Functionalized electrospun polymer nanofibers for treatment of water contaminated with uranium†

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Uranium (U) contamination of drinking water often affects communities with limited resources, presenting unique technology challenges for U⁶⁺ treatment. Here, we develop a suite of chemically functionalized polymer (polyacrylonitrile; PAN) nanofibers for low pressure reactive filtration applications for U⁶⁺ removal. Binding agents with either nitrogen-containing or phosphorous-based (e.g., phosphonic acid) functionalities were blended (at 1–3 wt%) into PAN sol gels used for electrospinning, yielding functionalized nanofiber mats. For comparison, we also functionalized PAN nanofibers with amidoxime (AO) moieties, a group well-recognized for its specificity in U⁶⁺ uptake. For optimal N-based (Aliquat® 336 or Aq) and P-containing [hexadecylphosphonic acid (HPDA) and bis(2-ethylhexyl)phosphate (HDEHP)] binding agents, we then explored their use for U⁶⁺ removal across a range of pH values (pH 2–7), U⁶⁺ concentrations (up to 10 μM), and in flow through systems simulating point of use (POU) water treatment. As expected from the use of quaternary ammonium groups in ion exchange, Aq-containing materials appear to sequester U⁶⁺ by electrostatic interactions; while uptake by these materials is limited, it is greatest at circumneutral pH where positively charged N groups bind negatively charged U⁶⁺ complexes. In contrast, HDPA and HDEHP perform best at acidic pH representative of mine drainage, where surface complexation of the uranyl cation likely drives uptake. Complexation by AO exhibited the best performance across all pH values, although U⁶⁺ uptake via surface precipitation may also occur near circumneutral pH values and at high (10 μM) dissolved U⁶⁺ concentrations. In simulated POU treatment studies using a dead-end filtration system, we observed U removal in AO-PAN systems that is insensitive to common co-solutes in groundwater (e.g., hardness and alkalinity). While more research is needed, our results suggest that only 80 g (about 0.2 lbs.) of AO-PAN filter material would be needed to treat an individual’s water supply (contaminated at ten-times the U.S. EPA maximum contaminant level for U) for one year.

Introduction

Uranium (U) contamination affects the drinking water of many consumers in the Four Corners region of the United States (Colorado, New Mexico, Arizona, and Utah), including indigenous communities such as the Navajo Nation.¹ Mining of U ore deposits in the region occurred between 1940 and 1980 but left a profound impact on the environment because
of the presence of thousands of abandoned and open mines. Over 500 abandoned mines containing residual U within waste rock are located on Navajo lands and contribute to U concentrations in unregulated water sources that exceed the U.S. Environmental Protection Agency Maximum Contaminant Level (US EPA MCL) of 30 μg L⁻¹. In surface waters and shallow aquifers, U is in the hexavalent state and forms the uranyl (UO₂²⁺) cation, which can further complex to ligands to form soluble species. Source waters in the region range from alkaline to circumneutral pH and are high in dissolved carbonate, leading to the formation of soluble uranyl complexes that can contribute to high concentrations of total U in drinking water sources. Some waters near mine waste sites can have pH values lower than 4 due to acid mine drainage.

For resource-constrained communities without reliable access to centralized water treatment systems, point-of-use (POU) and point-of-entry (POE) technologies are an attractive option for improving drinking water quality. For example, existing US EPA-approved small system compliance technologies (SSCT) for POU treatment of U⁶⁺ include ion exchange (IX) and reverse osmosis (RO) technologies, while activated alumina can also remove U⁶⁺ but is not listed as an SSCT. Although all of these approaches are capable of removing total U to levels below the US EPA MCL, these technologies can be difficult to use and maintain in underserved populations. For example, RO can involve high capital costs, requires high operating pressures with associated energy demand, and produces a concentrated waste brine that would need proper disposal. In places relying on unregulated water sources, as is the case in some locations within the Navajo Nation, such advanced technologies would be of limited value.

We have previously demonstrated the use of electrospinning to produce novel, chemically reactive membranes for simultaneous filtration of suspended particles and sequestration or destruction of dissolved chemical contaminants. For materials targeting dissolved metals, we have used surface-segregating surfactants, especially those with quaternary ammonium groups, to produce surface-functionalized polymeric nanofibers that effectively function as ion exchange materials. More recently, we have also used post-fabrication routes to introduce specific binding moieties on the nanofiber surface. For U⁶⁺, a popular moiety is amidoxime (AO), which is highly specific for the uranyl cation and can be produced via reduction of polyacrylonitrile (PAN), a polymer commonly used in electrospinning, with hydroxylamine (Table 1). Indeed, we have previously amidoximated PAN (AO-PAN) nanofibers for the selective concentration of U⁶⁺ to improve environmental sensing via surface enhanced Raman spectroscopy (SERS). However, we have not yet explored this material more generally for water treatment applications, where timescales of uptake, material capacity, performance across pH, and response to common co-solutes (e.g., alkalinity and hardness typical of U-contaminated waters) will likely influence performance.

Here, we fabricate various functionalized PAN nanofibers using electrospinning and explore their applications for the removal of U⁶⁺ from contaminated water supplies. To introduce U⁶⁺-specific binding sites, we not only used post-synthesis amidoximation of PAN nanofibers, but we also developed synthesis recipes integrating various N- and P-containing surfactants to produce functionalized nanofibers (Table 1). For N-containing surfactants, we focused on those with quaternary ammonium groups (e.g., tetrabutyl ammonium bromide or TBAB and Aliquat® 336) because these are analogous to strong base anion exchange sites and thus would be expected to electrostatically bind negatively charged U⁶⁺ complexes typical of circumneutral pH.

| Table 1 | (a) N-containing (quaternary ammonium) and (b) P-containing binding agents, as well as (c) the amidoximation reaction of PAN, used for U capture herein |

![Diagram]

- **N-containing (Quaternary Ammonium) Binding Agents**
  - Aliquat® 336 (Ag)
  - Tetrabutyl ammonium bromide (TBAB)

