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## Hypercrosslinked polymers for oil adsorption: the influence of porosity and fluorine incorporation

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The selective removal of oil and oil-based contaminants from water remains a critical challenge in environmental remediation. Here, we report a series of hypercrosslinked polymers with high surface areas and tuneable chemistries, achieving exceptional adsorption capacities for a variety of organic solvents, with a maximum capacity of  $>15 \text{ g g}^{-1}$  for chlorinated solvents. We describe how the adsorption capacities of the materials in pure organic solvents are governed by porosity rather than sample fluorine content and its associated hydrophobicity, challenging conventional design strategies. Oil/water separation tests of the most promising networks demonstrated the effective removal of toluene from water, achieving separation efficiencies of  $>99\%$ . The polymers also exhibit exceptional stability in organic solvents, allowing repeated use. This work establishes hypercrosslinked polymers as robust, scalable materials for efficient oil–water separation and advanced wastewater treatment.

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### Introduction

The remediation of oil pollution in aquatic environments presents broad technical and environmental challenges, owing to the complexity and scale of contamination events.<sup>1</sup> Marine transport, offshore drilling, and land-based runoffs collectively account for vast quantities of oil entering water bodies each year. Since 1970, an estimated 10 million tonnes of oil have been spilt into aquatic systems, including the release of around 4.9 million barrels of crude oil during the 2010 Deepwater Horizon incident.<sup>2</sup> The hydrophobic nature and low water solubility of oil-based contaminants mean they typically accumulate as a separate bulk phase, forming slicks or emulsions that are difficult to treat. Upon release, these hydrocarbons can spread over large surface areas, smothering coastlines, or spreading for hundreds of miles across bodies of water in a thin oil slick and disrupting delicate aquatic ecosystems.<sup>3</sup> Beyond ecological damage, oil spills also pose serious socioeconomic challenges and, as such, urgently require efforts toward more effective mitigation strategies.

Current oil remediation technologies face numerous limitations, particularly in separating immiscible oil from water. Skimmers and dispersants are commonly used, yet are energy-intensive, limited in effectiveness, and often cause secondary pollution.<sup>1</sup> Adsorbent materials capable of selectively capturing

large amounts of bulk-phase oils have garnered attention as attractive alternatives. Ideal sorbents should exhibit high uptake capacity, hydrophobicity, rapid adsorption, structural durability, and reusability. Achieving all these features simultaneously, while ensuring scalability and economic viability, remains a major challenge for adsorbent material design. Efforts have focused on developing various hydrophobic porous frameworks, including sponges,<sup>4,5</sup> membranes,<sup>6,7</sup> fabrics,<sup>8,9</sup> and aerogels,<sup>10</sup> However, many such systems are hampered by complicated synthesis routes, lack of durability, or challenging regeneration procedures.

Hypercrosslinked polymers (HCPs) may represent a promising alternative platform for oil sorption and water remediation. These amorphous polymer networks possess high surface areas, tuneable pore structures, swellability, and exceptional chemical stability, and can be synthesised from inexpensive, commercially available monomers.<sup>11–13</sup> HCPs have shown excellent performance in both gas phase adsorption, such as atmospheric water harvesting,<sup>14,15</sup> CO<sub>2</sub> capture,<sup>16</sup> and the capture of volatile organic compounds,<sup>17,18</sup> as well as in the adsorption of dissolved pollutants.<sup>19,20</sup> However, their potential for bulk-phase organic solvent uptake and direct oil/water separation remains relatively underexplored. Superhydrophobic monolithic adsorbents used for oil/water emulsion separation and oil recovery have been produced *via* the coating or anchoring of HCPs onto melamine formaldehyde sponges.<sup>5,21,22</sup> The coated sponges showed ultrahigh separation efficiency and outstanding durability. Elsewhere, Lin *et al.* fabricated hypercrosslinked microporous polymers *via* emulsion-templating, demonstrating that the incorporation of fluorine substantially enhanced both hydrophobicity and oil adsorption capacity.<sup>23</sup>

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Powdered or particulate adsorbents are particularly efficient in cleaning high viscosity or large-area thin oil spills when compared with monolithic adsorbents, however, the oil adsorption capacities of powders remain limited in comparison.<sup>24</sup> A powdered superhydrophobic covalent organic framework showed a crude oil uptake capacity of almost 12 g g<sup>-1</sup> after post-functional hydrophobisation *via* silane grafting.<sup>25</sup> Activated carbons have also shown promise for oil sorption, with lauric acid decorated biochar achieving an oil uptake capacity of 11 g g<sup>-1</sup> (ref. 26) and magnetite-loaded carbons offering facile separation using magnetic fields.<sup>27</sup> However, further work is needed in order to improve uptake capacities and unveil simpler routes to effective adsorbents.

In this work, we synthesise both non-functional and fluorine-containing HCPs by the simple self-polycondensation of 4,4'-bis(chloromethyl)biphenyl (BCMBP) and  $\alpha,\alpha'$ -dichloro-*p*-xylene (*p*DCX).<sup>28</sup> To investigate the impact of chemical composition on adsorption and separation performance, the fluorine-containing monomer 4-fluorobenzyl chloride (*p*FBC) is incorporated into the HCP structure in varying ratios. Given the hydrophobic nature of fluorine, its incorporation was expected to enhance oil adsorption and improve oil/water separation. The resulting materials are evaluated for their chemical composition, porosity, organic solvent uptake, and separation efficiency, offering insights into structure–property relationships in these HCP-based adsorbents.

