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). S1

Harmonic Bond

$$U = \frac{1}{2}p_0(r - p_1)^2$$
 [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/Ų) | 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

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

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

$$U = p_0(r - p_1)(r' - p_2)$$
 [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/Ų) | 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

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

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

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

| Pseudo atom | Type of bond-angle | po (°) | 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 module Materials Studio S5 (Table S6).

Table S6: Charges and Lennard-Jones parameters for all atoms of D4.

| Charge (e^{-}) | ϵ/κ_B (K) | σ (Å) |
|------------------|---------------------------|---|
| 1.321 | 202.429 | 3.826 |
| -0.763 | 30.213 | 3.118 |
| -0.889 | 52.873 | 3.431 |
| 0.2032 | 22.156 | 2.571 |
| | 1.321 -0.763 -0.889 | 1.321 202.429 -0.763 30.213 -0.889 52.873 |

1.2. Screening dataset. 1710 non-disordered MOFs with pore limiting diameters > 6 Å were screened from CoRE-MOF database, to which 29 well-known MOFs were added with proven stability and high porosity, including the MOF-74 isoreticular series ^{S6}, Cr-soc-MOF-1 ^{S7}, UiO-68 (UiO for University of Oslo), MOF-808, ^{S8} and a several structures originating from Dresden University of Technology (DUT) and Material of Institute Lavoisier (MIL). This additional list is given in Table S7.

The combined dataset of 1739 MOFs show a broad range of geometric and textural features: PLDs (6.0–71.5 Å), N₂ accessible SA (320–6770 m² g⁻¹), density (0.13–2.22 g cm⁻³), ϕ (0.42–0.94) and PVs (0.23–7.46 cm³ g⁻¹).

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

| MOFs | Reference | MOFs | Reference |
|---------------------------|-----------|------------------------------|-----------|
| RAVWAO | S6 | DUT-5 | S9 |
| RAVWES | S6 | DUT-51-Zr | S10 |
| RAVWIW | S6 | DUT-67-Zr | S11 |
| RAVWOC | S6 | MIL-68(AI) | S12 |
| RAVWUI | S6 | Cr-soc-MOF-1 | S7 |
| RAVXAP | S6 | MIP ^[4] -177 | S13 |
| RAVXET | S6 | MIP-200 | S14 |
| RAVXIX | S6 | Zr-IPA ^[5] | S15 |
| MIL-125 | S8 | Ni-BPM ^[6] | S16 |
| MOF-808-acetate | S8 | Ni-BPP ^[7] | S16 |
| MOF-808-formate | S8 | Ni-TPM ^[8] | S16 |
| NU ^[1] -1000 | S8 | Ni-TPP ^[9] | S16 |
| UiO ^[2] -68 | S8 | Ni-MOF-74 | S16 |
| Zr6-AzoBDC ^[3] | S8 | PCN ^[10] -224(Ni) | S17 |

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

 $[3] Azo BDC:\ azobenzene dicarboxy late;$

[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;

 $[9] \mathsf{TPP} \colon \mathsf{triphenyl\text{-}para}; \ [10] \mathsf{PCN} \colon \mathsf{Porous} \ \mathsf{coordination} \ \mathsf{network};$

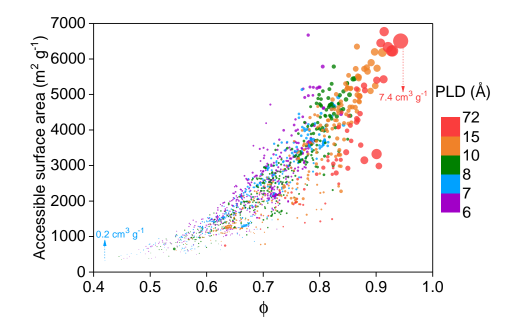


Fig. S1: Overview of the diversity of the MOF database with PLD > 6 Å in terms of void fractions, accessible surface areas and PLDs. Data points are color coded by PLDs of MOFs. Pore volumes of all structures are represented by size.

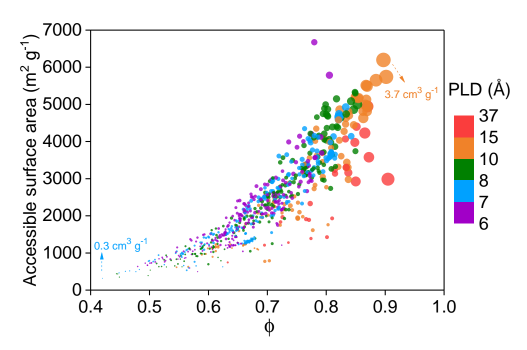


Fig. S2: Overview of the diversity of the hydrophobic MOFs database in terms of void fraction and accessible surface areas. Data points are color coded by PLDs of MOFs. Pore volumes of all structures are represented by size.