- **P-containing Binding Agents**
  - Bis(2-ethylhexyl) phosphate (HDEHP)
  - Tributylphosphate (TBP)
  - Hexadecylphosphonic acid (HPDA)
  - Diallyl dimethylphosphonate (DAMP)
  - n-Octadecylphosphonic acid (CMPO)

- **Amidoximation Reaction**
  - PAN + NH₂OH·HCl → AO-PAN
P-containing binding agents included surfactants with phosphoric or phosphonic acid groups [e.g., di(2-ethylhexyl)phosphoric acid (HDEHP) and heptadecylphosphonic acid (HDPA)] that form strong complexes with U$^{6+}$ and may ultimately promote more extensive removal via surface precipitation. We also explored the integration of commercially available P-based extractants [diamyl amyl phosphonate (DAAP), n-octyl[phenyl]-N,N,N-diisobuty carbamoylmethylphosphine oxide (CMPO), and tributyl phosphate (TBP)] that are commonly marketed (e.g., Eichrom’s TRU Resin with CMPO) to separate U$^{6+}$ from complex media in nuclear waste streams. For example, phosphate esters such as TBP have been used extensively in the nuclear fuel cycle to selectively extract U$^{6+}$. We also explored the integration of commercially available P-based extractants [diamyl amyl phosphonate (DAAP), n-octyl[phenyl]-N,N,N-diisobuty carbamoylmethylphosphine oxide (CMPO), and tributyl phosphate (TBP)] that are commonly marketed (e.g., Eichrom’s TRU Resin with CMPO) to separate U$^{6+}$ from complex media in nuclear waste streams. For example, phosphate esters such as TBP have been used extensively in the nuclear fuel cycle to selectively extract U$^{6+}$. We also explored the integration of commercially available P-based extractants [diamyl amyl phosphonate (DAAP), n-octyl[phenyl]-N,N,N-diisobuty carbamoylmethylphosphine oxide (CMPO), and tributyl phosphate (TBP)] that are commonly marketed (e.g., Eichrom’s TRU Resin with CMPO) to separate U$^{6+}$ from complex media in nuclear waste streams. For example, phosphate esters such as TBP have been used extensively in the nuclear fuel cycle to selectively extract U$^{6+}$.

After nanofiber synthesis and characterization of their physical and chemical properties, we tested eight different functionalized materials for U$^{6+}$ uptake to identify the most promising candidates for further material development. The performance of the most promising materials for U$^{6+}$ capture was then explored across a range of pH, dissolved U$^{6+}$ concentrations, and water chemistries, including in a dead-end, flow through filtration system typical of low-pressure POU water treatment. Outcomes of this work help to establish the viability of functionalized nanofiber filters as low pressure water treatment technologies for use in areas afflicted by U$^{6+}$ contamination of limited freshwater resources.

Materials and methods

Reagents

A complete list of reagents can be found in the ESI. Nanoﬁbers of PAN (MW 150,000, Aldrich) were fabricated by electrospinning on a support layer of polyvinylidene difluoride (PVDF; MW 180,000, Aldrich). Binding agents (Table 1) included N-containing tetrabutylammonium bromide (TBAB; Sigma Aldrich) and Aliquat® 336 (Aq; Alfa Aesar) and P-based binding agents tributyl phosphate (TBP; Sigma Aldrich), diamyl amyl phosphate (DAAP; Sigma Aldrich), CMPO (Carbosynth; 98%), bis(2-ethylhexyl) phosphate (HDEHP; 97%, Aldrich), and hexadecyl phosphonic acid (HDPA; 97%, Aldrich). Amidoximation of PAN used hydrazylamine hydrochloride (98%, Aldrich) and sodium hydroxide (97.0%, Fisher Scientific).

Electrospinning

Full details of nanofiber synthesis are provided in the ESI. Nanofiber mats were synthesized on a custom-built electrospinning rig described in our previous work. PAN nanofibers were deposited on a layer of PVDF nanofibers to enhance the mechanical stability of the materials; PVDF was electrospun first, and after completion of the PVDF layer, a layer of PAN was subsequently deposited via sequential electrospinning. The resulting bilayer material contained 50 wt% PAN and 50 wt% PVDF. For production of AO-PAN, the two-layer polymer structure was reacted with hydroxylamine according to the amidoximation procedure described in the ESI which was adapted from our earlier work.

Mats with surfactant-based binding agents followed the same synthesis procedure but used PAN sol gel precursor solutions containing the desired binding agent. Most surfactant-functionalized mats were prepared using a precursor solution with 7 wt% PAN and up to 3 wt% of binding agent dissolved in DMF (all wt% are reported relative to the total weight of sol gel). Because HDPA exhibited limited solubility in DMF, mats containing HDPA were only prepared with 6 wt% PAN and either 0.5 or 1 wt% HDPA dissolved in DMSO. All solutions were then stirred at 60 °C for 2 h at 700 RPM to ensure complete dissolution of the surfactants and a homogenous precursor solution.

Nanofiber characterization

Nanofiber morphology and diameter were determined through imaging with scanning electron microscopy (SEM; S-4800, Hitachi). Fourier transform infrared spectroscopy (FTIR; Nicolet™ iS™ 50 FTIR Spectrometer) was used to examine all functionalized nanofiber formulations to confirm the presence of PAN and the binding agent. Most pore volume and specific surface area were determined by N$_2$-BET adsorption isotherms on a Quantachrome NOVA 4200e Analyzer. X-ray photoelectron spectroscopy (XPS) was performed using a Kratos Axis Ultra spectrometer to characterize the near surface region (∼5–10 nm) of the functionalized nanofibers before and after U$^{6+}$ uptake experiments. Additional materials characterization details can be found in the ESI.

