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A nanotrap infused ultrathin hybrid composite material for rapid and highly selective entrapment of ⁹⁹TcO₄⁻†

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⁹⁹Tc is one of the potentially toxic radioactive substances owing to its long half-life and a high degree of environmental mobility. Hence, the sequestration of ⁹⁹Tc from radioactive waste has become enormously important and a contemporary research priority. However, selective extraction of this species in its stable oxoanionic form $(^{99}\text{TcO}_4^{-})$ is very challenging on account of bottlenecks such as low charge density, less hydrophilic nature, etc. Herein, an ultrathin hybrid composite material has been strategically designed and fabricated by covalent anchoring of a chemically stable amino functionalized nanosized cationic metalorganic polyhedron with a positively charged robust ionic covalent organic framework. The resulting thinlayer-based hybrid composite presented multiple exfoliated exposed interactive sites, including a Zr(iv)secondary building unit, amine and triaminoquanidine functional groups, which can selectively interact with TcO₄ oxoanions through a synergistic combination of electrostatic, H-bonding and various other supramolecular interactions. Thus synthesized function-tailored composite, by virtue of its multiple unique characteristics, manifested an ultrafast and very selective, high distribution coefficient ($\sim 10^6$ mL g⁻¹), as well as recyclable entrapment of TcO_4^- oxoanions from the complex mixture of superfluous (\sim 5000-fold) other interfering anions in both high and ultra-trace concentrations along with simulated nuclear waste and from different water systems. Dynamic flow-through experiments were conducted with the membrane of the hybrid material in simulated wastewater, which reduced the concentration of ReO₄-(surrogate of radioactive TcO_4) to below the WHO permissible level with rapid sequestration kinetics and excellent selectivity over excessive competing anions.

Introduction

⁹⁹Tc prevails as one of the most hazardous fission products in the nuclear fuel cycle because of its long half-life, potential radiation threat, high environmental mobility and strong redoxactive nature. To is a long lived $(t_{1/2} = 2.13 \times 10^5 a)$ β -emitting radionuclide (E = 294 keV) that is both chemically toxic and hazardous.2-4 The low charge density and high symmetry of the oxoanionic species of ⁹⁹Tc (⁹⁹TcO₄⁻) result in its extremely high solubility in water (11.3 M at 20 °C). Hence, 99TcO₄ can easily migrate through the natural water system and its migration is not affected by the abundance of the majority of natural materials. Therefore, attempts to develop several effective technologies toward sequestration and management of radioactive waste from water are governed by concerns about the environment and human health as well as the establishment of a safe nuclear sector. Among the explored technologies, solid sorbents operating via adsorption followed by ion exchange have demonstrated significant promise as established TcO₄ remediation technologies due to their easy, safe, low-cost and robust mechanism processes. 6-12 Along this line, several materials, that include ion-exchange resins and polymeric networks,13,14 inorganic cationic materials, and cationic metal-organic frameworks (MOFs),15 have been explored as 99TcO4 segregating materials. Albeit, most cases do not present the desired combination of a suitable sorbent material and are often associated with either suboptimal capacity12,16,17 and selectivity18 or

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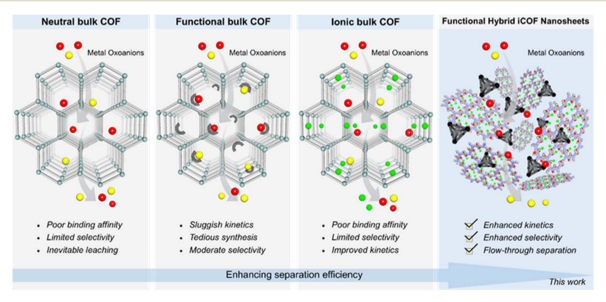
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poor removal kinetics. 12 Therefore, the overall efficiency of these materials as potential adsorbents is limited, particularly due to their poor selectivity. This is because plenty of co-existing competing anions, such as NO₃⁻, SO₄²⁻, CO₃²⁻, ClO₄⁻, Br⁻ and Cl-, are in large excess with TcO4-, which can act as interfering analytes, in both natural water systems and nuclear wastewater.13-15 On the other hand, although commercially available polymeric anion exchange resins have a significant advantage in TcO₄ uptake selectivity, their poor radiation resistance has thwarted their practical applicability.19 Additionally, TcO₄ can be extracted from aqueous solution with better adsorption kinetics, uptake capacity, reasonable selectivity, and strong radiation resistance capability using cationic metal-organic frameworks (MOFs). 15,20-22 However, the major disadvantage associated with these materials is their instability in chemical environments essential for practical use. Consequently, considerable attention has been paid to the development of effective regenerable materials with significantly better segregation capability in order to overcome these issues. Having said that, although covalent organic frameworks (COFs), in various capacities such as neutral/unmodified, ionicity or chelating functional groups have shown capability toward extracting toxic metal oxo-anions from water they are thwarted from real time utility by their moderate sorption capacity as well as suboptimal trapping kinetics and selectivity (Scheme 1). 10,23-27 Therefore, as an ongoing battle, improvement in terms of rapid kinetics, high selectivity and capacity towards TcO4oxoanions at both high and ultra-trace concentrations, from nuclear wastewater sources to point-of-use, is still needed for practical application, which is mostly unexplored due to the lack of rational synthetic design strategies.

In this sense, innovative functional porous materials have been identified as potential grounds for a variety of cutting-edge applications due to their predesigned adjustable reticular structure. The synthesis of composite materials can result in the

combined benefits and maximized performance of several individual functional materials.²⁸⁻³¹ Toward the development of innovative and multipurpose platforms in order to achieve desired applications, hybrid composite materials have come to the forefront and garnered a lot of research attention globally.32-34 Although, there has been remarkable advancement in the field of composite materials in recent years,35 hybrid composite materials have not been that much explored yet.³⁶ Along this line, the combined qualities of two individual active porous material based hybrids enable the composite to function as an excellent adsorbent.37 In this context, metal-organic polyhedra (MOPs), zero-dimensional crystalline materials, made up from metal ions (or metal clusters) and organic linkers, and covalent organic frameworks (COFs), crystalline porous organic materials which are assembled from covalent linking of pure organic linkers, could be an ideal platform as two individual porous materials for the hybrid.38,39 These two are widely known porous materials for a variety of applications such as gas storage, sensing, catalysis, photoelectric conversion, energy storage and biological activities.38-40 However, due to unavoidable aggregation-induced active site obstruction upon guest removal in the solid-state, MOPs are often unable to attain their full potential in the aforementioned applications.41 To date, only a few strategies have been documented to overcome such a prime issue by encasing MOPs inside porous materials (silica and MOFs)42,43 but hybridization of MOPs with other porous materials such as COFs has remained a difficult challenge. Nevertheless, covalent organic framework (COF) powder hybridized with cationic cages has been found to exhibit improved kinetics and moderate selectivity in the removal of different oxoanions from water.36 However, the intrinsically twodimensional stacked architecture of the neutral COF matrix of the nanocomposite limits the full exposure or utilization of their complete surface for adsorption, especially towards the active nanotraps present inside the pores. Moreover, the robust



Scheme 1 Schematic illustration of the advantages of MOPs@iCOF hybrid composite nanosheet materials over pristine neutral bulk COF and functionalized and ionic COF materials, towards rapid and selective oxoanion sequestration.

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heterogeneous nature with non-monolithic applicability of bulk hybrid COF powder has thwarted the large-scale practical flowthrough separation application. Now, to address these cardinal issues, we reinvestigated the molecular-level engineering of the host matrix to fabricate an effective composite material for highly efficient metal-oxoanion sequestration. In this regard, the construction of selective binding site containing active nanotrap (MOP) embedded COF-based ultrathin stable hybrid composite materials could be an ideal solution for device-based (such as membranes) excellent dynamic sequestration of specific metal oxoanions with ultrafast kinetics and very selective efficiency (Scheme 1). Considering this, here we hypothesized that the utilization of a triaminoguanidine (TAG) moiety based ionic COF (iCOF) could be an excellent choice as an efficient host matrix for threading guest-MOPs through the hybridization strategy to fabricate multifunctionalized hybrid nanocomposites. This is particularly important owing to their thin 2D structure (exfoliated) and high stability and the presence of excess free counter anions, which can be exchanged with targeted TcO₄ oxoanions.

