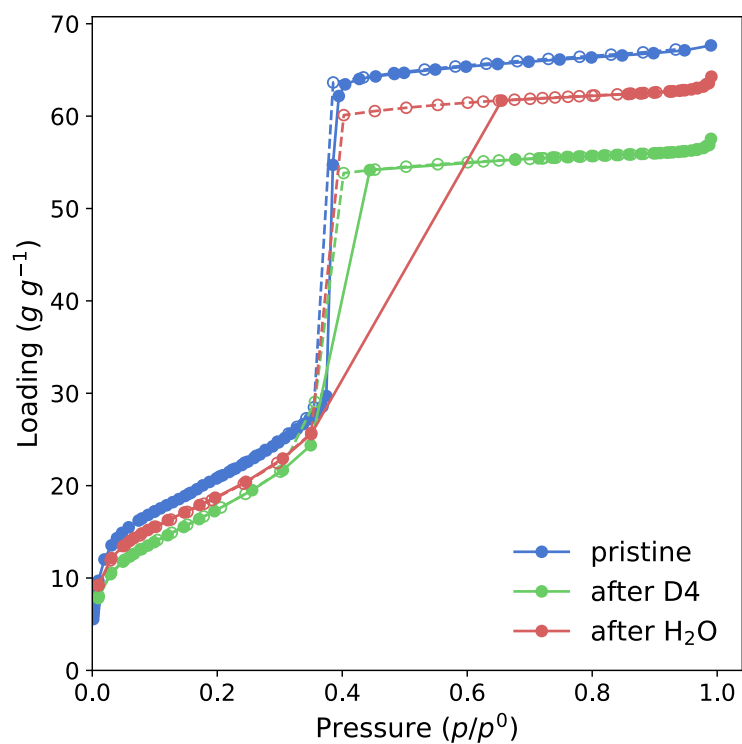


Fig. S8: Nitrogen sorption isotherms at 77 K for the pristine PCN-777, alongside with those measured on samples after D4 and water sorption.

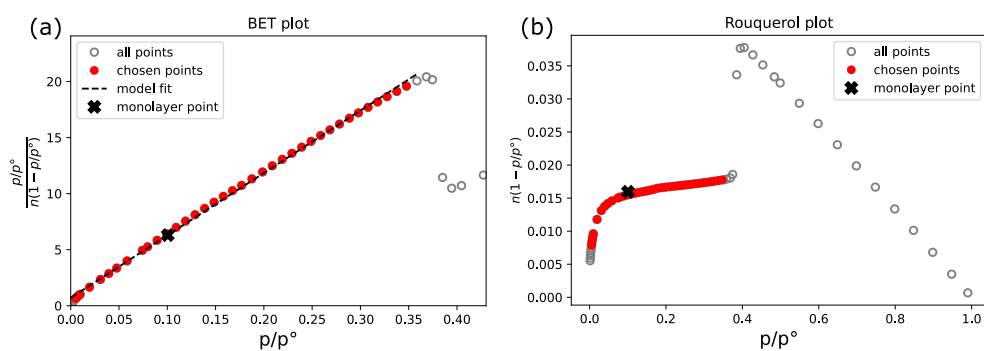


Fig. S9: BET and Rouquerol plots displaying selection of applicable isotherm points for the pristine PCN-777 isotherm.

3. D4 sorption experiments

3.1. D4 benchmarking with known MOFs. Isotherms were recorded on benchmark materials MIL-101(Cr) and DUT-4 using the same methodology detailed in the main manuscript.

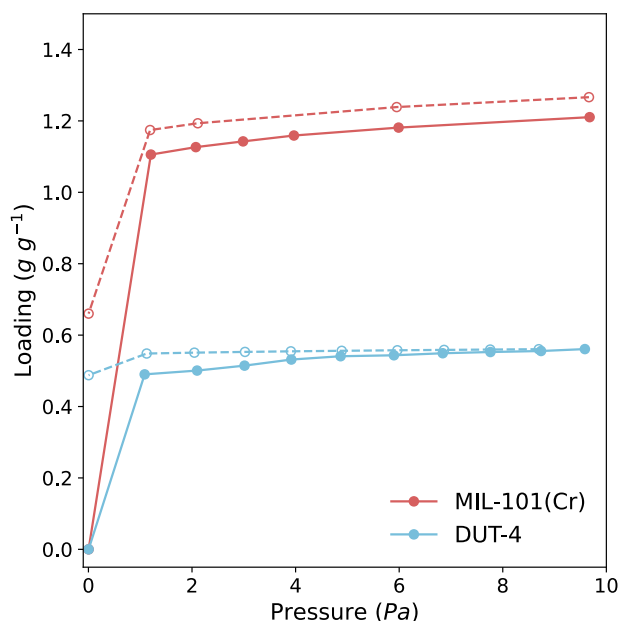


Fig. S10: D4 isotherms recorded on samples of MIL-101(Cr) (red) and DUT-4 (blue), used to validate our computational methodology for predicting total D4 capacity. Note the different desorption behavior (open symbols) of the two materials under secondary vacuum: partial desorption for MIL-101(Cr) and no desorption for DUT-4.

3.2. Isosteric heat of sorption of D4. A further isotherm was recorded at 313 K (40 °C) to allow for the calculation of the isosteric heat of adsorption through the Clausius-Clapeyron equation, as depicted in Fig. S11.

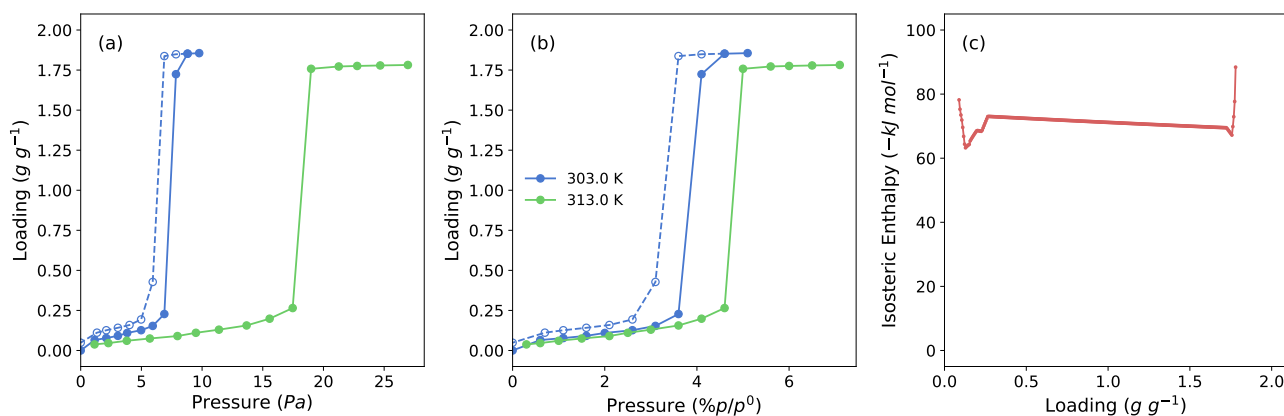


Fig. S11: D4 sorption isotherms on PCN-777 recorded at 303 K (blue) and at 313 K (green) in an absolute (a) and relative (b) pressure scale. (c) The calculated isosteric heat of adsorption as a function of D4 uptake.

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