Uranium uptake experiments

Initial comparison of different binding agents. To identify the most promising binding agents, uptake experiments with functionalized nanofibers were conducted with 10 μM total U$^{6+}$ at either pH 2 (Milli-Q Ultrapure water adjusted with 5 N HNO$_3$) or pH 6.8 (10 mM HEPES, which is commonly used in environmental and biological systems, including our prior work with U$^{6+}$ uptake and sensing on AO-PAN). These pH values were chosen for their relevance to U$^{6+}$ treatment systems, simulating remediation of U-contaminated acid mine drainage (pH 2) and treatment of U-contaminated drinking water sources (pH 6.8). Solutions of U$^{6+}$ were prepared by diluting a 1000 mg L$^{-1}$ depleted uranium nitrate (SPEX CertiPrep) stock to the desired initial concentration, typically 1 or 10 μM (0.24 or 2.4 ppm, or mg L$^{-1}$, as U$^{6+}$, respectively), in a 50 mL plastic conical vial. To initiate an uptake study, a functionalized PAN mat was added to a conical vial at a mass loading of 0.25 g L$^{-1}$ (∼0.25 cm$^2$ L$^{-1}$) and then incubated while mixing. Rate experiments confirmed that uptake of U was relatively rapid in all systems (see Fig. S1), with most uptake achieved in the initial 2 h. Because a small amount of U$^{6+}$ uptake continued over longer timescales in some
systems, all sorption studies were allowed to proceed for 16 h, at which point there was no significant change in solution concentration over time for any system (i.e., equilibrium). In all instances, minimal pH drift was observed (<0.1 pH units) during the 16 hour reaction period, after which mats were removed from the solution and analyzed for their sorbed U⁶⁺ content as described below.

**pH edge and isotherm experiments.** For the most promising materials identified from initial uptake studies, more in depth pH edge and isotherm experiments were conducted using experimental systems identical to those described above. For pH edge experiments, systems were assembled at initial pH values between 2 and 7. To avoid any influence of different buffers at different pH values, all experiments were conducted in Milli-Q Ultrapure water that was adjusted to the desired pH with either 5 M NaOH or HNO₃. pH edge experiments were conducted at initial U⁶⁺ concentrations of both 1 and 10 μM, and all systems were allowed to react for 16 hours to achieve equilibrium. At the conclusion of the experiment, the final pH value of each reactor was recorded to measure pH drift during incubation, which was typically <0.5 pH units.

For sorption isotherms, initial U⁶⁺ solutions were prepared at concentrations of 0.1, 0.5, 1, 5 and 10 μM U. Sorption isotherms were conducted at either pH 2 (Milli-Q Ultrapure water adjusted with either 5 N NaOH or 5 N HNO₃) or pH 6.8 (10 mM HEPES). All other experimental conditions are as described previously for pH edge systems (e.g., 16 h incubation period).

**Simulated POU filtration.** The performance of optimal nanofiber mat formulations was evaluated in a dead-end, flow-through filtration system (Fig. S2†) to simulate their application in water treatment. The filter holder (Cole-Parmer) had a 25 mm outer diameter with an active filtration area of 3 cm². Mats were cut to fit within this holder and typically weighed between 10–13 mg per layer of mat (depending on the formulation). To increase the mass of nanofibers used for treatment, thicker filters were created by stacking multiple layers of material with the same effective filtration area, thereby increasing the residence time for U-containing influent within the nanofiber mat. Flow-through conditions were created using influent flowrates of 0.4 or 0.8 mL min⁻¹ driven by a 60 mL syringe loaded on a syringe pump (New Era Pump Systems, Inc.). These flowrates produce fluxes ranging from 80 to 160 LMH (0.4 to 0.8 mL min⁻¹ assuming 3 cm² active area), which correspond to the high range for ultrafiltration (80 LMH) and low end of microfiltration (160 LMH). Filters were preconditioned with 20 mL of a 10 mM HEPES solution (pH 6.8) followed by either 120 or 240 mL of a 1 μM U⁶⁺ (0.24 ppm) solution in 10 mM HEPES at pH 6.8. Additional tests were conducted using solutions with 500 mg L⁻¹ Ca²⁺ and 500 mg L⁻¹ HCO₃⁻ to evaluate the influence of environmentally relevant ions. Effluent was collected in 4 mL samples for every 10 mL of filtered solution for analysis of dissolved U by ICP-MS as described below.

**Analytical methods**

For batch uptake systems, U analysis was conducted via liquid scintillation counting (LSC) using a ²³²U radiotracer (NIST traceable standard, Eckert & Ziegler) with a 3.5 Bq spike per 20 mL of solution. The activity of solutions was measured by adding 2 mL aliquots from each reactor to 10 mL of Ecolite scintillation cocktail (MP Biomedicals) in a 20 mL scintillation vial. Sorbed uranium was measured by moving the mat from the reactor and placing it into a 20 mL scintillation vial with 10 mL of Ecolite scintillation cocktail. Vials were shaken and left overnight to dark adapt (energy in scintillation cocktail from light is able to leave) and provide ample time for the polymer mats to dissolve in the scintillation cocktail. Samples were then counted on a liquid scintillation counter (LSC; Packard 1600CA Tri-Carb liquid scintillation analyzer) for 40 minutes. The range of 100 and 2000 keV was used to exclude beta signals produced by daughter isotopes of ²³⁸U, ²³⁴Th and ²³⁴Pa. Generally, in samples collected from equilibrated experimental systems, LSC analysis of both solution phase and sorbed uranium indicated complete mass balance.

ICP-MS analysis was used to analyze the effluent for U⁶⁺ collected from the flow through systems. Effluent samples analyzed by ICP-MS analysis were acidified with 2% HNO₃ (trace metals grade, Aldrich) and filtered with 0.45 μm filters prior to analysis on an Agilent Technologies 7900 ICP-MS. Argon was used as the carrier gas in low matrix mode and no collision gas was used. Mass-to-charge ratios of 7, 89, and 205 were used for tuning of the instrument prior to running calibration standards and samples in triplicate. ²⁰⁹Bi (Inorganic Ventures) was used as the internal standard at a concentration of 10 ppb. We note, initial studies indicated that trace amounts of some surfactants likely leached from the functionalized mats during uptake experiments, and this dissolved surfactant residual present in samples interfered with ICP-MS analysis. To avoid this interference, all surfactant-functionalized materials were washed with DI water prior to use in uptake experiments where samples required ICP-MS analysis (e.g., flow through systems). The washing procedure involved placing 5 mg of a functionalized PAN mat in a 50 mL conical vial with 10 mL of Milli-Q Ultrapure water. Vials were mixed end over end for 24 hours, while three changes of the water were performed over that time interval.