Taking all the above facts into consideration, herein an efficient anion exchangeable ultrathin hybrid composite material (IPcomp-8, (where "IP" stands for IISER Pune and "comp" stands for composite)) has been synthesized through covalent grafting of amino-functionalized nanosized cationic Zr-MOP with a chemically stable positively charged COF (iCOF) (Fig. 1). We have strategically designed and grafted the NH₂-

functionalized cationic MOPs into the ionic COF matrix, where the excess of exchangeable Cl⁻ anions from both host-iCOF and guest-Zr-MOP, free -NH2 functional groups of the MOP, and the triaminoguanidine (TAG) moiety of the iCOF accelerate the anion exchange process due to the electrostatic and hydrogenbonding interactions with the radioactive oxoanions (TcO₄⁻) (Fig. 1). Most importantly, the Cp₃Zr₃O(OH)₃ secondary building unit (SBU) of the MOP can selectively interact with TcO₄ through an exchange with its labile hydroxyl groups. 44-46 Also, the iCOF matrix tightly harboured the MOP molecules keeping most of the active sites well exposed for favorable interactions. In addition to this, the covalent positioning of the cationic MOP into the cationic COF architecture results in easy and self-exfoliation into ultrathin hybrid nanosheets, owing to the strong interlayer ionic repulsion at play. This further allows full exposure of all the multifunctional active sites originating from both the Zr-MOP and iCOF matrix, further triggering rapid mass diffusion and high selectivity towards target specific TcO₄ anions in water. Therefore, IPcomp-8 was found to exhibit ultrafast and highly selective separation of radioactive TcO₄ or ReO₄ (surrogate) anions from the contaminated samples in the presence of large excesses (\sim 10 to \sim 5000 fold) of other competing anions in both high (~50 ppm) and ultralow (~1000 ppb) concentrations. The composite also demonstrated TcO₄ entrapment from a wide range of pH and various irradiation doses with fast adsorption kinetics and the sorption was found to be reproducible. It could significantly lower the

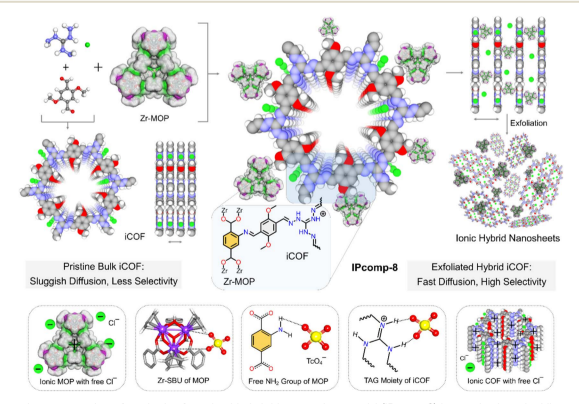


Fig. 1 Schematic representation of synthesis of an ultrathin hybrid composite material (IPcomp-8) by covalently embedding amino group functionalized Zr-MOP with iCOF and pristine iCOF. Various multifunctional features of IPcomp-8 showing ion-exchange electrostatic and functional group mediated interactions with radioactive oxoanions (color code: gray: carbon, white: hydrogen, red: oxygen, green: chlorine, light blue: nitrogen, violet: zirconium).

concentration of $\mathrm{ReO_4}^-$ to well below the WHO-set permitted level for drinking water (10 ppb).¹ Moreover, taking advantage of ultrathin nanosheets of the developed composite, we further fabricated a hybrid membrane, which demonstrated a highly efficient flow-through $\mathrm{ReO_4}^-$ extraction from contaminated water. Such a swift and effective adsorption performance by cooperative multifunctionalities of IPcomp-8 was further supported by several experimental and theoretical calculation studies. Our findings point to a promising area for tackling issues with $\mathrm{TcO_4}^-$ management and nuclear waste around the world.

Results and discussion

Initially, a discrete, chemically and hydrolytically stable, aminogroup functionalized, nanosized (~2 nm) cationic MOP ({[Cp₃- $Zr_3O(OH)_3]_4(NH_2-BDC)_6\}\cdot Cl_4$) was synthesized by reacting Zr(IV)metal with a 2-aminoterephthalic acid ligand and was characterized according to the previous literature report (Scheme S1 and Fig. S1-S5†).47 The triaminoguanidine (TAG) functionalized chemically robust cationic COF (iCOF) was also synthesized and thoroughly characterized following previously reported literature (Schemes S2, S3 and Fig. S6-S9†). 48 After synthesizing these pristine host matrix (iCOF) and guest nanotraps (Zr-MOP), the cationic Zr-MOP/iCOF hybrid material (IPcomp-8) was fabricated via a multicomponent condensation reaction by reacting NH2-functionalized Zr-MOP with the precursors of iCOF by a solvothermal method (Fig. 1) (see the ESI† for a detailed synthetic procedure). The free amino groups of the MOP can serve as an extra precursor, to react with dimethoxyterephthalaldehyde and triaminoguanidine monomers of the iCOF to fabricate covalently linked hybrid composite porous materials (IPcomp-8). By varying the weight of the MOP content, a series of hybrid materials have been synthesized. After the synthesis, thorough characterization studies of the hybrid composite have been performed in order to explore its detailed structural insights. The structural analysis of the composite material was deduced through a combination of powder X-ray diffraction (PXRD), Fourier transform infrared spectroscopy (FT-IR), X-ray photoelectron spectroscopy (XPS), nitrogen gas sorption studies, field emission scanning electron microscopy (FESEM), transmission electron microscopy (TEM), atomic force microscopy (AFM), and others. The formation of the crystalline structure of the composite material was established from PXRD studies (Fig. 2a). The PXRD analysis of IPcomp-8 indicated identical diffraction peaks as the pristine iCOF, further validating the formation and retention of the crystalline structural integrity of the iCOF backbone in the composite.48 However, a decrease in intensity of the peak corresponding to the 100 plane and the broadening of the 001 plane peak have been observed. This could be due to the random displacement of the 2D layers of the iCOF matrix in IPcomp-8. As a result, the reflection corresponding to the 100 plane becomes weak. The broader peak at 2θ ($\sim 25^{\circ}$) is due to the lack of π - π stacking between the iCOF layers of IPcomp-8, which is substantially influenced by the reduction of stacked layers in the composite. 49 FT-IR analysis was utilized to investigate the formation of the

composite between the guest NH_2 -MOP and host iCOF matrix. The FT-IR spectrum of IPcomp-8 indicated the appearance of a new prominent absorption band at \sim 1631 cm $^{-1}$ corresponding to C=N stretching (imine bonds) and disappearance of an absorption band at 1672 cm $^{-1}$ corresponding to C=O stretching (aldehyde), as well as the presence of bands in low wavenumber regions, corresponding to metal–carboxylate bonds of the MOP, indicating a successful condensation reaction and grafting of Zr-MOPs into the iCOF matrix (Fig. 2b).^{46–50} Also, the bands of N-H stretching at \sim 1439 and \sim 3423 cm $^{-1}$ confirm the presence of few free NH_2 functional groups in the composite material (Fig. S10†).

The XPS survey spectra of the composite clearly indicated the presence of all relevant elements, including C, N, O, Zr, and Cl, as compared to pristine iCOF, validating the grafting of Zr-MOP in the porous structure of the iCOF (Fig. 2c). Moreover, the deconvoluted XPS spectra of the Zr 3d energy level of IPcomp-8 were found to be shifted towards lower binding energy compared to Zr 3d of the pristine MOP, indicating interaction between the guest MOP and host iCOF matrix (Fig. 2d and S11†).51 The thermogravimetric analysis (TGA) of IPcomp-8 illustrated an improved thermal stability of the MOP molecules when anchored into the iCOF matrix with no disintegration of the structure up to 400 °C (Fig. S12†). The presence of featured signals of both MOP and iCOF in the ¹³C solid-state CP-MAS NMR spectrum of the composite material denoted that IPcomp-8 consisted of both MOP and iCOF structures (Fig. S13†). Thereafter, the nitrogen gas sorption isotherm was recorded at low temperature (77 K) to assess the porous nature of the nanocomposite in comparison to the pristine material. The isotherm profile of IPcomp-8 revealed relatively less uptake of N2 gas at low relative pressure, showing that the composite is less porous. Also, the sorption data exhibited significantly lower gas uptake and thus lower surface area of IPcomp-8 than the pristine iCOF, which imply the successful incorporation of Zr-MOP inside the iCOF matrix as well as exfoliation into ultrathin nanosheets (Fig. 2e).52 Furthermore, in order to evaluate the ionic properties of IPcomp-8, zeta potential measurement has been performed. The surface charge of IPcomp-8 was found to be positive with a zeta potential value of +49.3 \pm 2.1 mV, which inferred its high cationic nature (Fig. 2f). Importantly, the surface charge of the pristine cationic MOP and iCOF was found to be lower than that of the composite. The nanoscale level surface morphology of IPcomp-8 was examined by the FESEM, TEM and AFM analyses. The high-resolution FESEM images of IPcomp-8 showed robust sheet like morphology with plenty of small thin layers (Fig. 2g and S14†), while the pristine MOP and iCOF showed cube like microcrystalline and bulk-flake like morphology with micron to submicron layers, respectively (Fig. S14†). 48,51,52 Similar thin sheet like morphology was found in the case of TEM analysis (Fig. S15†). The fast Fourier transformation (FFT) images from the high resolution TEM (HRTEM) experiment showed a highly crystalline nature with clear lattice fringes and a selected area electron diffraction (SAED) pattern of the IPcomp-8 structure (Fig. 2h(insets)).53 Additionally, the energy dispersive X-ray (EDX) examination of IPcomp-8 revealed a Zr/Cl ratio of ~2.9 (in agreement with the crystallographic