## Experimental

### Materials

4,4'-Bis(chloromethyl)biphenyl (BCMBP, 95%),  $\alpha,\alpha'$ -dichloro-*p*-xylene (*p*DCX, 98%), FeCl<sub>3</sub> (97%), Sudan III ( $\geq 85\%$ ), 1,2-dichloroethane (DCE,  $\geq 99.0\%$ ), methanol ( $\geq 99.8\%$ ), toluene ( $\geq 99.7\%$ ), and *n*-heptane ( $\geq 99\%$ ) were purchased from Sigma-Aldrich. 4-Fluorobenzyl chloride (*p*FBC,  $>98\%$ ) and benzotri-fluoride (BTF,  $>98\%$ ) were purchased from Tokyo Chemical Industry. All chemicals were used as received.

### Hypercrosslinked polymer synthesis

Monomer(s) (details shown in Table 1) and DCE (30 mL) were added to a 100 mL two-neck round bottom flask and stirred at  $\sim 100$  rpm. After monomer dissolution, FeCl<sub>3</sub> (0.73 g, 4.5 mmol, 1.5 eq.) was added and the mixture was refluxed at

80 °C for 14 h. After cooling to room temperature, the resulting solid was filtered with a Büchner funnel, washed with 100 mL of methanol on the filter, and then by Soxhlet extraction with methanol for a further 22 h. The washed HCPs were placed in a fume cupboard overnight to allow excess solvent to evaporate, ground into a fine powder using a pestle and mortar, and dried in a vacuum oven at 80 °C overnight, yielding brown/dark brown powders. Networks derived from BCMBP and *p*DCX are named BP-HCP(-XF) and X-HCP(-XF), respectively, where XF represents the theoretical mol% of *p*FBC, if present.

### Characterisation

Fourier-transform infrared (FTIR) spectra were collected with a Tensor II FTIR spectrometer (Bruker) equipped with a Bruker Platinum-ATR Convenience sampling module at room temperature. Each spectrum was measured with 32 scans in the range of 400 to 4000 cm<sup>-1</sup> with a resolution of 4 cm<sup>-1</sup> and was recorded and analysed using the OPUS 7.5 software.

Thermogravimetric analysis (TGA) was conducted using a Discovery TGA (TA instruments). Between 3 to 10 mg of sample was placed into a platinum sample pan, equilibrated at 30 °C, then heated to 800 °C at a ramp rate of 10 °C min<sup>-1</sup> in air. Samples were held at the target temperature for 5 min before being allowed to cool. The TGA data were recorded and analysed using the Discovery TGA@Lab program in TRIOS software.

X-ray photoelectron spectroscopy (XPS) was performed on a Nexsa Photoelectron Spectrometer (Thermo Scientific). All measurements were performed using Al-K $\alpha$  X-rays with a spot size of 400  $\mu$ m. The survey spectra were taken at a pass energy of 200 eV from 1350 to 0 eV with a resolution of 1 eV. The high-resolution C 1s, O 1s, F 1s, and Cl 2p spectra were collected at a pass energy of 50 eV with a resolution of 0.1 eV. C 1s was collected in the range from 298 to 279 eV, O 1s from 545 to 525 eV, F 1s from 698 to 678 eV, and Cl 2p from 210 to 190 eV.

The N<sub>2</sub> adsorption–desorption isotherms were measured at  $-196$  °C using a 3Flex (Micromeritics), or a TriStar II (Micromeritics) for sample repeats. Before the measurement, 50 to 100 mg of sample was degassed at 120 °C under N<sub>2</sub> flow for 14 h using a FlowPrep 060 (Micromeritics). The networks' specific surface areas (SSA) were determined *via* the Brunauer–Emmett–Teller (BET) method in the relative pressure  $P/P_0$  range 0.05 to 0.3. Pore size distributions (PSDs) were determined from the N<sub>2</sub> adsorption branch of the isotherm using quenched solid density functional theory (QSDFT) and a carbon model with slit/cylindrical pores. The total pore volume  $V_P$ , micropore volume  $V_{Mic}$ , and mesopore volume  $V_{Mes}$  were determined from the cumulative pore volumes calculated using QSDFT in the relevant pore width ranges. When calculating  $V_P$  and  $V_{Mes}$ , isotherm points from  $P/P_0 > 0.97$  were excluded to avoid the effects of bulk N<sub>2</sub> condensation.

Dynamic vapour sorption (DVS) experiments were performed on a DVS-Discovery (Surface Measurement Systems). Approximately 10 mg of sample was weighed into a quartz crystal pan. Each experiment was performed at 25 °C and either deionised water (HPLC grade) or toluene (HPLC grade) was used to generate the desired relative pressure. Each ana-

**Table 1** Monomeric composition of all synthesised HCPs

Network	Monomer feedstock composition			
	Monomer 1	<i>n</i> (mmol)	Monomer 2	<i>n</i> (mmol)
BP-HCP	BCMBP	3	—	—
BP-HCP-25F	BCMBP	3	<i>p</i> FBC	1
BP-HCP-50F	BCMBP	3	<i>p</i> FBC	3
X-HCP	<i>p</i> DCX	3	—	—
X-HCP-25F	<i>p</i> DCX	3	<i>p</i> FBC	1
X-HCP-50F	<i>p</i> DCX	3	<i>p</i> FBC	3



lysis commenced at a relative pressure of 0.0  $P/P_0$  and was increased to the desired  $P/P_0$  for sorbate uptake. Isotherms were recorded up to 0.9  $P/P_0$ , using a step increment of 0.1. Each step was equilibrated until the weight change was below 0.005%  $\text{min}^{-1}$  prior to data point collection.