**Results and discussion**

**Comparison of functionalized PAN Nanofibers for U⁶⁺ uptake**

Initial uptake studies explored the performance of PAN nanofibers functionalized with different P- and N-containing binding agents as a function of their wt% in PAN at pH 2 and pH 6.8 (Fig. 1). Of the P-functionalized materials, integration of HDEHP (at pH 2) and HDPA (at pH 2 and 6.8) resulted in the greatest uptake (between 35–55% uptake for 0.25 g L⁻¹ of nanofibers and initially 10 μM U⁶⁺), with U⁶⁺ binding on other P-containing materials (e.g., TBP, CMPO,
sites in the fibers after electrospinning, as some binding also cannot rule out differences in the location of the binding.

10.25 g L⁻¹ of each mat. Experiments were conducted in 10 mM HEPES at pH 6.8 and water acidified to pH 2 with HNO₃.

and DAAP) being very limited (~10%) or negligible at both pH values. Performance of HDEHP was effectively invariant over the concentrations in PAN we explored (1 and 3 wt%); thus, all additional work with HDEHP was conducted at 1 wt% to minimize the amount of reagent needed for synthesis. For HDPA-containing materials, optimal performance was observed at a concentration of 0.5 wt% in PAN.

For N-containing binding agents, U⁶⁺ uptake was only observed at pH 6.8, with no detectable uptake at pH 2. Generally, Aq-containing materials outperformed those with TBAB. Although comparable uptake of U was observed for 1 wt% in PAN of either Aq or TBAB, increasing the concentration to 2 wt% resulted in higher uptake with Aq but lower uptake for TBAB-containing materials. We have previously found that the mass ratio of quaternary ammonium surfactant to polymer can influence the performance of the functionalized PAN.¹⁷ Accordingly, additional studies exploring the influence of Aq concentration between 0.5 to 4 wt% were conducted, revealing maximum U⁶⁺ uptake at 2 wt% in PAN. All additional uptake studies were conducted at this optimal Aq loading.

AO-PAN exhibited the greatest uptake at pH 6.8, nearly double of the removal displayed by either HDPA- or Aq-containing materials at the same pH value. At pH 2, AO-PAN also produced measurable uptake of U⁶⁺ at a level comparable to that observed for HDEHP-containing materials but below the extent of removal achieved with HDPA-functionalized PAN.

We note that beyond basic characterization to ensure all materials tested in Fig. 1 were comparable in morphology (i.e., nanofibers), no additional investigations were performed to better understand differences in the performance of P- and N-containing binding agents. Several possibilities exist, including differences in the chemistry of the binding sites available on the functionalized PAN (e.g., the relative affinity for U⁶⁺ uptake on different P-containing functionalities). We also cannot rule out differences in the location of the binding sites in the fibers after electrospinning, as some binding agents may surface segregate to a greater extent than others. Thus, it is certainly possible that the performance of materials with low U⁶⁺ uptake in Fig. 1 could be further optimized, but such work is beyond the scope of the current study.

Characterization of optimally performing nanofibers

Key characterization details for optimal nanofiber formulations including AO-PAN and those containing either 2 wt% Aq, 0.5 wt% HDPA, or 1 wt% HDEHP are summarized in Fig. 2. Overall, there were only a few notable differences between the functionalized PAN nanofibers explored for U uptake. The average diameter for all functionalized PAN nanofibers was between 110–160 nm, but the typical standard deviation from the nanofiber distribution (see histograms in Fig. S3†) indicates all diameters are statistically equivalent. Moreover, there was no obvious influence of increasing functionalization, either based on wt% of embedded P- or N-containing binding agents or amidoximation, on the average or distribution of nanofiber diameters. Similarly, the specific surface area for all materials fell between 11 and 21 m² g⁻¹ (with most being statistically equivalent based on the standard deviation from replicate analyses), with no clear trends in surface area based upon the amount or type of integrated binding sites. There were some modest differences in the pore volume of different materials, especially for HDEHP-containing nanofibers that exhibited pore volumes [13 (±5) × 10⁻³ cm³ g⁻¹] considerably lower than the other materials [for example, 45 (±5) × 10⁻³ cm³ g⁻¹ for PAN]. We speculate this could be an indication that HDEHP preferentially locates within the pore structure of PAN, blocking pore access. Analysis via FTIR (Fig. S4†) was consistent with expectations for PAN-based polymers, but typically revealed little evidence of the different functionalization routes we employed except for HDEHP and amidoximation. This is not necessarily surprising because FTIR is a bulk characterization technique and most binding agents were present at a relatively low wt% in the functionalized nanofibers.

pH-Dependent U uptake

At an initial U⁶⁺ concentration of 10 μM, PAN with 0.5 wt% HDPA exhibited among the highest uptake (>60% of total U⁶⁺) over the entire pH range (Fig. 3a). AO-PAN achieved its lowest removal at pH 2 (~40%), but U⁶⁺ removal increased with pH, producing relatively high and constant removal between pH 3 and 7 (~80%). PAN with 1 wt% HDEHP exhibited the opposite behavior relative to AO-PAN; its highest uptake was at pH 2 (~50%), but U⁶⁺ removal decreased markedly at pH 3 (~20%) and was maintained at this low level for all higher pH values explored. Finally, PAN with 2 wt% Aq exhibited the lowest removal overall [between 0–15%], but U⁶⁺ uptake did modestly increase with increasing pH values.