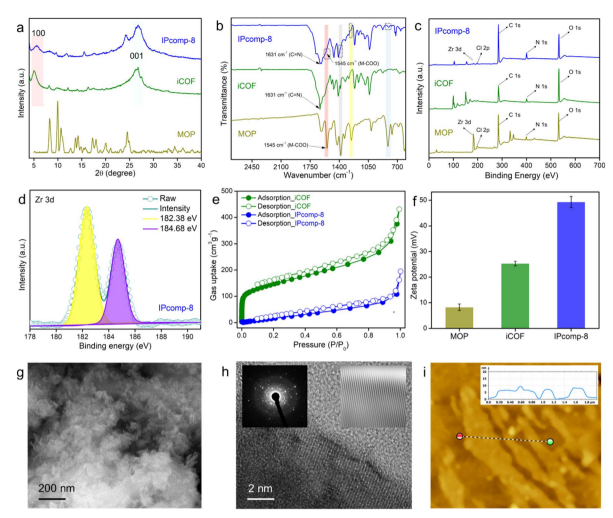


Fig. 2 (a) PXRD profile, (b) FTIR spectra, and (c) XPS survey spectra of IPcomp-8 along with the pristine materials, (d) deconvoluted Zr 3d XPS spectra of IPcomp-8, (e) N₂ sorption profile of iCOF and IPcomp-8 at 77 K, (f) comparison of zeta potential values of MOP, iCOF and IPcomp-8, (g) FESEM and (h) HRTEM images of IPcomp-8, (inset: lattice fringes and SAED images), and (i) AFM image and the corresponding height profile of IPcomp-8.

information of the Zr(IV)-MOP)47 confirming the existence of stable cationic MOP molecules in the iCOF matrix (Tables S1-S5†). The EDX and high-angle annular dark-field (HAADF) TEM elemental mapping of the composite materials from TEM experiments demonstrated the even distribution of the MOPs' constituent elements (Zr distribution) along with the other relevant elements (Fig. S16 and Tables S1-S5†). Moreover, the AFM experiment has been performed to explore the surface morphology of IPcomp-8 with greater details. The AFM images of IPcomp-8 demonstrated an ultrathin sheet like morphology with an average height of \sim 4 to 7 nm (Fig. 2i). However, the AFM morphology of the pristine iCOF exhibited bulk sheets with an average height of ~40 nm (Fig. S17†). These data clearly indicated the ultrathin morphology (nanosheets) of the hybrid composite. The formation of such an ultrathin structure of IPcomp-8 was mainly attributed to the exfoliation of the 2D layers of iCOF in the hybrid composite induced by intermolecular ionic repulsion between the adjacent iCOF layers, which is further originated due to the presence of charged (cationic)

MOP molecules inside the iCOF structure as well as the cationic backbone of the iCOF matrix (Fig. 1).52-55 Moreover, the optical characteristics of the hybrid composite also helped understand the presence of luminescent amino-functionalized MOP within the iCOF matrix. The hybrid composite's UV-vis diffuse reflectance spectroscopy (DRS) spectra revealed distinct absorption peaks for both the MOP and iCOF, demonstrating the presence of both components in the hybrid composite (Fig. S18†). Apart from these characterization studies, detailed control experiments have been performed to investigate the stability and a leaching test of the covalently bound (through the imine bond formation between free amine groups of MOPs and the aldehyde group of iCOF, Fig. 1) MOP molecules with iCOF was conducted, which indicated successful formation of the Zr-MOP/iCOF hybrid composite materials (IPcomp-8) (see ESI notes 1 and 2, Fig. S19-S24†). Furthermore, the hybrid composite's stability to irradiation was studied by analysing gamma ray irradiated samples using PXRD and FT-IR analysis (Fig. S25 and S26†).

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Capture studies

IPcomp-8 displayed a stable colloidal suspension in water for long time with the characteristic Tyndall effect owing to its intrinsic ionic nature (Fig. S27†).55 The presence of excess free exchangeable Cl anions and multiple interaction sites, including the Zr(IV) secondary building unit (SBU) and amino functional group of the MOP and cationic TAG moiety of the iCOF in the hybrid composite material enthused us to examine its sequestration capability toward toxic/hazardous oxoanions in water. $^{56-58}$ Metal derived radioactive oxoanion, TcO_4^- , and its surrogate oxoanion ReO₄ have been chosen for the sequestration study. Initially, to optimize the performance, we carried out the capture study with a series of compounds (composites synthesized with different amounts of MOPs) in ReO₄ solution (\sim 25 ppm) treated for 5 min. From the result of the inductively coupled plasma mass spectroscopy (ICP-MS) analysis it was found that the composite made by taking \sim 15 mg of MOP

(IPcomp-8) showed the best ReO₄ sequestration performance (Fig. S28†). Thereafter, all the following metal-oxoanion sequestration studies have been performed with IPcomp-8. Efficient extraction of radioactive metal pollutants from nuclear waste systems being a key-targeted criterion, at first, the effect of contact time of IPcomp-8 towards TcO₄ has been evaluated by batch adsorption experiments. The kinetics study of TcO₄ capture was conducted with the radiometry method where IPcomp-8 was found to exhibit ultrafast sequestration performance as within less than 5 min of contact time, almost >93% removal efficiency was observed (Fig. 3a). Notably, the fast sorption kinetics of IPcomp-8 is comparable to that of the other record holding state-of-the-art adsorbents and substantially faster than that of the majority of the reported sorbents. In this context, it is important to note that the majority of the widely used adsorbents for TcO₄⁻ sorption, including commercial anion-exchange resins, Purolite A532E and A530E,59 cationic

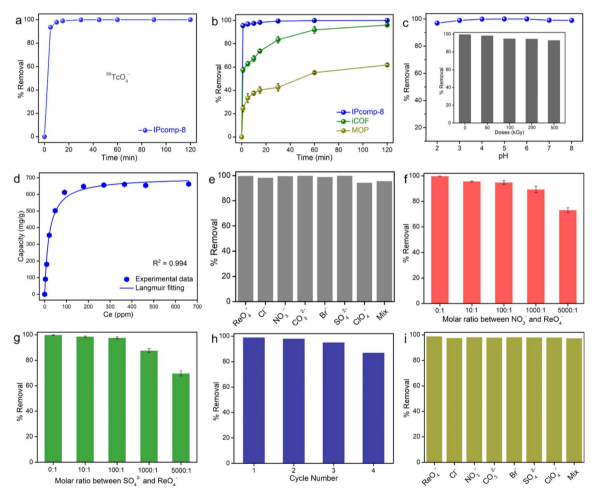


Fig. 3 % Removal of (a) TcO_4^- and (b) ReO_4^- anions in different contact times by IPcomp-8, MOP and iCOF, conditions: $[Re]_{initial} = 25$ ppm and $m_{sorbent}/V_{solution} = 1$ g L^{-1} . (c) Removal efficiency of IPcomp-8 towards TcO_4^- at different pH and (inset) different doses of γ -irradiation, and (d) Langmuir isotherm fitting model for ReO_4^- capture, conditions: contact time = 24 hours and $m_{sorbent}/V_{solution} = 1$ g L^{-1} . (e) Result of selectivity for ReO_4^- removal efficiency in the presence of different anions with high concentrations, conditions: $[Re]_{initial} = 25$ ppm, contact time = 24 hours, and $m_{sorbent}/V_{solution} = 1$ g L. (f and g) Selectivity result for ReO_4^- in the presence of excess NO_3^- and SO_4^{2-} anions, respectively, and (h) recycle test result of IPcomp-8 for ReO_4^- ions, conditions: $[Re]_{initial} = 25$ ppm, contact time = 24 hours, and $m_{sorbent}/V_{solution} = 1$ g L^{-1} . (i) Selectivity of IPcomp-8 towards ReO_4^- capture at an ultralow concentration, conditions: $[Re]_{initial} = 1000$ ppb, contact time = 24 hours, and $m_{sorbent}/V_{solution} = 1$ g L^{-1} .