### Oil adsorption

To determine oil adsorption capacity, approximately 45 mg of HCP was suspended in solvent (10 mL) in a covered glass vial and stirred at 250 rpm for 1 h. After stirring, HCPs were vacuum filtered in a Büchner funnel to remove excess solvent, removed from the filter paper, and quickly reweighed to minimise solvent evaporation. The mass of the recovered adsorbents after drying was typically 35–45 mg, indicating a mass loss in filtration of up to 10 mg. The networks were then dried at 120 °C for at least 16 h and reweighed. Each measurement was repeated 5 times to ensure reproducibility, and adsorption capacity  $k_D$  was calculated using eqn (1):

$$k_D = \frac{m_A - m_D}{m_D} \quad (1)$$

where  $m_A$  and  $m_D$  are the weight of the sample after adsorption and after drying, respectively.

For comparison, the adsorption capacity  $k_B$  based on the dry weight of the sample before adsorption was calculated according to eqn (2), following previous reports:<sup>5,22,23,27</sup>

$$k_B = \frac{m_A - m_B}{m_B} \quad (2)$$

where  $m_B$  is the weight of the sample before adsorption.

### Oil/water separation experiments

To demonstrate the separation of immiscible organic solvent from water, toluene and distilled water were selected as organic and aqueous phases, respectively. To visualise the separation, toluene was dyed using fat-soluble dye Sudan III (0.01 g  $\text{mL}^{-1}$ ). The oil/water mixture was prepared in a 25 mL graduated cylinder using 0.5 mL of Sudan III dyed toluene and 20 mL of distilled water, followed by the addition of samples in portions while shaking gently until no red organic phase could be observed. The samples considered in this test were BP-HCP (88.1 mg) and BP-HCP-25F (87.0 mg). After adsorption, samples were removed by vacuum filtration and the oil/water separation efficiency SE was calculated using eqn (3), as reported by Sarkar *et al.*:<sup>29</sup>

$$SE = \frac{m_{Wr}}{m_{Wi}} \times 100\% \quad (3)$$

where  $m_{Wi}$  and  $m_{Wr}$  were the initial weight of water and the weight of the remaining water after separation, respectively. The adsorption efficiency AE of samples during oil/water separation was calculated using eqn (4):

$$AE = \frac{m_{St}}{m_{Sa}} \times 100\% \quad (4)$$

where  $m_{St}$  and  $m_{Sa}$  were the theoretical (based on their isolated adsorption capacities) and actual masses of the HCPs used in the separation experiment, respectively. Additionally, 20 mL of water alone was used as a control to measure any potential loss during vacuum filtration.

## Results and discussion

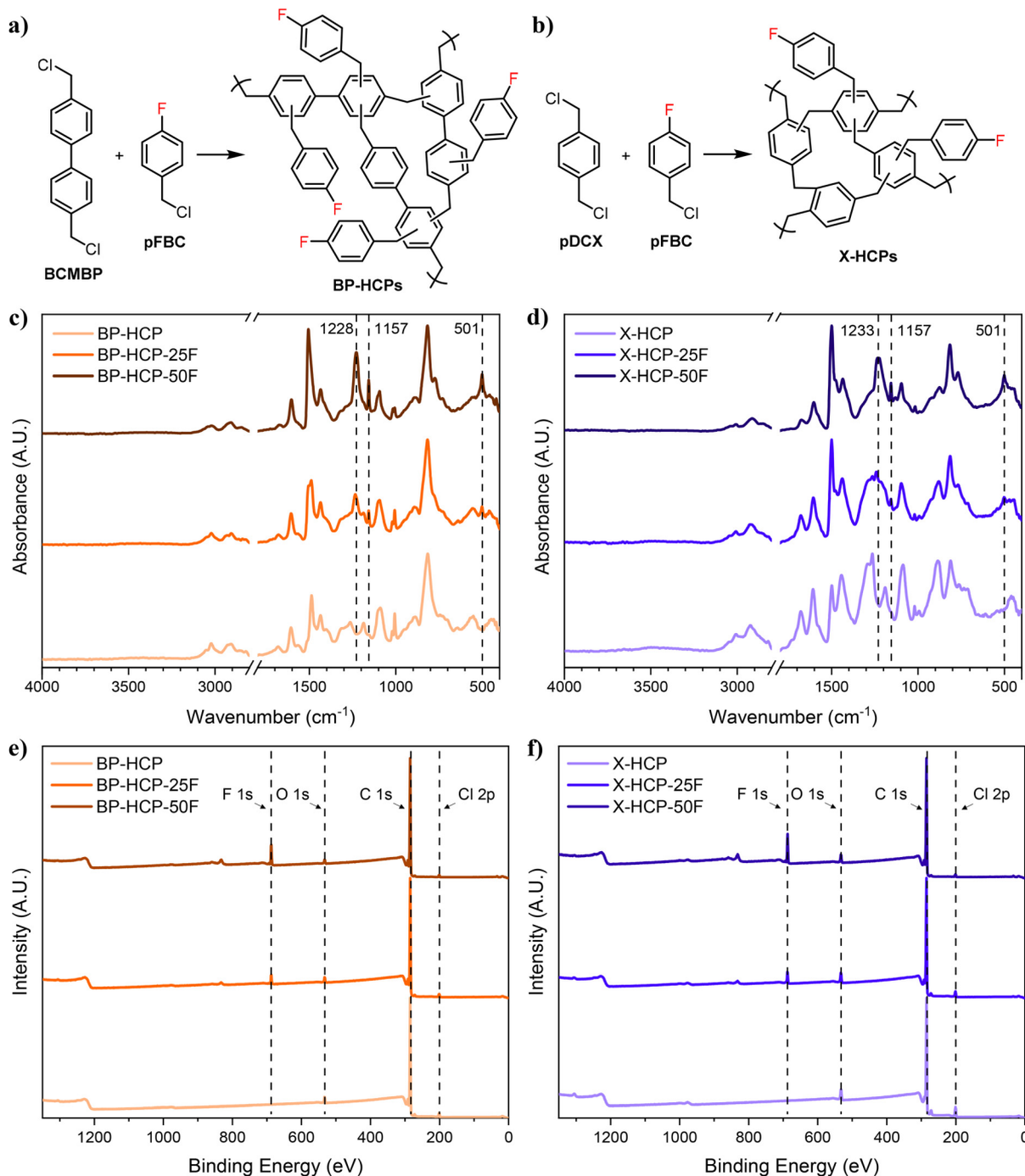
### Synthesis and characterisation of HCPs