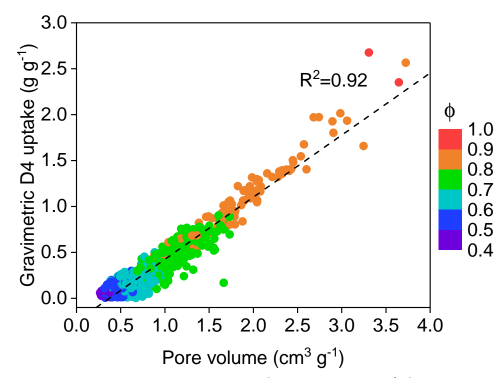


Fig. S3: The relation of gravimetric D4 uptake of the 811 hydrophobic MOFs (g g^{-1}) and their pore volumes (cm³ g^{-1}), color coded by void fraction of the MOFs.

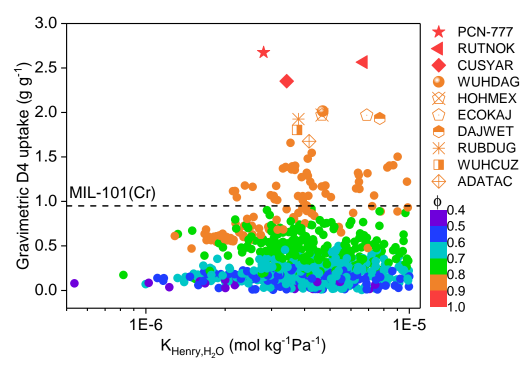


Fig. S4: Predicted D4 uptake performance at 298 K for the hydrophobic MOFs database plotted as a function of their computed Henry constant for water, color coded by void fraction, ϕ . Top performing 10 candidates are represented by different symbols in the legend to the right.

| MOF | Details |
|-------------------|---|
| FOTNIN (PCN-777) | Organic ligand: 4,4',4'-s-triazine-2,4,6-triyl-tribenoic acid Metal site: Zr PLD: 28.36 Å SA: $2990\mathrm{m^2g^{-1}}$ Density: $0.27\mathrm{gcm^{-3}}$ PV: $3.31\mathrm{cm^3g^{-1}}$ ϕ : 0.90 Gravimetric D4 uptake: $2.68\mathrm{gg^{-1}}$ Volumetric D4 uptake: $0.72\mathrm{gcm^{-3}}$ |
| RUTNOK (IRMOF-76) | Organic ligand: 4,7-bis(4-carboxylphenyl)-1,3-dimethylbenzimidazium-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 m² g⁻¹ Density: 0.29 g cm⁻³ PV: 2.99 cm³ 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: $5000\mathrm{m^2g^{-1}}$ Density: $0.32\mathrm{gcm^{-3}}$ PV: $2.74\mathrm{cm^3g^{-1}}$ ϕ : 0.87 Gravimetric D4 uptake: $1.97\mathrm{gg^{-1}}$ Volumetric D4 uptake: $0.63\mathrm{gcm^{-3}}$ |
| 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 |

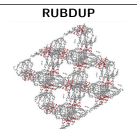
DAJWET

Organic ligand: tetrakis (4-carboxylatophenyl) porphyrin

Metal site: Mg PLD: 26.59 Å SA: $5000 \text{ m}^2 \text{ g}^{-1}$ Density: 0.28 g cm^{-3} PV: $3.06 \text{ cm}^3 \text{ g}^{-1}$

 ϕ : 0.87

Gravimetric D4 uptake: $1.93\,\mathrm{g\,g^{-1}}$ Volumetric D4 uptake: $0.54\,\mathrm{g\,cm^{-3}}$



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

WUHCUZ (NU-1103)

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: $1.80\,\mathrm{g\,g^{-1}}$ Volumetric D4 uptake: $0.54\,\mathrm{g\,cm^{-3}}$

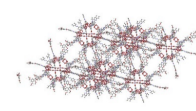
ADATAC

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: $5130 \text{ m}^2 \text{ g}^{-1}$ Density: 0.34 g cm^{-3} PV: $2.57 \text{ cm}^3 \text{ g}^{-1}$ ϕ : 0.87

Gravimetric D4 uptake: $1.68\,\mathrm{g\,g^{-1}}$ Volumetric D4 uptake: $0.57\,\mathrm{g\,cm^{-3}}$



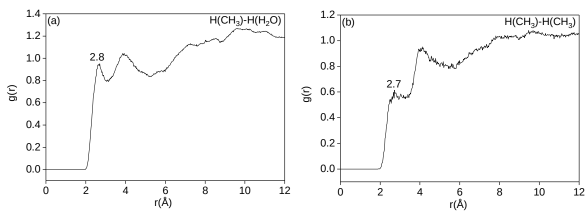


Fig. S5: All-atom averaged radial distribution functions between (a) H atom from CH_3 group of D4 molecules and H atom from coordinated water of the framework at 10% total loading and (b) H atom from CH_3 groups of D4 at 100% loading.