MOFs,60-62 inorganic materials,11,63 etc., were found to take a comparatively longer time to reach sorption equilibrium. This ultrafast exchange kinetic result of IPcomp-8 can help quickly respond to a nuclear incident and lower the risk posed by a ⁹⁹TcO₄ leakage, which further encouraged us to carry out detailed TcO₄ - sequestration studies. Now, owing to the high radioactive nature and low availability of 99TcO₄-, the following high concentration batch anion-exchange experiments have been performed with nonradioactive surrogate ReO₄ owing to their identical chemical behaviour. For this, the timedependent sequestration study was performed by taking IPcomp-8 in 25 ppm of stock ReO₄ - solution. After a certain contact time interval, the compound was filtered and the residual concentration was measured. The result from ICP-MS data indicated the significant lowering of the Re(vII) concentration in very little time, as almost 95% ReO₄ removal efficiency was observed within less than 1 min in the case of IPcomp-8 (Fig. 3b, and S29†). Furthermore, the sorption kinetics towards ReO₄⁻ was found to follow the pseudosecond order kinetics model with a high correlation coefficient of R^2 0.999 for IPcomp-8. The rate constant value of the hybrid showed an ultrafast sorption rate than pristine materials (Fig. S28†). Inspired by this result as well as in order to evaluate the advancement of the hybrid composite (IPcomp-8) over the individual components towards improved TcO₄ sequestration, similar uptake study was conducted with the pristine MOP and iCOF. To our delight, in this comparison study, the composite material was found to demonstrate higher removal efficiency than both the bare MOP and iCOF (Fig. 3b, and S29†). Within less than 1 min of contact time, IPcomp-8 exhibited ~95% removal efficiency, whereas MOP and iCOF exhibited significantly lower uptake efficiencies, at \sim 25% and \sim 57%, respectively. This outcome demonstrates the hybrid composite's superiority over distinct substances in metal oxoanion separation applications (Fig. S30-S32†).

The pH of the nuclear wastewater or industrial wastewater system may vary from acidic to alkaline depending upon the environment of the discharge system. Therefore, it is important to check the sequestration efficiency of a sorbent material in a wide pH range. Interestingly, IPcomp-8 was found to eliminate TcO₄ across a wide range of pH without losing its actual efficiency (Fig. 3c). In addition, considering the strong ionizing field in nuclear waste remediation, the sequestration of TcO₄ by IPcomp-8 was evaluated after irradiation with high doses of γ-irradiation. To our delight, the hybrid exhibited excellent elimination of TeO_4^- even after exposing γ -irradiation, further indicating high radiation tolerance ability (Fig. 3c(inset)). Next, on account of the enormous inventory of 99TcO4- in used nuclear waste, the adsorption capacity of an adsorbent is a fundamental parameter in order to understand its full efficiency. The saturation capacity of IPcomp-8 for ReO₄ was estimated by carrying out the adsorption isotherm experiment. In a typical test, a range of initial concentrations (20–800 ppm) of ReO4- aqueous solution was treated with IPcomp-8 for a contact period of 24 h to reach sorption equilibrium. The uptake of ReO₄⁻ by IPcomp-8 was found to follow the Langmuir isotherm model with the high correlation coefficient value,

which indicated the monolayer sorption mechanism (Fig. 3d). When the sorption capacities were plotted as a function of equilibrium concentration, the maximum sorption capacity of ReO_4^- by IPcomp-8 was found to be \sim 632 mg g⁻¹. However, a reduction in the capacity was noticed when measuring the maximum adsorption capacity of the IPcomp-8 sample after γ ray irradiation towards high concentrations of ReO₄ (Table S6†). It should be mentioned that apart from the kinetics and capacity, selectivity of an adsorbent is another highly important parameter to realize its potential towards real-time application. Therefore, the efficacy of an adsorbent towards a specific analyte in the presence of an excess of other interfering analytes should be tested. According to earlier literature, it has been observed that the majority of studies have mainly focused on uptake capacity, but very little attention has been paid to the selectivity aspect in the presence of other interfering analyte anions, which is crucial for sequestration from the actual wastewater system. 19,62-65 Nuclear or industrial wastewater contains anions such as Cl⁻, NO₃⁻, Br⁻, SO₄²⁻, ClO₄⁻, etc., in enormous excess amounts, which interfere in the adsorption process.⁵⁷⁻⁶² Initially, to check the influence of these aforementioned interfering anions in this adsorption process, a similar ReO₄ capture experiment has been performed by taking a binary mixture of ReO₄⁻ as the target anion along with these anions individually in a similar higher concentration. To our delight, it was found that the ReO₄ sorption efficiency by IPcomp-8 remained intact (almost 98%) in the presence of other competing anions (Fig. 3e). Inspired by this result, we were motivated further to test the efficiency of IPcomp-8 towards ReO_4^- in the presence of surplus concentrations (\sim 10, \sim 100, \sim 1000, and \sim 5000-fold) of two specific interfering analytes such as NO₃⁻ and SO₄²⁻, which are generally omnipresent with TcO₄ or ReO₄ in wastewater. By performing a similar sorption test, we observed no significant effect of both NO₃⁻ and SO₄²⁻ competing ions on the sequestration efficiency of IPcomp-8 up to a ~10-fold concentration (Fig. 3f and g). Even though the concentration of NO_3^-/SO_4^{-2-} is in a \sim 100-fold excess, IPcomp-8 exhibited as high as >95% removal percentage towards ReO₄. In the case of excessive concentrations (\sim 1000 and \sim 5000-fold), the ReO₄ removal efficiency of IPcomp-8 was found to diminish (Fig. 3f and g). However, IPcomp-8 was still found to exhibit \sim 89% and \sim 74% ReO₄ removal efficiency in the presence of \sim 1000 and \sim 5000-fold NO₃⁻, respectively (Fig. 3f). In addition to this, the composite also demonstrated superior removal efficiency towards ReO₄ in the presence of SO₄²⁻ anions, which is highly competitive owing to its high charge/ size ratio (Fig. 3g). The distribution coefficient value (K_d) of an adsorbent is an important parameter, which is used to explore its binding affinity towards a specific analyte.66 The K_d values of IPcomp-8 for ReO₄ in the presence of these aforementioned competing anions were further calculated, which exhibited as high as in 10⁶ order for all cases (Table S7†). All these results validate the highly selective nature of IPcomp-8 towards TcO₄ radioactive anions, making it a suitable adsorbent with potential for practical application. Such excellent selectivity of IPcomp-8 for TcO₄⁻ or ReO₄⁻ originates from the multifunctional properties of the composite, such as Zr(IV)-SBU, free NH₂-

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groups and TAG moieties. Next, to be an ideal adsorbent, recyclability is another imperative requirement. The hybrid material loaded with ReO₄⁻ anions was regenerated using saturated Na₂SO₄ solution by stirring it for ~10 hours. Even after four cycles, it was discovered that IPcomp-8 could be easily recycled without significantly losing its sorption effectiveness, demonstrating exceptional regeneration capability (Fig. 3h).