Six HCPs were synthesised using either BCMBP or *p*DCX as a framework monomer/crosslinker and *p*FBC as a fluorine-containing monomer (Fig. 1a and b). All networks were synthesised *via* an  $\text{FeCl}_3$  catalysed Friedel–Crafts alkylation in 1,2-dichloroethane. Further synthesis details are provided in the Experimental section. The resulting polymers were named according to their monomeric composition, as detailed in Table 1. Networks derived from BCMBP, which contain a biphenyl-based skeleton, are designated BP-HCP(-XF), where X represents the theoretical mol% of *p*FBC. For networks synthesised from *p*DCX, the notation X-HCP(-XF) is used. All HCPs were brown powders produced in yields of >86%. We also attempted the self-polycondensation of *p*FBC alone, however poor yields prevented us from pursuing this polymer further.

The FTIR spectra of BP-HCPs (Fig. 1c) displayed the characteristic absorption peaks of the 4,4'-dimethylbiphenyl framework, with BP-HCP-25F and BP-HCP-50F exhibiting three additional peaks at 1228, 1157 and 501  $\text{cm}^{-1}$ . The peaks at 1228 and 1157  $\text{cm}^{-1}$  were assigned to C–H in-plane bending vibration in the fluorinated aromatic ring, and the peak at 501  $\text{cm}^{-1}$  was assigned to the characteristic  $\text{C}_{\text{Ar}}\text{–F}$  bending vibration, confirming the incorporation of fluorine. The  $\text{C}_{\text{Ar}}\text{–F}$  stretching vibration expected at around 1330  $\text{cm}^{-1}$  was likely overlapped by other signals and was not observed. Increasing *p*FBC content led to a greater intensity of peaks associated with fluorine, indicative of BP-HCP-50F containing the highest fluorine content of the BP-HCPs. A similar trend was observed in the X-HCP series (Fig. 1d), where X-HCP-25F and X-HCP-50F exhibited the same three additional peaks at 1233, 1157 and 501  $\text{cm}^{-1}$ , relative to X-HCP, confirming successful *p*FBC incorporation. Again, peaks associated with fluorine increased with expected *p*FBC content, indicating X-HCP-50F contained the highest fluorine content of the X-HCPs. The FTIR spectra of the BP-HCPs after 6 months storage in ambient conditions were almost the same as their corresponding pristine spectra (Fig. S1), confirming the excellent stability of HCPs.

XPS was conducted to understand the surface chemistry and relative fluorine content of all HCPs. Survey spectra (Fig. 1e and f) confirmed the presence of fluorine in all networks produced from monomer feeds containing *p*FBC, as well as the presence of residual terminal chlorine. The *p*FBC was incorporated effectively with increasing *p*FBC monomer ratios, with BP-HCP-50F and X-HCP-50F reaching fluorine contents of  $6.0 \pm 0.4$  at% and  $7.1 \pm 0.1$  at%, respectively. The fluorine content of all networks was in slight excess of the theoret-





**Fig. 1** (a) Reaction scheme for the synthesis of BP-HCPs, (b) reaction scheme for the synthesis of X-HCPs, (c) FTIR spectra of BP-HCPs, (d) FTIR spectra of X-HCPs, (e) XPS survey spectra of BP-HCPs, and (f) XPS survey spectra of X-HCPs. FTIR spectral data between 1800 and 2800  $\text{cm}^{-1}$  are omitted for clarity as no peaks were present in this region.

tical value (Table S1), likely due to the molecule's ability to effectively stabilise the carbocation formed upon the loss of Cl, aiding crosslink formation to the BCMBP monomer. However, the fluorine substituent is weakly deactivating toward electrophilic aromatic substitution and likely suppresses further propagation upon incorporation into the polymer. The presence of oxygen in all HCPs was ascribed to the attack of trace

water at the carbocation generated during the Friedel–Crafts reaction. In the fluorine-decorated HCPs, the high-resolution F 1s spectra displayed a peak at 687.5 eV, assigned to the C<sub>Ar</sub>-F in the pFBC (Fig. S2). An additional peak at ~694 eV was observed, which does not correspond to known C-F species but is close to the binding energy of SF<sub>6</sub> (693.5 eV).<sup>30</sup> A plausible explanation for this feature may be residual sulfur con-



tamination, a common impurity in organic materials originating from crude oil cracking.

