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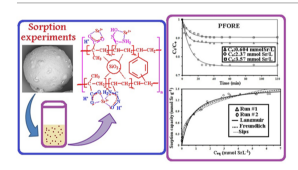
Controlled bi-functionalization of silica microbeads through grafting of amidoxime/methacrylic acid for Sr(II) enhanced sorption

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HIGHLIGHTS

- Bi-functionalization of silica particles increases Sr(II) sorption properties.
- Methacrylic (80%)/amidoxime (20%) substitution ratio is optimum.
- Maximum sorption capacity reaches
 1.57 mmol Sr g⁻¹ at pH 8.
- Total Sr(II) desorption and weak loss in sorption properties at the fifth recycling.
- The sorbent is remarkably selective to Sr(II) in seawater.

GRAPHICAL ABSTRACT



ABSTRACT

Silica microspheres are functionalized through the grafting of either polymethacrylic acid or amidoxime groups (mono-functional composite, PMAA/SiO $_2$ and PAO/SiO $_2$, respectively) or in combination (bi-functional composite, PAO $_x$ MAA $_{100-x}$ /SiO $_2$). The materials are characterized by textural analysis, TGA, SEM-EDX, elemental analysis, FTIR, XPS, pH $_{PZC}$ in order to confirm the synthesis route, identify the reactive groups and support the interpretation of sorption mechanisms toward Sr(II). The sorption of Sr(II), at optimum pH 8, involves amine groups and hydroxyl/carboxyl groups in the bi-functional composite sorbent. The comparison of sorption properties shows that maximum sorption requires a majority of carboxylic groups from polyacrylic compared with amidoxime functions; optimum formulation: PAO $_{20}$ MAA $_{80}$ /SiO $_2$ sorbent. Sorption isotherms are efficiently fitted by the Langmuir and Sips equations: the maximum sorption capacity reaches around 1.38 mmol Sr g $^{-1}$. Kinetic profiles for sorption are modeled using the pseudo-first order rate equation; equilibrium is achieved within 40–60 min. Complete desorption of Sr(II) using 1 M HCl solutions is even faster; equilibrium time being

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15–20 min. The sorbent can be recycled for 5 cycles with a limited decrease in the sorption performance, and full desorption. In fixed-bed column, the breakthrough curves are fitted by the Thomas equation. In multi-component solutions, the sorbent shows a preferred selectivity according the sequence: K(I) > Ba(II) > Na(I) > Ca(II). Tested on seawater, the sorbent shows high distribution ratio for Sr(II), B(III) and Mo(VI)

1. Introduction

The removal of strontium from water bodies has recently retained increasing attention from scientific community due to Fukushima Daiichi NPP accident [1,2]. The dissemination of radioactive strontium may cause serious health and life hazards due to accumulation in soils [3] and bioaccumulation in the food chain [4]. Apart this accidental dispersion of hazardous ⁹⁰Sr, the metal is used in fireworks, optical materials including TV tube monitor, ferrite magnets, ceramics and the manufacturing of some toothpastes [5]. Majors threats for human health concern direct radiotoxicity (after nuclear accident), bones diseases (associated with Ca replacement with Sr) causing different types of cancer, anemia, and leukemia [5].

Precipitation coupled with microfiltration has been investigated for the recovery of Sr(II) [6,7]. Solvent extraction was also used [8,9]. However, sorption processes have retained much more attention, playing with a wide diversity of supports. Carbon-based sorbents have been investigated for Sr(II) uptake from surface waters [10–13]. Biosorbents and biopolymers showed also promising properties for strontium uptake. [14-18]. Zeolite, silica and clays have been used under their raw form [19-22], after chemical modification [23-27], magnetite incorporation [5,28], or extractant impregnation [29]. Sorbents impregnated with Prussian Blue and analogues have also been successfully tested for Sr(II) removal [30,31]. Metal and bi-metallic nanoparticles have been applied to the recovery of cesium and strontium from simulated effluents contaminated with radioelements [32,33]. Functionalized polymer resins were widely investigated for Sr(II) recovery [34-36].