Overall, the IPcomp-8 hybrid material is superior to conventional adsorbents due to its rapid capture efficiency with high adsorption capacity and unparalleled selectivity toward TcO₄ or ReO₄. Due to accidental discharge in a real water body the concentration of the toxic analyte exists in a very low concentration (<1 ppm).¹ Therefore, it is important to explore the ability of the adsorbent to sequestrate the target analyte at trace concentrations. By virtue of the rapid and efficient TcO₄ or ReO₄ oxoanion adsorbing behaviour of IPcomp-8, treatment of ReO₄ contaminated wastewater was performed, by mimicking the real time scenario. Moreover, an ultralowconcentration (~1000 ppb) ReO₄ sequestration study in the presence of an abundance of other competing ions was also carried out and monitored using ICP-MS analysis to show the material's effectiveness. Initially, in the kinetics study, IPcomp-8 was found to exhibit ultrafast sequestration efficiency towards ${
m ReO_4}^-$ achieving ${\sim}99\%$ removal efficiency within less than 1 min (Fig. S33†). Consequently, the concentration of ReO₄ was found to decrease rapidly, and the final concentration was found to be much below the WHO permitted level of drinking water (Fig. S34†).67 Furthermore, the kinetic study was found to follow pseudo-second order kinetics with a high rate constant value (Fig. S35†). Moreover, the selectivity of IPcomp-8 in low concentrations was also investigated taking all the aforementioned coexisting anions. From the selectivity test, it was found that IPcomp-8 exhibits superior ReO₄ capture efficiency even in the presence of all other anions in equal as well as highly excess concentrations (Fig. 3i, S36 and S37†). All these results of efficient and selective entrapment of radioactive TcO₄ oxoanions indicated the potential of IPcomp-8 towards real-time nuclear wastewater treatment application.

Adsorption mechanism studies

After evaluating the capture performance, the underlying sorption mechanistic insight was further explored experimentally and theoretically in order to understand the multiple interactions that are responsible for such efficient TcO4sequestration performance by IPcomp-8. For this, the following post capture characterization studies have been performed. At first the FTIR analysis of ReO₄⁻ loaded IPcomp-8 displayed the appearance of a sharp peak at ∼911 cm⁻¹ corresponding to Re-O asymmetric stretching of ReO₄⁻ anions, which supports the binding of ReO₄ with IPcomp-8 (Fig. 4a).⁵⁷⁻⁶² The thermogravimetric analysis of the ReO₄ loaded material showed similar thermal stability compared to the pristine IPcomp-8 (Fig. S38†). Thereafter, the XPS analysis was performed to understand the nature of interaction between the positively charged backbone of the composite material and the ReO₄⁻ oxoanion. The Re 4f binding energy peak was clearly visible in the XPS survey spectra

of samples that had undergone ReO₄ treatment, indicating that Re was present on the surface of IPcomp-8 (Fig. 4b). The deconvoluted XPS spectra of the Re 4f core-level confirmed the existence of Re in their +7 oxidation state (ReO₄⁻) (Fig. 4c).⁵⁷⁻⁶² Furthermore, extended X-ray adsorption fine structure (EXAFS) study was performed in order to understand the local structure around the Re atoms in ReO₄ loaded IPcomp-8. The fitting analysis of the raw data and Fourier transforms of Re L3 edge K³-weighted spectra of ReO₄ loaded IPcomp-8 indicated a Re-O shell with a radial distance (R) of 1.734 Å and the coordination number of Re-O was found to be 4 (Fig. 4d and e). These values were found to be close to the matric parameters of the referenced material (Table S8 and Fig. S39†).68 A further detailed EXAFS analysis of the Zr metal center of the ReO₄⁻ treated IPcomp-8 supported the stable existence of the MOP within the iCOF matrix of the composite after the capture test (ESI note 3, Table S9 and Fig. S40-S42†). Importantly, it was discovered that the EXAFS results and the XPS data correlated, confirming that Re species maintain their +7 oxidation states and that the adsorption process does not include Re(vII) reduction. 69 This finding demonstrated unequivocally that the strong electrostatic interactions between the cationic structure of IPcomp-8 and ReO₄ were responsible for the excellent removal efficiency.

Next, we have also performed an isothermal titration calorimetry (ITC) experiment to evaluate thermodynamic feasibility of the favourable interactions between the compound and the analyte in an aqueous medium (ESI note 4†). The ITC experiment quantitatively provides the change in enthalpy, entropy and Gibbs free energy associated with the adsorption driven by the ion-exchange process between the adsorbent and adsorbate.70 The recorded thermogram depicted a high association constant with an exothermic binding event along with the parameter. This result indicated that the adsorption of ReO₄⁻ by IPcomp-8 was favourable as the Gibbs free energy of the process was found to be negative with a positive association constant and positive entropy (Fig. 4f and S43†). Thus, the ITC data clearly demonstrated the similarities between the thermodynamic parameters and the experimental findings of strong interaction of IPcomp-8 and ReO₄-. Furthermore, the HRTEM image of ReO₄⁻ treated IPcomp-8 showed identical morphological configuration with retained crystalline nature (Fig. S44†). This further explored the robust nature of the hybrid composite material towards wastewater treatment. The EDX elemental mapping analysis from the TEM experiment of ReO₄ aIPcomp-8 was performed, which indicated the appearance of homogeneous distribution of Re metal ions along with other relevant elements throughout the structure of the composite (Fig. 4g, h, and S45†). All these above investigations and characterization studies were carried out to support the possible ReO₄ or TcO₄ binding mechanism with IPcomp-8. Previously in the literature it was well established that Zr(IV) SBU and amino functional groups can act as active sites for selective oxoanion binding.46,71 As a result, the composite material's cationic guest (NH2-MOPs) has a significant impact on the selective trapping of TcO₄. Having said that, pristine MOP molecules undergo self-aggregation,41 which inhibits the functional groups from interacting with the incoming

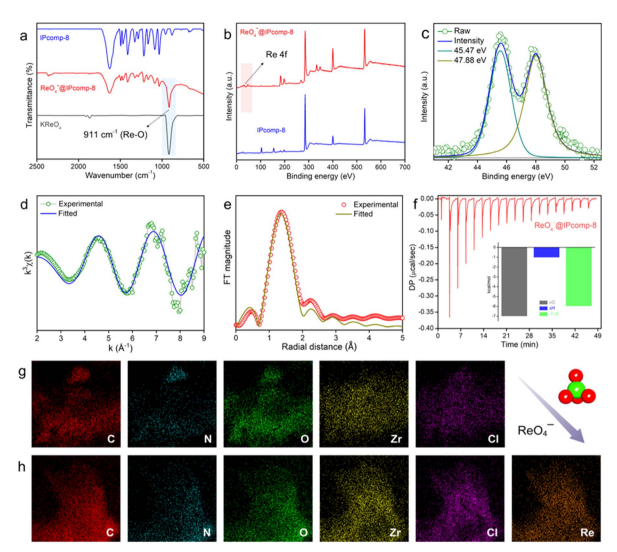


Fig. 4 (a) FT-IR spectra of ReO₄⁻ loaded IPcomp-8, (b) XPS survey spectra and (c) deconvoluted Re 4f core-level XPS spectra of IPcomp-8 and ReO₄-@IPcomp-8, (d) raw and (e) Fourier transforms of Re L3 edge EXAFS spectra of ReO₄- treated IPcomp-8, (f) ITC thermogram with its corresponding parameters, and (g and h) TEM EDX elemental mapping of IPcomp-8 and ReO₄⁻@IPcomp-8, respectively.

oxoanions; hence this results in the reduction of the capture efficiency. In the IPcomp-8 hybrid composite material nanosized MOPs are covalently tethered with the layers of the iCOF that restricts the self-aggregation process and consequently exposed more number of active binding sites for selective interaction with metal oxoanions. Moreover, when the MOP molecules bind to the iCOF it is exfoliated driven by electrostatic repulsions and forms ultrathin hybrid nanosheets. These nanosheets bear a greater number of exposed interactive sites with excessive free exchangeable Cl ions for the amplified strong electrostatic interactions between the cationic host matrix and anionic metal based oxoanions (ReO₄ or TcO₄) which enables ultrafast and selective capture efficiency. Additionally, the TAG functional moiety of the selected chemically stable iCOF network synergistically helped with the selective interaction with targeted metal oxo pollutants. IPcomp-8 exhibited highly efficient selective removal of ReO₄⁻ oxoanions, from the contaminated water system. The multifunctional nature of the hybrid ultrathin nanosheet composite material

can be rationalized to the efficient sequestration performance. Among them, the ultrathin nanosheet structure of the composite aids in the rapid diffusion of oxoanions through its exfoliated open porous structure towards the main active sites: guest-amino functionalized Zr(w)-SBU-based cationic MOPs (Fig. 5a). Thereafter, the oxoanion TcO_4^-/ReO_4^- interacts with the Zr-SBU and the free -NH2 group of the cationic MOP, followed by exchange with Cl ions of both the guest (MOP) and host matrix (iCOF). Such interactions can be attributed to the dispersive interactions or supramolecular interactions or dipole interactions of the TcO₄⁻/ReO₄ with the functional groups of MOP molecules as well as the backbone of the iCOF matrix of the composite material. These selective strong interactions were further validated using theoretical calculations using a density functional theory (DFT) study, which quantitatively supported the experimental values. The interactions between the cationic MOP and fragment of iCOF (both abbreviated as U+) in IPcomp-8 and anions such as TcO₄ or ReO₄, SO₄, NO₃, and others were investigated. The calculations showed that the Zr(iv)-SBU