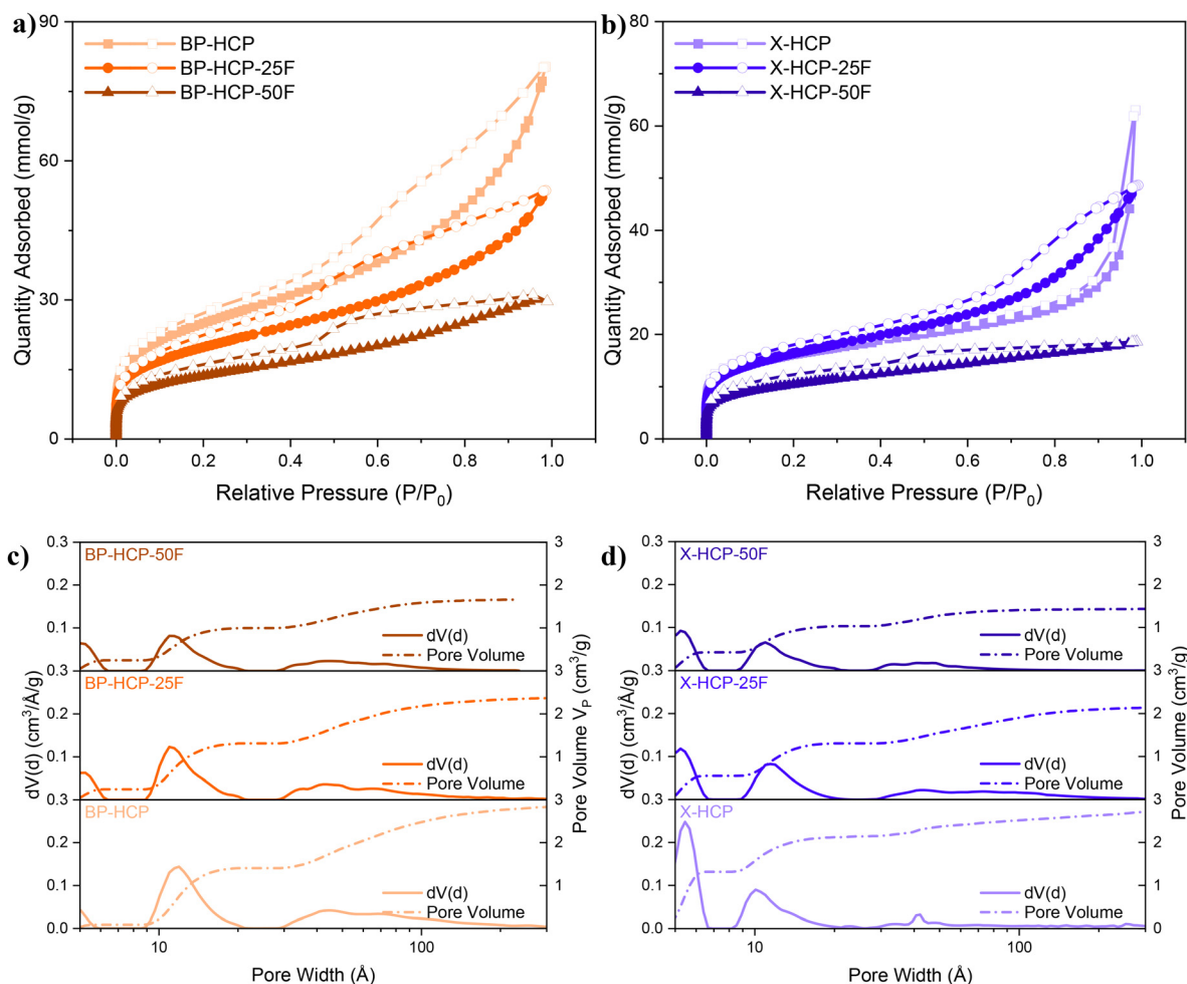
The high-resolution C 1s spectra of BP-HCPs and X-HCPs were deconvoluted into three peaks  $\sim 284.8$  eV, 287.0 eV and 291.4 eV, corresponding to  $sp^2/sp^3$  C, the combination of C–O and C–Cl, and a low-intensity  $\pi \rightarrow \pi^*$  shake-up feature,<sup>30</sup> respectively (Fig. S3). In fluorine-functionalised HCPs, C–F was expected to produce a peak at 286.0–286.5 eV but was not clearly distinguishable due to spectral overlap with C–O and C–Cl.

The TGA curves of all HCPs showed similar thermal stability (Fig. S4). All networks increased in weight at around 240 to 260 °C, indicative of oxidation,<sup>15</sup> before the onset of thermal decomposition at around 340 to 360 °C. All samples were decomposed completely at approximately 580 to 600 °C. The introduction of fluorine into the HCP networks did not noticeably alter their thermal stability.

$N_2$  adsorption–desorption isotherms of all samples were collected at  $-196$  °C. According to IUPAC classification,<sup>31</sup> all networks displayed a combination of type I and type IV isotherms (Fig. 2a and b), consistent with typical HCPs.<sup>14,32</sup> The

type I behaviour demonstrated by a steep uptake at low  $P/P_0$  indicates microporosity, while the type IV component identified by hysteresis during desorption reflects mesoporous capillary condensation.