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 ^{S18}, with all textural characteristics as stated in reference.

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

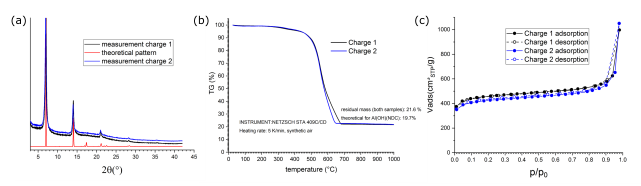


Fig. S6: Characterization of the DUT-4 sample, in duplicates as black and blue: (a) PXRD, alongside simulated pattern in red (b) TGA curves and (c) N_2 physisorption isotherms at 77 K.

2.3. PCN-777. Synthesis

Procedure and activation KRICT

Characterisation

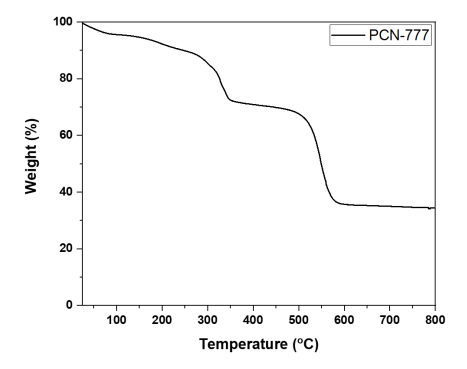


Fig. S7: Thermogravimetric curve recorded on as-synthesised PCN-777.

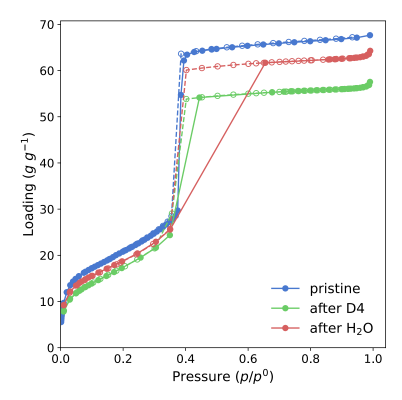


Fig. S8: Nitrogen sorption isotherms at 77 K for the pristine PCN-777, alongside those measured on samples after D4 and water sorption.

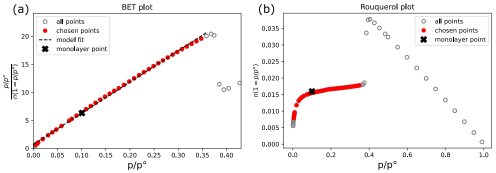


Fig. S9: BET and Rouquerol plots displaying selection of applicable isotherm points for the pristine PCN-777 isotherm.

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.

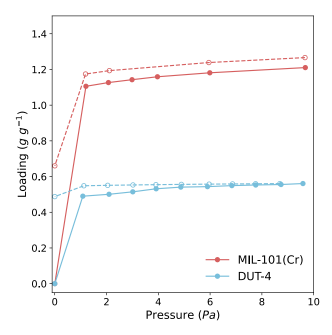


Fig. S10: D4 isotherms recorded on samples of MIL-101(Cr) (red) and DUT-4 (blue), used to validate our computational methodology for predicting total D4 capacity. Note the different desorption behavior (open symbols) of the two materials under secondary vacuum: partial desorption for MIL-101(Cr) and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

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

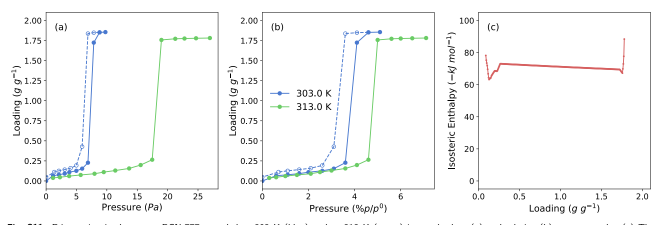


Fig. S11: D4 sorption isotherms on PCN-777 recorded at 303 K (blue) and at 313 K (green) in an absolute (a) and relative (b) pressure scale. (c) The calculated isosteric heat of adsorption as a function of D4 uptake.

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