Composite organic/inorganic resins have shown promising perspectives combining the mechanical stability of silica core (for example) and highly reactive and selective functionalities of the polymer layer (which can be tailored for specific metal target) [27]. In the case of Sr(II) sorption, the coating of silica core may be based on the grafting of polymers bearing carboxylic-acid moieties [27] (maximum sorption capacity, q_m: increased with temperature up to 92 mg Sr g⁻¹). Indeed, carboxylic acid groups proved being efficient in enhancing the sorption properties of different supports [37] (q_m: 58.5 mg Sr g⁻¹), including directly on carboxylate groups of alginate beads [38] (q_m: 110 mg Sr g⁻¹). Amidoxime reactive groups have also shown high affinity for Sr (II) on synthetic polymers [39], on chitosan-based supports (q_m: 153 mg Sr g^{-1} , [40]) or composite algal/PEI beads (q_m : 189 mg Sr g^{-1} , [41]). Several studies over the last decades have demonstrated the benefits of bi-functionality on metal sorption properties [42-51]. In the case of sulfonic/carboxylic resin, Younis et al. [43] reported sorption capacities as high as 175 mg Sr g⁻¹. This work contributes to the development of bi-functional sorbents for the sorption of Sr(II). Silica particles are coated with polymethacrylic acid (PMAA/SiO₂), polyacrylonitrile (PAN/SiO₂), or amidoximated polyacrylonitrile (PAO/SiO₂) (monofunctionalized silica); the sorption properties are compared with a series of bi-functionalized silica composite (with different fractions of methacrylic acid and amidoxime moieties, PAO_xMAA_{100-x}/SiO₂).

The major outcomes of this work consist of the synthesis of mechanically stable sorbents (based on silica core), with enhanced reactivity (due to the combined effects of methacrylic acid and amidoxime bifunctionality). The optimization of the ratio of reactive moieties brings new insights in the promising perspectives opened by bi-functionality.

In the first part of the work, the materials are characterized using BET and TGA analysis, elemental analysis, FTIR and XPS

spectroscopies, and titration (for the determination of pH_{PZC}). In the second part of the work, the sorption of Sr(II) is investigated through the study of pH effect, uptake kinetics, sorption isotherms, metal desorption and sorbent recycling, and sorption in multi-component solutions (sorption selectivity) in batch reactors. The effect of flow rate on Sr(II) sorption in dynamic system (fixed-bed columns) is also investigated. Finally, $PAO_{20}MAA_{80}/SiO_2$ composite sorbent is tested for the removal of Sr(II) from seawater; in an attempt to simulate the recovery of $^{90}Sr(II)$ from radioactive contaminated effluent.

2. Materials and methods

2.1. Materials

Strontium nitrate, potassium peroxydisulfate, sodium pyrosulfite, ethanol (absolute), sodium carbonate anhydrous, and salts of barium chloride dihydrate (BaCl $_2$ ·2H $_2$ O), potassium chloride (KCl), sodium chloride (NaCl) and anhydrous calcium chloride (CaCl $_2$), used for preparing multi-component solutions, were supplied by Guangdong Guanghua Sci-Tech Co., Ltd., Shantou, Guangdong, China. Stock solutions were prepared at the concentration of 1 g L $^{-1}$ using ultrapure water (UPW). Working solutions were diluted to the assigned concentration using UPW just before use. Hydroxylammonium chloride, methyl cellulose (MC) and polyvinyl alcohol (PVA) were purchased from Shanghai Test Sihewei Chemical Co., Ltd. (Shanghai, China). Silica micro-beads were supplied by Asahi Chemicals, Co Ltd. (Osaka, Japan). Acrylonitrile (AN), divinylbenzene (DVB), nitric acid and hydrochloric acid were purchased from Sinopharm Chemical Reagent Co., Ltd. (Shanghai, China).

Seawater for testing Sr(II) removal from complex solutions was collected from Fangchenggang, a prefecture-level city in the south of Guangxi Zhuang Autonomous Region, People's Republic of China, is located at the following coordinates: 21°36′N 108°18′E (see MAP AM1, in Additional Material Section).

2.2. Synthesis of sorbents

2.2.1. Mono-functionalized/SiO2 sorbent

2.2.1.1. Synthesis of polyacryl-amidoxime/SiO $_2$ sorbent. A three-necked round bottom flask, heated using a silicone oil bath, was equipped with a condenser and a mechanical stirrer, while the third necked was connected to a dropping funnel. For reducing synthesis costs, experimental conditions were adapted: water was used as the solvent in the polymerization reaction, while $K_2S_2O_8$ and $Na_2S_2O_5$ were selected as redox initiator.