With increasing *p*FBC content, BP-HCPs exhibited a steady decline in porosity. The BET-specific surface areas ( $SSA_{BET}$ ) decreased from  $1748 \pm 78$  in BP-HCP to  $960 \pm 114$   $m^2 g^{-1}$  in BP-HCP-50F, with corresponding reductions in the total pore volume  $V_P$ , micropore volume  $V_{Mic}$ , and mesopore volume  $V_{Mes}$  (Table 2). On the other hand, porous properties remained relatively unchanged between X-HCP and X-HCP-25F, with  $SSA_{BET}$  of  $1089 \pm 105$   $m^2 g^{-1}$  and  $1140 \pm 94$   $m^2 g^{-1}$ , respectively, and an apparent increase in  $V_{Mes}$  from  $0.61$   $cm^3 g^{-1}$  to  $0.84$   $cm^3 g^{-1}$  was observed. However, at the highest *p*FBC loading (X-HCP-50F), porosity declined significantly, resulting in a  $SSA_{BET}$  of  $691 \pm 69$   $m^2 g^{-1}$  and  $V_{Mes}$  of  $0.40$   $cm^3 g^{-1}$ . Overall, X-HCP networks derived a larger portion of their  $V_P$  from micropores than their BP-HCP counterparts (Table 2). Pore size distributions were determined using the QSDFT model, displaying an abundance of  $\sim 1$  nm wide micropores and broad mesoporosity in all networks (Fig. 2c and d).



**Fig. 2** (a)  $N_2$  adsorption–desorption isotherms, and (c) pore size distribution and cumulative pore volume of BP-HCPs. (b)  $N_2$  adsorption–desorption isotherms, and (d) pore size distribution and cumulative pore volume of X-HCPs.



**Table 2** Porous properties of all HCPs including BET-specific surface areas  $SSA_{\text{BET}}$ , total pore volumes  $V_{\text{p}}$ , micropore volume  $V_{\text{Mic}}$ , mesopore volume  $V_{\text{Mes}}$ , and ratio of  $V_{\text{Mic}}$  to  $V_{\text{Mes}}$ 

Sample	$SSA_{\text{BET}}$ ( $\text{m}^2 \text{g}^{-1}$ )	$V_{\text{p}}$ ( $\text{cm}^3 \text{g}^{-1}$ )	$V_{\text{Mic}}$ ( $\text{cm}^3 \text{g}^{-1}$ )	$V_{\text{Mes}}$ ( $\text{cm}^3 \text{g}^{-1}$ )	$V_{\text{Mic}}/V_{\text{Mes}}$
BP-HCP	$1748 \pm 78$	2.85	1.40	1.45	1.0
BP-HCP-25F	$1403 \pm 182$	2.38	1.31	1.07	1.2
BP-HCP-50F	$960 \pm 114$	1.66	0.99	0.67	1.5
X-HCP	$1089 \pm 105$	2.73	2.12	0.61	3.5
X-HCP-25F	$1140 \pm 94$	2.14	1.30	0.84	1.6
X-HCP-50F	$691 \pm 69$	1.43	1.03	0.40	2.6

### Oil adsorption performance and oil/water separation performance of HCPs

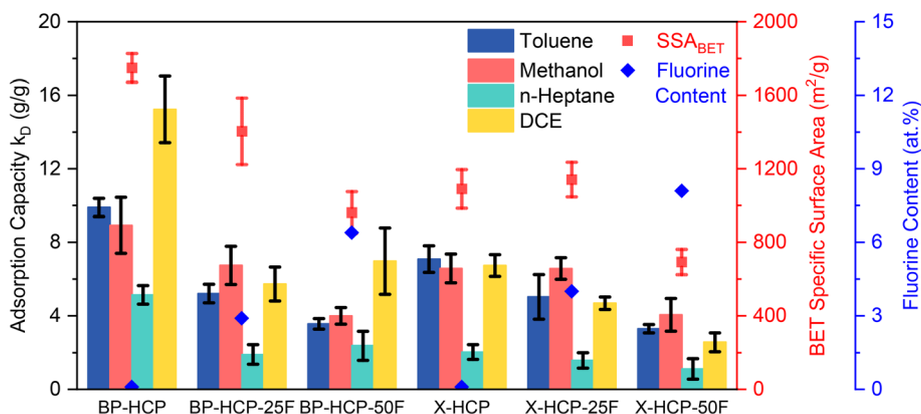
To evaluate the adsorption performance of HCPs toward organic solvents, toluene, methanol, *n*-heptane, and DCE were selected as representative compounds of aromatic hydrocarbons, alcohols, linear alkanes, and halides, respectively. Adsorption experiments were conducted by suspending HCP samples in pure solvent and stirring for 1 h before removal of excess solvent by vacuum filtration. The HCPs were weighed immediately after filtration to minimise evaporation losses and then again after complete drying.

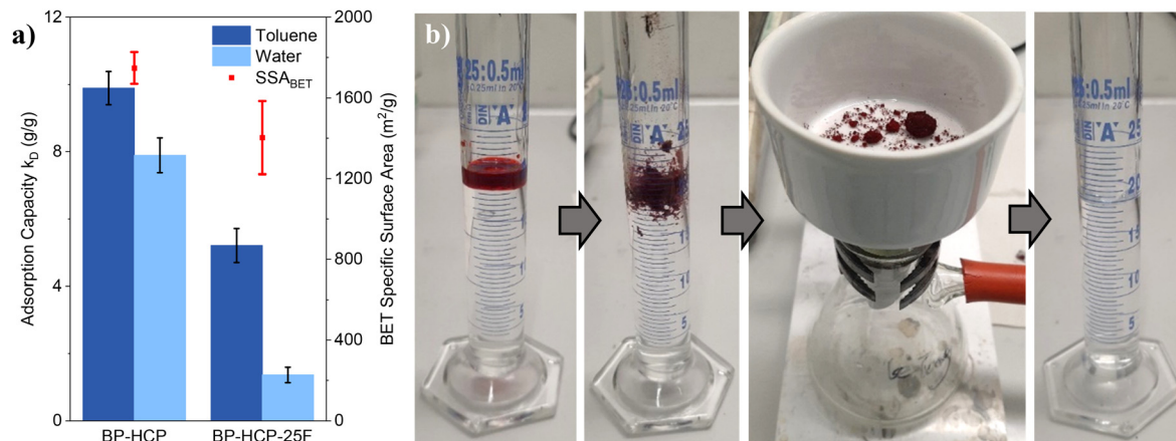
Adsorption capacity was calculated using two methods; the first,  $k_{\text{B}}$ , follows conventional literature approaches and considers the initial sample weight before adsorption. However, as the powdery HCPs in this work were recovered by vacuum filtration, some sample loss during filtration led to an underestimation of adsorption capacity. To correct for this loss, a second adsorption capacity,  $k_{\text{D}}$ , was calculated using the dry sample weight after adsorption, ensuring fairer comparison between networks. The  $k_{\text{D}}$  values were consistently only slightly higher than the corresponding  $k_{\text{B}}$  values (Table S2) and, therefore, remain comparable to  $k_{\text{B}}$  values in the literature. As such,  $k_{\text{D}}$  is used in the following discussion as a measure of adsorption capacity.