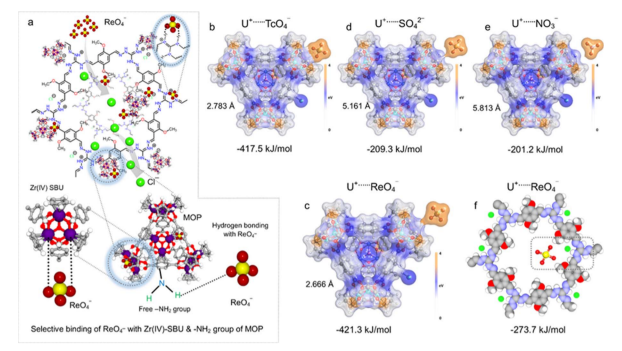


Fig. 5 (a) Illustration of mechanistic insight of selective ReO_4^- capture through different interactions by IPcomp-8. Density functional theory (DFT) calculations: (b-f) energy-optimized adsorption complexes for interactions between TcO_4^- or ReO_4^- with Zr(w)-SBU of the MOP and iCOF structural units with their corresponding binding energies.

of the MOPs selectively interacts with TcO_4^- and ReO_4^- over other anions with their stable adsorption structure at binding energies of -417.5 and -421.3 kJ mol^{-1} , respectively (Fig. 5b and c). The binding energies for other anions were less than those of TcO_4^- and ReO_4^- (Fig. 5d, e and S46†). Other bindings through a hydroxyl group (OH $^-$), hydrogen of the $^-$ Cp ring and the amino functional group ($^-$ NH $_2$) of the MOP are provided in the ESI file (Fig. S47 and S48).† In addition, the cationic backbone of the iCOF structure interacts with ReO_4^- anions through hydrogen bonding with the TAG moiety at a binding energy of $^-$ 273.7 kJ $^-$ kJ $^-$ (Fig. 5f). Such a difference between the binding energies of $^-$ 0 through $^-$ 1 or $^-$ 1 or $^-$ 2 with $^-$ 2 with $^-$ 3 over other anions (such as $^-$ 3 NO $_3^-$ 3, SO $_4^{-}$ 4, Cl $_3^-$ 3 RF $_3^-$ 4 etc.) through a robust ion-exchange mechanism.

Membrane based continuous flow-through sequestration

Inspired by such excellent radioanion sequestration efficiency as well as the robust ultrathin 2D nanosheet nature of the hybrid COF composite material (IPcomp-8), we further sought to explore the potential of the material for large-scale continuous flow-through based TcO_4^- or ReO_4^- extraction by developing a hybrid membrane. At first an IPcomp-8 based stable hybrid membrane was fabricated by a typical mixed-matrix-membrane technique (Fig. 6a) (see Section S9.1†).⁵³ The thus developed hybrid membrane was found to possess a smooth surface and compact packing without significant defects on the surface, as revealed from the SEM images of the membrane (Fig. 6b and S49†). In addition, the 2D cross-sectional images from the microscale X-ray computed tomography (CT)

experiment indicated a similar smooth surface morphology of the membrane (Fig. S50†). Moreover, the volume rendered 3D CT images and its corresponding color-coded images of the membrane showed distribution of pores throughout the undulated surfaces on one side and relatively smoother ones on the other side (Fig. 6c–e and S51†). Task-specific applications such as sequestration of specific elements/molecules/ions through membranes hold great promise towards industrial application.⁷² Therefore, the extraction performance of the IPcomp-8 membrane for ReO₄⁻ removal from contaminated samples was first evaluated (see Section S9.2†).

In a typical experiment, a ~ 1000 ppb ReO_4^- feed was charged through the hybrid membrane in a dynamic mode and the concentration of the membrane passed solution was measured by ICP-MS (Fig. S52†). As shown in Fig. 6f, after the treatment by a flow-through dynamic process the membrane was found to reduce the concentration of the ReO₄⁻ solution to below ~ 10 ppb. As a result of the adsorption-based membrane separation, the IPcomp-8 membrane was found to exhibit more than ~93% removal efficiency up to 170 mL of ReO₄ feed solution (Fig. 6g). After that, the removal efficiency started to decrease and was observed to be intact as >70% up to 315 mL of contaminated solution. However, after that, the removal efficiency of the membrane was significantly decreased because of blocking of most of the active sites through binding with ReO₄. Moreover, the recycling ability of the membrane was further investigated using saturated Na₂SO₄ solution, which greatly regenerated the capture performance of the hybrid membrane. In the recycling test, the membrane was found to demonstrate more than ~75% removal efficiency up to 250 mL of feed ReO₄

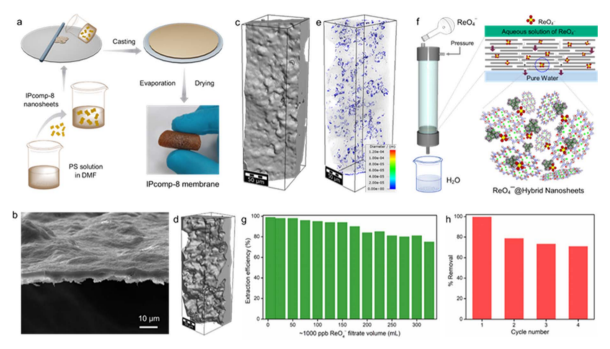


Fig. 6 (a) Fabrication procedure and digital image of the IPcomp-8 based hybrid membrane. (b) SEM image, (c and d) volume rendered 3D X-ray tomography images of the IPcomp-8 based membrane. (e) Visualization of pore distribution with a colour-scale CT image. (f) Schematic representation of flow-through selective extraction of ReO_4^- oxoanions by the IPcomp-8 based hybrid membrane. (g) Extraction efficiency of IPcomp-8 towards ReO₄ with different amounts of feed volume. (h) Regeneration result of the membrane, conditions: [Re]_{initial} = 1000 ppb, pH = 6.9.

solution in all four cycles (Fig. 6h). From the perspective of practical application, this study demonstrated the exceptional segregation effectiveness of the IPcomp-8 membrane towards TcO₄ oxoanions in both environmental and industrially contaminated water. The cooperative impact of the exposed ultrathin hybrid nanosheets, in addition to the presence of Zr-SBU, free -NH₂ groups, and exchangeable Cl⁻ ions in the overall nanocomposite, is thought to be responsible for the IPcomp-8 membrane's excellent selective separation performance.

Conclusions

In conclusion, we have strategically designed and successfully synthesized a covalently linked MOP@iCOF cationic hybrid composite material (IPcomp-8), which demonstrated ultrafast and selective sequestration of TcO₄ oxoanions in water. This work demonstrates an optimization of a rationally designed strategy to remarkably enhance the sequestration kinetics and selectivity towards pertechnetate segregation from wastewater systems. Structural and optical characterization studies have been thoroughly carried out to gain detailed insights into the novel composite material. The major targeted criterion being the selective entrapment of TcO₄⁻, careful anchoring of nanosized discrete cationic MOP molecules based on a Cp₃Zr₃-O(OH)₃ SBU into the chemically stable charged iCOF matrix produced unprecedented selectivity and ultrafast kinetics for TcO₄⁻/ReO₄⁻ capture. Notably, in the dynamic sorption technique, the membrane form of the hybrid composite illustrated

unparalleled absorption ability toward TcO₄⁻/ReO₄⁻ from wastewater samples. This newly developed hybrid cationic material may be a possible contender for wastewater treatment due to its rapid uptake capability, excellent selectivity, and satisfactory recyclability. We believe that the current discoveries will pave the way for the creation of a range of efficient innovative sorbent materials for use in other environmentally important separation applications.