Notably, at a lower initial U⁶⁺ concentration of 1 μM (Fig. 3b), different pH-dependent removal trends were observed for some, but not all, materials. While trends in U⁶⁺
uptake at 1 μM were comparable to those observed at higher initial U⁶⁺ for HDEHP- and Aq-containing PAN, AO-PAN and HDPA-containing PAN exhibited different pH-dependent performance. For AO-PAN, this difference was only observed at higher pH values (pH > 5). Specifically, whereas removal was relatively constant (~80%) above pH 5 in 10 μM U⁶⁺ systems, uptake decreased steadily from pH 5 (~80%) to pH 7 (~40%) in 1 μM U⁶⁺ systems. A much greater difference in performance between low and high concentration U⁶⁺ systems was observed with HDPA-functionalized PAN. While removal at 10 μM U⁶⁺ was greater than 60% across all pH values, the removal in 1 μM U⁶⁺ systems was greatest at pH 2 (~50% U⁶⁺) and decreased steadily until pH 4 (~10%), above which uptake was minimal.

A possible explanation for the difference in performance between 1 μM and 10 μM U⁶⁺ systems for AO-PAN and HDPA-containing materials could be removal via surface precipitation at high initial U⁶⁺ concentrations, similar to processes previously reported to occur on mineral phases and functionalized polymers. For AO-PAN, for example, the exact binding mechanism for uranium to amidoxime is still widely disputed, with arguments for either monodentate (binding with either N or O) or bidentate (binding to both N and O) complexes in prior investigations. Pekel et al. suggested
that deprotonation of the imine group was important for chelation to uranyl by exchange of $H^+$ with $UO_2^{2+}$ while Hirotsu et al.\textsuperscript{37} reported that ligand exchange (and ion exchange of $H^+$ depending on pH) occurs during uranyl uptake. In both 1 and 10 $\mu$M U systems at pH 2, similar uranium removal occurs ($\sim$40%) with 100% of the uranyl species being $UO_2^{2+}$, suggesting that the high $H^+$ concentration competes with $UO_2^{2+}$ in the amidoxime group. A decrease in the $H^+$ concentration (i.e., increasing pH between 3 and 5) results in much greater ($\sim$80%) $UO_2^{2+}$ removal. Hydrolysis of $UO_2^{2+}$ starts at pH 4, and it is no longer the dominant species by pH 5 (see Guillaumont et al.\textsuperscript{38} and speciation diagrams for 1 and 10 $\mu$M $UO_2^{2+}$ solutions in Fig. S5\textsuperscript{1}). In 10 $\mu$M systems, speciation diagrams suggest that insoluble $UO_2(OH)_2\cdot H_2O$ is the dominant species by pH $\sim$ 5.5, and we suspect this species may be precipitating on the surface based upon the high ($\sim$80%) removal still observed at pH 6 and 7 in 10 $\mu$M $UO_2^{2+}$ systems. In contrast, the decrease in removal with increasing pH observed in 1 $\mu$M $UO_2^{2+}$ systems ($\sim$70% at pH 6 and $\sim$40% at pH 7) may be indicative of speciation changes that occur above pH 5; we expect $UO_2OH^+$ to be the dominant form at pH 5 and 6 ($\sim$50% and $\sim$40% of total $UO_2^{5+}$, respectively) followed by $UO_2(OH)_2\cdot H_2O$ at pH 7 ($\sim$90% of total U). Because ligand exchange is expected to occur in these regions, the lower uptake may also be due to slower kinetics involved with ligand exchange.\textsuperscript{39}

For HDHP-functionalized PAN, uranyl phosphate complexes are known to have very low log $K_{sp}$ values (∼49.00 to −53.33)\textsuperscript{40} compared to that of hexavalent uranyl hydroxides (∼21.75 to −24.10)\textsuperscript{41} and uranyl carbonates (∼13.29 to −14.91),\textsuperscript{40} which enables uranium phosphates to precipitate in even acidic solutions. This behavior has been seen before with phosphate-functionalized TiO$_2$, where an insoluble sodium autunite (NaUO$_2PO_4$) complex formed after uranium sorption in acidic solutions (pH 2).\textsuperscript{42} In this earlier work, the mechanism of uptake was described as a combination of adsorption and surface complexation that shifts to surface precipitation;\textsuperscript{42} such a scenario may also be likely for $UO_2^{5+}$ removal on HDHP-functionalized nanofibers in our 10 $\mu$M $UO_2^{5+}$ systems, whereas only adsorption and surface complexation occur in our 1 $\mu$M $UO_2^{5+}$ systems.

For HDEHP- and Aq-containing nanofibers, trends in pH-dependent removal lend insight into their mechanism of $UO_2^{5+}$ binding. For example, the sorption capacity of HDEHP-containing materials is reduced considerably above pH 3. HDEHP has a $K_{sp}$ of 1.47,\textsuperscript{43} and thus will become increasingly more deprotonated (i.e., more anionic) from pH 2 to pH 3. Over this same pH range, $UO_2^{5+}$ removal decreases from 50% to $\sim$10% in both 1 and 10 $\mu$M systems. Thus, $UO_2^{5+}$ removal does not appear to proceed via a purely electrostatic mechanism (i.e., positively charged $UO_2^{2+}$ bound by negatively charged HDEHP sites), suggesting that $UO_2^{5+}$ uptake may also occur by exchange of $H^+$ during uranyl coordination, which has been previously observed by Kiwan and Amin.\textsuperscript{44} Moreover, hydrolysis of the $UO_2^{2+}$ cation should not affect $UO_2^{5+}$ uptake with HDEHP because hydrolysis products are not abundant until pH 4 for solutions containing 1 or 10 $\mu$M $UO_2^{5+}$ (see Guillaumont et al.\textsuperscript{38} and Fig. S5\textsuperscript{1}). As a final consideration, the chemical differences between HDEHP and HDPA may also lend insight regarding the mechanism of $UO_2^{5+}$ uptake. HDEHP contains only one hydroxyl group available for $UO_2^{5+}$ binding, whereas HDPA has multiple hydroxyls that may allow it to chelate and precipitate $UO_2^{5+}$ in a manner similar to the phosphate anion.