The adsorption capacities of the HCPs were compared with their  $SSA_{\text{BET}}$  and  $V_{\text{p}}$  (Fig. 3 and Table S2). In the BP-HCP networks, adsorption capacities toward toluene and methanol decreased in line with  $SSA_{\text{BET}}$  and  $V_{\text{p}}$ . Specifically, adsorption

dropped from  $9.89 \pm 0.49$  to  $3.55 \pm 0.29 \text{ g g}^{-1}$  for toluene and from  $8.92 \pm 1.52$  to  $3.99 \pm 0.45 \text{ g g}^{-1}$  for methanol, when comparing BP-HCP and B-HCP-50F. However,  $k_{\text{D}}$  did not follow the same trend for *n*-heptane and DCE; although both BP-HCP-25F and BP-HCP-50F exhibited lower adsorption than BP-HCP, a slight increase in uptake was observed in BP-HCP-50F. The higher fluorine content in BP-HCP-50F may enhance DCE adsorption *via* halogen interactions, while also improving *n*-heptane uptake due to its strong hydrophobicity. The hydrophobicity of BP-HCP-50F was assessed and compared to BP-HCP by the collection of dynamic vapour sorption isotherms of both toluene and water (Fig. S5). Although BP-HCP displayed a higher total uptake capacity for both water and toluene, the ratio of toluene:water total uptake was significantly larger in BP-HCP-50F (20:1) compared to BP-HCP (6:1), indicating increased hydrophobicity with the inclusion of fluorine.

Among the X-HCP polymers, the adsorption capacities for all four solvents again correlated closely to  $SSA_{\text{BET}}$  and  $V_{\text{p}}$  (Table S2), remaining similar across X-HCP, X-HCP-25F, and decreasing for X-HCP-50F. In the X-HCPs, no positive effect on adsorption capacity was observed with the inclusion of fluorine. Overall, adsorption capacities correlated more closely with porosity rather than fluorine content, reinforcing the importance of pore structure in organic solvent uptake. The solvent adsorption capacity of our best performing material, BP-HCP, is compared with various oil adsorbents in the literature in Table S3.

**Fig. 3** Adsorption capacities  $k_{\text{D}}$  of HCPs for different organic solvents compared to their BET-specific surface area  $SSA_{\text{BET}}$  and fluorine content.



**Fig. 4** (a) Adsorption capacities  $k_D$  of BP-HCP and BP-HCP-25F toward toluene and water, compared to their BET-specific surface areas. (b) Removal process of Sudan III dyed toluene from water using BP-HCP-25F.

The adsorption capacities  $k_D$  of BP-HCP and BP-HCP-25F toward distilled water were  $7.89 \pm 0.52$  and  $1.36 \pm 0.23$   $g\ g^{-1}$ , respectively (Fig. 4a). The water adsorption performance of BP-HCP-25F was significantly lower than that of BP-HCP, caused by the combination of a reduction in  $SSA_{BET}$  and  $V_P$ , as well as the hydrophobicity of the introduced *p*-fluorobenzyl groups, which may hinder water infiltration into the HCP pores.<sup>24,26</sup> Due to good apparent oil/water selectivity in isolated systems, BP-HCP and BP-HCP-25F were selected for testing in toluene/water separation experiments.

To visualise the oil/water separation process, Sudan III dye was added to toluene, and a model oil/water mixture was prepared by adding 0.5 mL of dyed toluene to 20 mL of distilled water. Both BP-HCP and BP-HCP-25F successfully adsorbed and removed the toluene from the water (Fig. 4b), demonstrating their applicability as adsorbents for oil/water separation. The adsorption efficiencies (AE) of BP-HCP and BP-HCP-25F reached up to 49.5% and 95.1%, respectively. The improved AE of BP-HCP-25F suggests that the increased hydrophobicity of the network improves efficiency in the toluene/water separation; however, this is more likely due to the addition of excess BP-HCP. It should be noted that this experiment was conducted as a proof of principle to observe oil/water separation

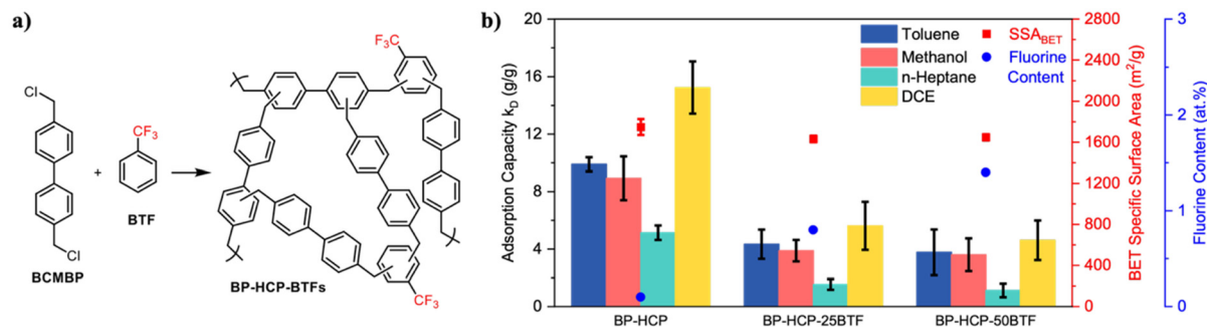
visually. We will seek to determine the limits of oil/water separation in future work.