Data availability

All data used for this study are available in the paper and ESI files.†

Author contributions

W. M. conceived and designed the project. W. M., S. F. and D. M. conducted all the experiments and analyzed all the data. W. M. wrote the manuscript, while S. F., S. L., D. M. helped finalize the draft. S. F. designed the graphical art in the figures. A. S. performed all the radioactive studies. R. R. U. did the EXAFS measurement and fitting analysis. A. T. performed the tomography experiment. M. M. S. performed the theoretical studies. S. K. G. supervised the project. All authors have given approval to the final version of the manuscript.

Conflicts of interest

The authors declare no competing interests.

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Notes and references

- 1 D. Banerjee, D. Kim, M. J. Schweiger, A. J. Kruger and P. K. Thallapally, *Chem. Soc. Rev.*, 2016, **45**, 2724–2739.
- 2 J. G. Darab and P. A. Smith, *Chem. Mater.*, 1996, **8**, 1004–1021.
- 3 International Atomic Energy Agency, *Nuclear Power Reactors* in the World, Reference Data Series No. 2, IAEA, Vienna, 2018.
- 4 K. H. Lieser, Radiochim. Acta, 1993, 63, 5-8.
- 5 B. Gu and K. E. Dowlen, An Investigation of Groundwater Organics, Soil Minerals, and Activated Carbon on the Complexation, Adsorption, and Separation of Technetium-99, Publication No. 4502, Environmental Sciences Division, ORNL/TM-13154, Oak Ridge, Tennessee, 1996.
- 6 Q. Sun, B. Aguila and S. Ma, *Trends Chem.*, 2019, 1, 292–303.7 S. R. J. Oliver, *Chem. Soc. Rev.*, 2009, 38, 1868–1881.
- 8 A. Ravi, A. S. Oshchepkov, K. E. German, G. A. Kirakosyan, A. V. Safonov, V. N. Khrustalev and E. A. Kataev, *Chem. Commun.*, 2018, 54, 4826–4829.
- 9 J. Li, L. Zhu, C. Xu, D. Zhang, M. A. Silver, P. Li, L. Chen, Y. Li, D. Zuo, H. Zhang, C. Xiao, J. Chen, J. Diwu, O. K. Farha, T. E. Albrecht-Schmitt, Z. Chai and S. Wang, *J. Am. Chem. Soc.*, 2018, 9, 93007.
- 10 Y. Wang, J. Lan, X. Yang, S. Zhong, L. Yuan, J. Li, J. Peng, Z. Chai, J. K. Gibson, M. Zhai and W. Shi, *Adv. Funct. Mater.*, 2022, 32, 2205222.
- 11 S. Wang, P. Yu, A. B. Purse, M. J. Orta, J. Diew, W. H. Casey, B. L. Phillips, E. V. Alekseev, W. Depmeier, D. T. Hobbs and T. E. Albrecht-Schmitt, *Adv. Funct. Mater.*, 2012, 22, 2241–2250.
- 12 K. Kang, S. Lie, M. Zhang, L. Li, C. Liu, L. Lei, X. Dai, C. Xu and C. Xiao, *Adv. Funct. Mater.*, 2022, **32**, 2208148.
- 13 C. Xiao, A. Khayambashi and S. Wang, *Chem. Mater.*, 2019, **31**, 3863–3877.
- 14 X. Wang, L. Chen, L. Wang, Q. Fan, D. Pan, J. Li, F. Chi, Y. Xie, S. Yu, C. Xiao, F. Luo, J. Wang, X. Wang, C. Chen, W. Wu, W. Shi, S. Wang and X. Wang, *Sci. China: Chem.*, 2019, 62, 933–967.

- 15 M. Hao, Y. Liu, W. Wu, S. Wang, X. Yang, Z. Chen, Z. Tang, Q. Huang, S. Wang, H. Yang and X. Wang, *EnergyChem*, 2023, 5, 100101.
- 16 A. Celik, S. C. Roy, M. A. Quintero, K. Taylor-Pashow, D. Li, M. G. Kanatzidis, X. Zhu and S. M. Islam, ACS Appl. Eng. Mater., 2023, 1(7), 1711–1718.
- 17 D. Li, N. B. Shustova, C. R. Martin, K. Taylor-Pashow, J. C. Seaman, D. I. Kaplan, J. W. Amoroso and R. Chernikov, J. Environ. Radioact., 2020, 222, 106372.
- 18 Y. Wang, M. Xie, J. Lan, L. Yuan, J. Yu, J. Li, J. Peng, Z. Chai, J. K. Gibson, M. Zhai and W. Shi, *Chem*, 2020, **10**, 2796–2809.
- 19 K. M. Long, G. S. Goff, S. D. Ware, G. D. Jarvinen and W. H. Runde, *Ind. Eng. Chem. Res.*, 2012, **51**, 10445–10450.
- 20 D. Sheng, L. Zhu, C. Xu, C. Xiao, Y. Wang, Y. Wang, L. Chen, J. Diwu, J. Chen, Z. Chai, T. E. Albrecht-Schmitt and S. Wang, *Environ. Sci. Technol.*, 2017, 51, 3471–3479.
- 21 S. Dong, Z. Yu, L. Guo, Y. Yang, C. Tu, R. Krishna and F. Luo, *Small*, 2023, 2304054.
- 22 L. Zhu, D. Sheng, C. Xu, X. Dai, M. A. Silver, P. Li, Y. Wang, Y. Wang, L. Chen, C. Xiao, J. Chen, R. Zhou, C. Zhang, O. K. Farha, Z. Chai, T. E. Albrecht-Schmitt and S. Wang, J. Am. Chem. Soc., 2017, 139(42), 14873–14876.
- 23 Q. Sun, L. Zhu, B. Aguila, P. K. Thallapally, C. Xu, J. Chen, S. Wang, D. Rogers and S. Ma, *Nat. Commun.*, 2019, 10, 1646.
- 24 P. Zhang, Z. Wang, S. Wang, J. Wang, J. Liu, T. Wang, Y. Chen, P. Cheng and Z. Zhang, *Angew. Chem.*, 2022, 61, e202213247.
- 25 B. Wang, J. Li, H. Huang, B. Liang, Y. Zhang, L. Chen, K. Tan, Z. Chai, S. Wang, J. T. Wright, R. W. Meulenberg and S. Ma, *ACS Cent. Sci.*, 2024, 10(2), 426–438.
- 26 Y. Wang, M. Xie, J. Lan, L. Yuan, J. Yu, J. Li, J. Peng, Z. Chai, J. K. Gibson, M. Zhai and W. Shi, *Chem*, 2020, 6, 1–14.
- 27 H. Peng, B. Jiang, F. Li, J. Gong, Y. Zhang, M. Yang, N. Liu and L. Ma, Chem. Mater., 2023, 35(6), 2531–2540.
- 28 C. R. Kim, T. Uemera and S. Kitagawa, *Chem. Soc. Rev.*, 2016, 45, 3828–3845.
- 29 C.-P. Li, H.-R. Li, J.-Y. Ai, J. Chen and M. Du, *ACS Cent. Sci.*, 2020, **6**, 2354–2361.
- 30 G. Distefano, H. Suzuki, M. Tsujimoto, S. Isoda, S. Bracco, A. comotti, P. Sozzani, T. Uemura and S. Kitagawa, *Nat. Chem.*, 2013, 5, 335–341.
- 31 R. K. Deshpande, J. L. Minnaar and S. G. Telfer, *Angew. Chem., Int. Ed.*, 2010, **49**, 4598–4602.
- 32 Y. Peng, M. Zhao, B. Chen, Z. Zhang, Y. Uang, F. Dai, Z. Lai, X. Cui, X. Tan, C. Tan and H. Zhang, *Adv. Mater.*, 2018, 30, 1705454.
- 33 F.-M. Zhang, J.-L. Sheng, Z.-D. Yang, X.-J. Sun, H.-L. Tang, M. Lu, H. Dong, F.-C. Shen, J. Liu and Y.-Q. Lan, *Angew. Chem.*, 2018, 130, 12282–12286.
- 34 T. Kitao, Y. Zhang, S. Kitagawa, B. Wang and T. Uemura, *Chem. Soc. Rev.*, 2017, **46**, 3108–3133.
- 35 M. Muschi and C. Serre, *Coord. Chem. Rev.*, 2019, **387**, 262–
- 36 S. Mollick, S. Fajal, S. Saurabh, D. Mahato and S. K. Ghosh, *ACS Cent. Sci.*, 2020, **6**, 1534–1541.
- 37 Y. Liang, X. Yang, X. Wang, Z.-J. Guan, H. Xing and Y. Fang, *Nat. Commun.*, 2023, **14**, 5223.