2.2.1.1.1. Synthesis of polyacrylonitrile/SiO $_2$ sorbent – PAN/SiO $_2$ (Scheme 1a). The initiator solution was prepared by dissolving 0.5 g $K_2S_2O_8$ and 0.2 g $Na_2S_2O_5$ in 10 mL UPW, under strong stirring, for 20 min at $22\pm 2\,^{\circ}C$. Ten grams of porous silica particles were immersed in the mixture of 50 mL PVA solution (1% w/w) and 6 mL MC solution (0.5% w/w), under strong stirring (250 \pm 5 rpm). In a second step, the suspension was drop wisely added to the solution containing both the crosslinker and the mixture of monomers (i.e., 1.4 g AN + 0.23 g of DVB). The reaction was maintained for 20 min at 40 °C, before increasing the temperature to 70 °C for 2 h; the final step took place at 90 °C for 7 h. The final product, polyacrylonitrile/SiO $_2$ (PAN/SiO $_2$), was obtained after washing with hot water, acetone, and dried at 50 °C under vacuum for 12 h.

2.2.1.1.2. Synthesis of polyacryl-amidoxime/SiO₂ sorbent – PAO/SiO₂

Scheme 1a. Synthesis route for the preparation of PAN/SiO₂.

(Scheme 1b). Four grams of PAN/SiO₂ were dispersed into 40 mL of water/ethanol mixture (1:1, v/v). Fifteen g of hydroxylammonium chloride were dissolved in 100 mL of water/ethanol solution (5:1, v/v) containing 7 g of Na₂CO₃. This solution was mixed with PAN/SiO₂ suspension for 20 min at 40 °C, before heating under reflux at 70 °C for 5 h. The final product (PAO/SiO₂) was dried under vacuum for 12 h.

2.2.1.1.3. Synthesis of polymethacrylic acid/ SiO_2 sorbent – PMAA/ SiO_2 (Scheme 2). The synthesis followed exactly the same procedure as described above substituting acrylonitrile with methacrylic acid. The same ratios of monomers to additives were respected. Scheme 2 reports the synthesis route (for polymethacrylic acid (PMAA) functionalization).

2.2.2. Bi-functionalized/SiO₂ sorbent

2.2.2.0.1. Synthesis of polyacrylonitrile/methacrylic acid sorbent with different ratios (Scheme 3). The bi-functionalization of SiO2 was processed first by simultaneously grafting PAN and PMAA moieties. Ten g of silica micro-beads were mixed with 50 mL of PVA solution (1%, w/w) completed with MC solution (6 mL, 0.5% w/w), followed by dropwise addition of $0.5\,g$ $K_2S_2O_8$ and $0.2\,g$ $Na_2S_2O_5$ (previously dissolved in 10 mL of water). The mixture was stirred (250 \pm 5 rpm) at room temperature (i.e., 22 ± 2 °C) for 20 min. The grafting of additional reactive groups was performed by dropwise addition of variable amounts of AN (i.e., 0.3 g, 0.7 g and 1.2 g) and MAA (i.e., 1.2 g, 0.7 g and 0.3 g) in the presence of 0.25 g of DVB. The reaction was thermally processed in three steps: at 40 °C for 20 min, followed by 2 h at 70 °C and finished at 90 °C for 7 h. The copolymer ratios varied to produce: PAN_{20}/MAA_{80} , PAN_{50}/MAA_{50} , and AN_{80}/MAA_{20} respectively. The products (PAN_xMAA_{100-x}/SiO₂) were washed with hot water, acetone, before being dried at 50 °C under vacuum for 12 h.

2.2.2.1. Synthesis of polyacryl-amidoxime/methacrylic acid sorbent with different ratios. The three samples of PAN_xMAA_{100-x}/SiO_2 were amidoximated by reaction with hydroxyl ammonium hydrochloride at 70 °C for 5 h, in the presence of sodium carbonate. Four g of polymer/SiO_2 composite were dispersed into a 1:1 water/ethanol solution (40 mL) before adding NH_3OH-Cl/Na_2CO_3 (dissolved in 5:1 water/ethanol solution) with the following mass ratios 12:4.8 g, 7.5:3 g; and 3:1.2 g for $PAN_{20}/MAA_{80},\ PAN_{50}/MAA_{50},\ and\ PAN_{80}/MAA_{20},$ respectively. The reaction was initiated at 40 °C, for 20 min. In a

Scheme 2. Synthesis route for the preparation of PMAA/SiO₂.