At both initial concentrations (1 and 10 $\mu$M), Aq-containing nanofibers produced a slight increase in $UO_2^{5+}$ uptake with increasing pH. This behavior likely reflects that uptake of $UO_2^{5+}$ by Aq is dependent on the fraction of anionic uranyl species present in solution. Aq is positively charged across the pH range investigated, and as a strong base ion exchanger, it has been shown to bind negatively charged uranium complexes.\textsuperscript{45–47} We therefore hypothesize that anion exchange is the main mechanism for uptake of $UO_2^{5+}$ on Aq-functionalized nanofibers, but further verification of this mechanism is warranted. In fact, for pure aqueous systems,
anionic uranyl species (e.g. $\text{UO}_2(\text{OH})_2$) should not be formed until $\sim$ pH 7 (see Guillaumont et al. and Fig. S5†). Furthermore, while negatively charged species can form in the presence of carbonate (e.g., $(\text{UO}_2)_2\text{CO}_3(\text{OH})_2$) can form as early as pH 4, these anionic carbonate species are only produced at dissolved CO$_2$ concentrations higher than those in our experimental systems.$^{48}$

Sorption isotherms for U on functionalized PAN nanofibers

To explore their capacity for U$^{6+}$ uptake, sorption isotherms were collected with AO-PAN and PAN containing either 2 wt% Aq, 0.5 wt% HDPA, or 1 wt% HDEHP (Fig. 4). Functionalized PAN nanofibers were tested over a range of U$^{6+}$ concentrations that varied from just below its MCL in drinking water ($\sim$0.1 μM) to the more extreme levels of U$^{6+}$ contamination that may be present in some affected water resources (10 μM). We only developed isotherms at the optimal pH value observed for each functionalized material in pHedge experiments. PAN nanofibers functionalized with HDPA and HDEHP were tested in acidic conditions (pH 2), whereas isotherms for Aq-containing PAN and AO-PAN were conducted at pH 6.8.

Overall, a trend of increasing solution phase U$^{6+}$ concentration resulted in increased sorbed U$^{6+}$ concentrations for all materials, and the extent of uptake generally agreed well with our other experimental results (see Fig. 1 and 3). To model U$^{6+}$ uptake, we used the empirical Freundlich isotherm model $[C_{\text{sat}} = K_f(C_{\text{aq}})^{1/n}]$, where $K_f$ is the Freundlich isotherm parameter and $n$ is the degree of linearity] because we observed no clear evidence consistent with surface site saturation (as would be expected for a Langmuir-type isotherm). The parameters for the Freundlich isotherms determined by non-linear regression analysis are summarized in Fig. 4 for each functionalized material. At pH 6.8, AO-PAN far exceeded the uptake of Aq-containing mats, and uptake on AO-PAN was clearly non-linear ($n = 1.5 \pm 0.2$). At pH 2, U$^{6+}$ sorption on HDPA-functionalized PAN ($n = 1.5 \pm 0.7$) was considerably greater than HDEHP-containing materials ($n = 0.8 \pm 0.2$), with model outputs indicating that sorption isotherms did not significantly differ from linearity over the range of U$^{6+}$ concentrations explored (although relatively large standard deviations in model fits were observed because of the modest degree of variability in uptake observed between two replicate isotherm experiments).

Even when uptake was clearly non-linear (e.g., AO-PAN), we did not achieve the sorption capacity of any materials using these isotherm conditions. At the highest initial U$^{6+}$ concentration explored of 10 μM (or 2.4 mg L$^{-1}$), corresponding concentrations for sorbed U$^{6+}$ were approximately 4 and 10 μg mg$^{-1}$ at pH 2 for HDEHP- and HDPA-containing nanofibers, respectively, and approximately 2 and 8 μg mg$^{-1}$ at pH 6.8 for Aq-containing and AO-PAN nanofibers, respectively. A prior investigation of AO-PAN nanofibers with a polystyrene core shell reported a maximum sorbed concentration of 130 μg mg$^{-1}$ (conditions: 1 g mat L$^{-1}$; pH 4).$^{47}$ Phosphate-functionalized polyethylene had a maximum sorbed concentration of 180 μg mg$^{-1}$ (conditions: 0.2 g mat L$^{-1}$; pH 8.2).$^{48}$ Strong base anion exchangers, similar to Aq, have not been used for U$^{6+}$ uptake in nanofibers but show high uptake in resins at $\sim$50 μg mg$^{-1}$ in groundwater (conditions: initial U$^{6+}$ concentration of 1200 μg L$^{-1}$; pH 6.5; flow through system).$^{47}$ Although many of these prior investigations report sorbed U$^{6+}$ concentrations that are greater than what we report for the functionalized nanofibers herein, we note that several of these earlier works used initial U$^{6+}$ levels far exceeding the concentrations used in our experimental systems. Thus, we cannot rule out that some of these
high levels of U\textsuperscript{6+} uptake may reflect U\textsuperscript{6+} removal via surface precipitation, as we suspect may occur at high U\textsuperscript{6+} and high pH on AO-PAN and HDPA-functionalized PAN, leading to greater removal via multi-layer growth of a separate U-containing solid phase.

To probe the nature of surface bound U\textsuperscript{6+}, XPS analyses were collected for all functionalized materials after U\textsuperscript{6+} uptake experiments conducted with an initial concentration of 10 \textmu{}M. XPS analysis of these reacted nanofiber mats detected the presence of U\textsuperscript{6+} on the surface of all functionalized materials (Table S1). High resolution U 4f spectra (Fig. S6†) contained signals corresponding to the U 4f\textsubscript{7/2} and U 4f\textsubscript{5/2} doublet on HDPA-, HDEHP-, Aq- and AO-PAN functionalized materials. However, while confirming the presence of U\textsuperscript{6+} on the surface of all functionalized nanofibers, XPS analysis was unable to provide any greater details regarding the nature of U\textsuperscript{6+} surface species or complexes.