The separation efficiencies (SE) of BP-HCP and BP-HCP-25F were determined by measuring the weight difference of the aqueous phase before and after separation, with both samples achieving 96% (Fig. S6). To account for potential water loss, a control experiment was conducted using water alone, allowing for the calculation of the actual separation efficiencies ( $SE_a$ ) according to eqn (5):

$$SE_a = 100\% - SE_c + SE_m \quad (5)$$

where  $SE_c$  represents the separation efficiency of the control and  $SE_m$  the separation efficiency of the sample. After correcting for filtration-induced water loss, the  $SE_a$  of BP-HCP and BP-HCP-25F were  $99.5 \pm 0.4\%$  and  $99.4 \pm 0.4\%$  (Fig. S6), respectively.

Despite the difference in oil/water selectivity between BP-HCP and BP-HCP-25F in isolated systems, the  $SE_a$  was unchanged in the two-phase system, likely due to the general preference of both networks for the organic phase over water, as well as differences in experimental conditions compared with the ideal adsorption measurements.



**Fig. 5** (a) Reaction scheme for the synthesis of BP-HCPs. (b) Adsorption capacities  $k_D$  of BTF containing HCPs toward various organic solvents, their BET-specific surface areas  $SSA_{BET}$ , and fluorine contents.



## Recovery of HCP adsorbents

The HCPs exhibited excellent resistance to organic solvents and water, as confirmed by FTIR spectra and porous properties of the recovered samples (Fig. S7 and Table S4). Minimal changes in their chemical composition and  $SSA_{\text{BET}}$  after adsorption suggest that the structural integrity of HCPs remained intact, highlighting their suitability for the adsorption of oil from water and adsorbent recycling. Further details are shown and discussed in the SI.

To more thoroughly assess the impact of fluorine incorporation on oil adsorption in pure solvent systems, we also produced HCPs using trifluorotoluene (BTF) in place of *p*FBC in a BP-HCP series of networks (reaction scheme shown in Fig. 5a). A similar trend was observed in these networks to those discussed herein, with respect to their characterisation, adsorption of organics (Fig. 5b), and regeneration (complete data set shown in section 2 of the SI). In the BTF series, the strongly electron-withdrawing  $-\text{CF}_3$  significantly reduces electrophilic reactivity, resulting in significantly reduced incorporation into the network compared to theoretical values (Table S5). Nevertheless, fluorine incorporation into HCPs again did not appear to benefit organic adsorption and, overall, BP-HCP remained the most promising candidate for oil/water separation.

## Conclusions

In this study, a series of hypercrosslinked polymers were synthesised containing various concentrations of 4-fluorobenzyl chloride. The synthesis approach was facile and yielded networks with BET-specific surface areas up to  $1750 \text{ m}^2 \text{ g}^{-1}$ , high thermal stability, and excellent resistance to organic solvents and water. Despite the controlled introduction of fluorine, the adsorption of pure organic solvents was primarily governed by porosity rather than chemical composition. Among the HCP networks, BP-HCP exhibited the highest porosity, with a surface area of  $1750 \text{ m}^2 \text{ g}^{-1}$  and a total pore volume of  $2.9 \text{ cm}^3 \text{ g}^{-1}$ , correlating well with its superior adsorption capacity for all solvents:  $9.9 \text{ g g}^{-1}$  for toluene,  $8.9 \text{ g g}^{-1}$  for methanol,  $5.0 \text{ g g}^{-1}$  for *n*-heptane, and  $15.2 \text{ g g}^{-1}$  for DCE. Furthermore, BP-HCP demonstrated excellent toluene/water separation with a separation efficiency of  $>99\%$ . Our results confirm that fluorine incorporation did not enhance the network capacities for solvent adsorption, reinforcing that pore volume and surface area are most crucial for the adsorption process. These findings highlight BP-HCP's potential as a robust, reusable adsorbent for high-capacity oil/water separation, while also emphasising the importance of pore structure in the design of high-performance HCP-based oil/solvent adsorbents.

## Conflicts of interest

There are no conflicts to declare.

## Data availability

The data supporting this article have been included as part of the SI.

The SI includes additional experimental data, tables, and figures supporting the main text. It provides XPS-derived surface elemental compositions, FTIR spectra after long-term storage, and detailed XPS core-level spectra for all polymers. Additional TGA curves,  $\text{N}_2$  adsorption–desorption isotherms, pore size distributions, and adsorption capacity data are presented, along with dynamic vapour sorption isotherms and oil/water separation efficiency measurements. Recovery studies are detailed through FTIR and porosity analyses of reused adsorbents. Comparative adsorption performance with literature materials is tabulated. Section 2 presents synthesis, characterisation, porosity, adsorption, and recovery data for trifluorotoluene-derived HCPs, BP-HCP-BTFs. See DOI: <https://doi.org/10.1039/d5lp00081e>.

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