Edge Article

- 38 C. S. Diercks and O. Yaghi, Science, 2017, 355, eaal1585.
- 39 A. J. Gosselin, C. A. Rowland and E. D. Bloch, *Chem. Rev.*, 2020, **120**, 8987–9014.
- 40 F. J. Rizzuto, K. S. Krbek and J. R. Nitschke, *Nat. Rev. Chem*, 2019, 3, 204–222.
- 41 S. Mollick, S. Fajal, S. Mukherjee and S. K. Ghosh, *Chem. Asian J.*, 2019, **14**, 3096–3108.
- 42 X. Qiu, W. Zhong, C. Bai and Y. Li, *J. Am. Chem. Soc.*, 2016, 138, 1138–1141.
- 43 Y.-H. Kang, X. D. Liu, N. Yan, Y. Jiang, X.-Q. Liu, L.-B. Sun and J.-R. Li, *J. Am. Chem. Soc.*, 2016, **138**, 6099–6102.
- 44 A. J. Howarth, M. J. Katz, T. C. Wang, A. E. Prats-Platero, K. W. Chapman, J. T. Hupp and O. K. Farha, *J. Am. Chem. Soc.*, 2015, 137, 7488–7494.
- 45 C. O. Audu, H. G. Nguyen, C.-Y. Chang, M. J. Katz, L. Mao, O. K. Farha, J. T. Hupp and S. T. Nguyen, *Chem. Sci.*, 2016, 7, 6492–6498.
- 46 R. Lv, J. Wang, Y. Zhang, H. Li, L. Yang, S. Liao, W. Gu and X. Liu, J. Mater. Chem. A, 2016, 4, 15494–15500.
- 47 G. Liu, Y. D. Yuan, J. Wang, Y. Cheng, S. B. Peh, Y. Wang, Y. Qian, J. Dong, D. Yuan and D. Zhao, *J. Am. Chem. Soc.*, 2018, 140, 6231–6234.
- 48 Z. Zhang, X. Dong, J. Yin, Z.-G. Li, X. Li, D. Zhang, T. Pan, Q. Lei, X. Liu, Y. Xie, F. Shui, J. Li, M. Yi, J. Yuan, Z. You, L. Zhang, J. Chang, H. Zhang, W. Li, Q. Fang, B. Li, X.-H. Bu and Y. Han, J. Am. Chem. Soc., 2022, 144, 6821–6829.
- 49 S. Chandra, S. Kandambeth, B. P. Biswal, B. Lukose, S. M. Kunjir, M. Chaudhary, R. Babarao, T. Heine and R. Banerjee, J. Am. Chem. Soc., 2013, 135, 17853–17861.
- 50 J. Liu, W. Duan, J. Song, X. Guo, Z. Wang, X. Shi, J. Liang, J. Wang, P. Cheng, Y. Chen, M. J. Zaworotko and Z. Zhang, J. Am. Chem. Soc., 2019, 141, 12064–12070.
- 51 S. Fajal, W. Mandal, S. Mollick, Y. D. More, A. Torris, S. Saurabh, M. Shirolkar and S. K. Ghosh, *Angew. Chem.*, *Int. Ed.*, 2022, 61, e202203385.
- 52 H. Singh, M. Devi, N. Jena, M. Iqbal, Y. Nailwal, A. D. Sarkar and S. K. Pal, *ACS Appl. Mater. Interfaces*, 2020, **12**, 13248–13255.
- 53 H. Yang, L. Yang, H. Wang, Z. Xu, Y. Zhao, Y. Luo, N. Nasir, Y. Song, H. Wu, F. Pan and Z. Jiang, *Nat. Commun.*, 2019, **10**, 2101
- 54 M. A. Khayum, S. Kandambeth, S. Mitra, S. B. Nair, A. Das, S. S. Nagane, R. Mukherjee and R. Banerjee, *Angew. Chem.*, *Int. Ed.*, 2016, 128, 15833–15837.
- 55 S. Mitra, S. Kadambeth, B. P. Biswal, A. K. Mohammed, C. K. Choudhury, M. Mehta, G. Kaur, S. Banerjee, A. Prabhune, S. Verma, S. Roy, U. K. Kharul and R. Banerjee, J. Am. Chem. Soc., 2016, 138, 2823–2828.

- 56 S. Rapti, A. Pournara, D. Sarma, I. T. Papadas, G. S. Armatas, A. C. Tsipis, T. lazarides, M. G. Kanatzidis and M. J. Manos, *Chem. Sci.*, 2016, 7, 2427–2436.
- 57 J. Li, B. Li, N. Shen, L. Chen, Q. Guo, L. Chen, L. He, X. Dai, Z. Chai and S. Wang, ACS Cent. Sci., 2021, 7, 1441–1450.
- 58 X. Li, Y. Li, H. Wang, Z. Niu, Y. He, L. Jin, M. Wu, H. Wang, L. Chai, A. M. Al-Enizi, A. Nafady, S. F. Shaikh and S. Ma, Small, 2021, 17, 2007994.
- 59 J. Li, L. Zhu, C. Xiao, L. Chen, Z. Chai and S. Wang, *Radiochim. Acta*, 2018, **106**, 581–591.
- 60 N. Shen, Z. Yang, S. Liu, X. Dai, C. Xiao, K. Taylor-Pashow, D. Li, C. Yang, J. Li, Y. Zhang, M. Zhang, R. Zhou, Z. Chai and S. Wang, *Nat. Commun.*, 2020, 11, 5571.
- 61 D. Sheng, L. Zhu, X. Dai, C. Xu, P. Li, C. Pearce, C. Xiao, J. Chen, R. Zhou, T. Duan, O. K. Farha, Z. Chai and S. Wang, *Angew. Chem., Int. Ed.*, 2019, 58, 4968–4972.
- 62 S. Dong, Z. Yu, L. Guo, Y. Yang, C. Tu, R. Krishna and F. Luo, *Small*, 2023, **19**, 2304054.
- 63 S. Wang, E. V. Alekseev, J. Diwu, W. H. Casey, B. L. Phillips, W. Depmeier and T. E. Albrecht-Schmitt, *Angew. Chem., Int. Ed.*, 2010, 49, 1057–1060.
- 64 S. Fajal, S. Dutta and S. K. Ghosh, *Mater. Horiz.*, 2023, 10, 4083–4138.
- 65 T. He, Y. T. Zhang, X.-J. Kong, L. Yu, X.-L. Lv, Y. Wu, Z.-J. Guo and J.-R. Li, ACS Appl. Mater. Interfaces, 2018, 10, 16650–16659.
- 66 W. Mandal, S. Fajal, S. Mollick, M. M. Shirolkar, Y. D. More, S. Saurabh, D. Mahato and S. K. Ghosh, ACS Appl. Mater. Interfaces, 2022, 14(17), 20042–20052.
- 67 L. H. Keith and W. A. Teillard, Environ. Sci. Technol., 1979, 13, 416–423.
- 68 A. K. Poswal, A. Agarwal, H. K. Poswal, D. Bhattacharyya, S. N. Jha and N. K. Sahoo, *J. Synchrotron Radiat.*, 2016, 23, 1–8.
- 69 R. Zhao, D. Chen, N. Gao, L. Yuan, W. Hu, F. Cui, Y. Tian, W. Shi, S. Ma and G. Zhu, Adv. Funct. Mater., 2022, 32, 2200618.
- 70 R. J. Drout, S. Kato, H. Chen, F. A. Son, K. Otake, T. Islamoglu, R. Q. Snurr and O. K. Farha, *J. Am. Chem. Soc.*, 2020, 142, 12357–12366.
- 71 R. J. Drout, K. Otake, A. J. Howarth, T. Islamoglu, L. Zhu, C. Xiao, S. Wang and O. K. Farha, *Chem. Mater.*, 2018, 30, 1277–1284.
- 72 F. Li, J. Zhui, P. Sun, M. Zhang, Z. Li, D. Xu, X. Gong, X. Zou, A. K. Geim, Y. Su and H.-M. Cheng, *Nat. Commun.*, 2022, 13, 4472.