Simulated treatment in flow through systems

Break through curves showing normalized U\textsuperscript{6+} concentration (\textit{i.e.}, effluent concentration normalized to influent concentration; \textit{C}_{\text{effluent}}/\textit{C}_{\text{influent}}) as a function of volume of water treated from dead-end filtration flow through systems are shown in Fig. 5 for AO-PAN and HDPA-containing nanofiber filters at pH 6.8. For such curves, we define filter exhaustion or complete breakthrough as when the effluent U\textsuperscript{6+} concentration is equal to that of the influent concentration (\textit{C}_{\text{effluent}} = \textit{C}_{\text{influent}}), which would mean either that the materials are saturated (\textit{i.e.}, all binding sites are occupied and thus not capable of removing any uranium) or that timescales for U\textsuperscript{6+} uptake on the remaining available binding sites are far slower than the residence time for U\textsuperscript{6+} in the nanofiber filter system. With an influent concentration of 1 \textmu{}M U\textsuperscript{6+} (240 \mu{}g L\textsuperscript{-1}) at pH 6.8, we note that any normalized concentration above \textasciitilde{}0.1 would be considered above the MCL for uranium (30 \mu{}g L\textsuperscript{-1}).

For AO-PAN (Fig. 5a), the lowest filter mass tested (13 mg) did not show complete breakthrough, but produced approximately constant, incomplete (~40%) removal of U\textsuperscript{6+} where \textit{C}_{\text{effluent}} was \textasciitilde{}60% of \textit{C}_{\text{influent}}. Increasing the mass of AO-PAN (from 13 mg to 26 mg by adding a second filter layer) resulted in effectively complete removal of U\textsuperscript{6+}. Based on these results, U\textsuperscript{6+} uptake on AO-PAN filters appears kinetically limited under our experimental conditions. At lower filter mass (13 mg), breakthrough was approximately steady state; complete saturation of the filter did not occur (\textit{i.e.}, there was always some residual capacity for U\textsuperscript{6+} removal), but U\textsuperscript{6+} was present in the effluent and the effluent concentration was not changing over time. Addition of more filter mass (from 13 mg to 26 mg) increased the contact time between the U\textsuperscript{6+}-containing solution and the AO-PAN filter, which in turn resulted in near-complete removal of U over the duration of the 120 mL filtration experiment. Notably, at the conclusion of the experiment with the 26 mg filter, the mass of U\textsuperscript{6+} captured was \textasciitilde{}1.1 \mu{}g mg\textsuperscript{-1} after treating 120 mL of water, which is well below the maximum sorbed concentration of \textasciitilde{}8 \mu{}g mg\textsuperscript{-1} observed in batch isotherm experiments with AO-PAN (see Fig. 3). This suggests that AO-PAN materials still have considerably more sites available for U\textsuperscript{6+} binding. A second run of a 26 mg filter over 240 mL of 1 \textmu{}M U\textsuperscript{6+} influent revealed removal of all influent uranium to levels that were below detection in the effluent and thus below the EPA MCL (Fig. S7† all \textit{C}_{\text{effluent}} values were below detection or 1 \mu{}g L\textsuperscript{-1} via our ICP-MS analytical method). Once again, this sample still had not reached saturation and the amount of U\textsuperscript{6+} bound on the reacted AO-PAN filter (~2.2 \mu{}g mg\textsuperscript{-1}) was only \textasciitilde{}25% of the max U\textsuperscript{6+} sorption found in batch. Once again, this supports kinetically limited U\textsuperscript{6+} removal in AO-PAN systems, where thicker filter materials or lower flow rates will produce higher residence times and better removal performance.

Fig. 5 Normalized concentration (effluent concentration divided by influent concentration) of U\textsuperscript{6+} as a function of the volume treated in a dead-end filtration setup with (a) AO-PAN and (b) 0.5 wt% HDPA-functionalized PAN. Experiments used an influent concentration of 1 \textmu{}M in 10 mM HEPES (pH 6.8) and a flowrate of 0.8 mL min\textsuperscript{-1} (160 LMH), unless otherwise indicated. For AO-PAN, results are shown for different masses (thicknesses) of filters (13 and 26 mg), replicate filters (1 and 2) and more complex solution chemistries (500 mg L\textsuperscript{-1} of Ca\textsuperscript{2+} or HCO\textsubscript{3}\textsuperscript{-} adjusted to pH 6.8). For HDPA-functionalized materials, four replicate experiments (1 through 4) with 20 mg filters are shown.
For AO-PAN filters, the presence of Ca\(^{2+}\) (as a competing ion to simulate hardness) and CO\(_3\)\(^{2-}\) (as a ligand for uranyl from alkalinity) had no influence on U\(^{6+}\) removal (Fig. 5a). Using higher mass filters (26 mg), there was no detectable U\(^{6+}\) in the filter effluent across 120 mL of treated volume for either influent solution. Sorbed uranium from the Ca\(^{2+}\) and CO\(_3\)\(^{2-}\) runs were nearly identical to the experiments performed in the absence of competing ions, with U\(^{6+}\) contents of \(\sim 1.2 \mu g \text{ mg}^{-1}\) for all three trials (as determined by LSC analysis of the reacted filter). The lack of interference from Ca\(^{2+}\) and CO\(_3\)\(^{2-}\) may be due to the chemical complexity of U\(^{6+}\) by AO groups on the surface of the mats as opposed to electrostatic interactions that could potentially be impacted by co-solute ions. It should also be noted that the pH of the solution varied from 6.8 to 7.5 over the course of experiments containing the CO\(_3\)\(^{2-}\) anion, suggesting that some HCO\(_3\)\(^{-}\) may have been scavenged by AO-PAN during the run by either amidoxime or residual nitrile groups.

Different behavior was observed in flow through experiments performed with the HDPA-functionalized filters. Results from four replicate experiments are shown in Fig. 5b. Partial U\(^{6+}\) removal was observed with each HDPA-containing filter, with detectable U\(^{6+}\) in most effluent samples across the four replicate studies. The degree of U\(^{6+}\) removal was also highly variable from one experiment to the next, with some systems routinely achieving more than 80% removal of influent U\(^{6+}\) (at 1 \(\mu M\)), while much less removal and more rapid breakthrough was observed in other instances. Another noteworthy feature observed in all systems was a period of increasing U\(^{6+}\) removal after an initial period of more rapid breakthrough, observed by the clear localized maxima in \(C_{\text{effluent}}/C_{\text{influent}}\) values in each of the four replicate experiments (see maxima after 40–80 mL of treated influent in Fig. 5b).

We propose that these unique U breakthrough profiles result from the mechanism of surface binding responsible for U\(^{6+}\) removal in HDPA-filter systems. From pH edge experiments at elevated U\(^{6+}\) concentration (10 \(\mu M\)), surface precipitation likely contributes to U\(^{6+}\) removal at near-neutral pH values. In contrast, from pH edge experiments at lower initial U\(^{6+}\) concentrations (1 \(\mu M\)), more limited removal was observed by HPDA-containing nanofibers at near-neutral pH, with any uptake presumably occurring via complexion between the phosphonic acid group on HDPA and soluble U\(^{6+}\) species. By analogy, we would expect initial removal in our flow through systems to occur via complexion but be relatively limited, consistent with the early periods of U\(^{6+}\) breakthrough observed in filter effluent. We would also expect that after some period of filter exposure to influent U\(^{6+}\) enough U\(^{6+}\) would be bound on the HDPA-functionalized surface to initiate formation of higher order U\(^{6+}\) species (e.g., dimers, trimers, oligomers and eventually a separate surface phase). If the rate of formation of these higher order species (resulting from surface bound U\(^{6+}\) interacting with dissolved U\(^{6+}\) species) is faster than the rate at which available HDPA sites form new surface complexes with dissolved U\(^{6+}\) species, we would anticipate the extent of U\(^{6+}\) removal in our filter systems to increase over time.

Such a biphasic mechanism for U\(^{6+}\) removal (i.e., first HDPA complexing U\(^{6+}\) followed by formation of higher order U\(^{6+}\) species through bound U-soluble U interactions) would likely explain the high variability observed in breakthrough curves for HDPA-containing filters in Fig. 5b. A critical point in the breakthrough curve will be when formation of higher order surface U\(^{6+}\) species begins, and it is likely the occurrence of such a transition point would be dependent on highly localized factors related to the flow path through the nanofiber filter. For example, if we consider the amount of U\(^{6+}\) mass accumulated in the filter over time (Fig. S8†), a clear increase in the rate of U\(^{6+}\) removal is observed between 40–60 mL of treated influent for all replicates, at which we suspect the transition from U\(^{6+}\)-complexation by HDPA to formation of higher order U\(^{6+}\) surface species occurs. Notably, however, in all cases, the surface U\(^{6+}\) concentration is relatively low (on the order of 1.2 \(\mu g \text{ mg}^{-1}\) mat or less; see Fig. S8†). Indeed, because of the relatively low loading of surface U\(^{6+}\), far less than observed for U-containing samples previously characterized spectroscopically (see Fig. S5†), we were unable to detect any surface U\(^{6+}\) via XPS on these reacted filters to further explore differences in bound species as a function of filter run time. We are currently exploring the use of other spectroscopic methods (e.g., XAFS) which may be better suited for examining the nature of bound U\(^{6+}\) in HDPA nanofiber filtration systems.

Environmental implications

In this work, we have produced various functionalized nanofibers for binding of U\(^{6+}\). Of the materials we explored, the strongest performance across all system conditions was AO-PAN, which has been widely used for capture of U\(^{6+}\) from various matrices. AO-PAN exhibited high U\(^{6+}\) capacity and sustained performance during filtration, even in the presence of more complex solution compositions (e.g., hardness and alkalinity). Based on our dead-end filtration experiments, and assuming that the average person consumes 2 L of water daily, our results suggest it would only require 80 g (about 0.2 lbs.) of AO-PAN filter material to treat water contaminated with 1 \(\mu M\) U\(^{6+}\) to levels below US EPA standards and our method of detection (e.g., 1 \(\mu g \text{ L}^{-1}\) via our ICP-MS analytical method) for one year.

While other materials exhibited less capacity for U\(^{6+}\) uptake, there still may be advantages to these alternative formulations. From a fabrication standpoint, amidoximation requires post-processing of electrospun PAN and uses highly concentrated and harsh reagents. The integration of N- and P-containing surfactants directly into the electrospinning sol gel affords more simplicity in filter fabrication, with less generation of chemical waste. Further, in applications of these materials to sequester and concentrate U\(^{6+}\) for biomonitoring or sensing, where information about solution phase speciation may be desirable, the ability to leverage different binding
agents to preferentially sequester separate U⁶⁺ species may be advantageous. For example, Aq and TBAB were included herein because of the prior use of N-containing functionalities in ion exchange, and thus these surfactants would be well-suited to specifically capture anionic U⁶⁺ species.

Future work is needed to better understand the nature of surface U⁶⁺ species on each of the most promising functionalized PAN nanofibers. Herein, the levels of surface-bound U⁶⁺ generated in our experimental systems prohibited extensive surface characterization. In particular, the mechanism of U⁶⁺ sorption on HPDA-functionalized nanofibers merits additional investigation based upon results from our flow through systems, which suggest that the surface U⁶⁺ species may change over time with increasing total U⁶⁺ bound to the nanofiber surface. Characterization of the bound U⁶⁺ species on HDPA and other functionalized nanofibers will be important to better predict long-term filter performance, including the potential for inadvertent U⁶⁺ release during water treatment applications and the potential for filter regeneration and reuse once saturation capacity is achieved.

Conflicts of interest
The authors have no conflicts of interest to declare.

Acknowledgements
Research reported in this work was supported by the National Institute of Environmental Health Sciences of the National Institutes of Health under award number R01ES027145. Additional support for MEC was provided through a National Research Traineeship (NRT) grant from the National Science Foundation (NSF) under award number DGE-1633098 and through an NSF Graduate Research Fellowship. The authors would also like to acknowledge the two anonymous reviewers whose comments improved the clarity and impact of this work.

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