$$\begin{array}{c} (80, 50, 20) \\ \text{CH}_2\text{--}\text{C} \\ \text{CH}_3 \\ \text{CH}_2\text{--}\text{CHCN} \\ (20, 50, 80) \end{array} \\ \begin{array}{c} K_2S_2O_8/Na_2S_2O_5 \\ \hline \\ Na_2S_2O_8/Na_2S_2O_5 \\ \hline \\ OC/2h \\ \hline \\$$

 $\begin{array}{l} n_1 \!\!=\!\! 20(0.9g)/n_2 \!\!=\!\! 80(3.6g); \; n_1 \!\!=\!\! 50(2.1g)/n_2 \!\!=\!\! 50(2.1g); \; n_1 \!\!=\!\! 80(3.6g)/n_2 \!\!=\!\! 20(0.9g) \; for \\ PAN_{20}/MAA_{80}, \; PAN_{50}/MAA_{50}, \; and \; PAN_{80}/MAA_{20}, \; respectively. \end{array}$

Scheme 3. Synthesis route for the preparation of PAN_xMAA_{100-x}/SiO₂.

second step, the temperature was raised to 70 °C (under reflux) and the reaction lasted for 5 h. Sorbents were filtered off and washed several times with hot water and ethanol and finally dried under vacuum at 50 °C for 12 h. Scheme 4 summarizes the synthesis route (including the generic formula of PAO_xMAA_{100-x} sorbent).

2.3. Characterization of materials

The morphology of sorbent particles was observed using a Phenom ProX scanning electron microscope (SEM, Thermo Fisher Scientific, Netherlands). Semi quantitative EDX analysis (integrated in the SEM equipment) was used for characterizing the sorbents before and after metal sorption. Textural investigation was performed using Micromeritics TriStar II (Norcross, GA, USA); the samples were degassed for 13 h at 100 °C. The specific surface area was determined using BJH methods for BET equation (on both sorption and desorption branches of nitrogen isotherm).

Chemical characterization was investigated using FTIR spectrometry (4000–400 $\rm cm^{-1}$) with an IRTracer-100 (Shimadzu, Tokyo, Japan). Dried samples were grinded and mixed with KBr and conditioned as KBr disc, prior to FTIR analysis. XPS analysis was performed using an ESCALAB 250XI+ instrument (Thermo Fischer Scientific, Inc., Waltham, MA, USA). Elemental analysis was carried out using a Vario EL cube element analyzer (Elementar Analysensysteme GmbH, Langenselbold, Germany) for quantification of C, H, and N elements in the sorbents. Thermal analysis of the sorbents (PAN $_{20}/MAA_{80}$ and PAO $_{20}/MAA_{80})/SiO_2$ were performed using a Netzsch STA 449 F3

Scheme 1b. Synthesis route for the preparation of PAO/SiO₂.

$$\begin{array}{c} OH \\ COH \\ CH_{3} \\ CH_{2} \\ CH_{3} \\ CH_{3} \\ CH_{2} \\ CH_{3} \\ CH_{2} \\ CH_{3} \\ CH_{3} \\ CH_{4} \\ CH_{2} \\ CH_{3} \\ CH_{5} \\ CH_{2} \\ CH_{5} \\ CH_$$

Scheme 4. Synthesis route for the preparation of PAO_xMAA_{100-x}/SiO₂.

Jupiter (NETZSCH-Gerätebau GmbH, Selb, Germany) (temperature ramp: 10 °C/min) under nitrogen atmosphere.

The pH-drift method was used for determining the $pH_{\rm PZC}$ value [52]. A fixed amount of sorbent (i.e., 100 mg) was dispersed under agitation for 48 h in a series of flasks containing 50 mL of 0.1 M or 1 M NaCl solutions with initial pH (pH_0) adjusted to 1–14. The final pH (pH_{eq}) was measured using a Mettler Toledo pH-meter (Colombus, USA) and the $pH_{\rm PZC}$ was determined by the value corresponding to: $pH_0=pH_{eq}.$

2.4. Sorption studies

Most of the experiments were carried out in batch process. A fixed amount of sorbent (m, g) was mixed with a given volume of solution (V, L), containing strontium (C_0 , mmol Sr L $^{-1}$) for 48 h. The sorbent dosage (SD, g L $^{-1}$) was defined as SD = m/V. The standard temperature was $22 \pm 2\,^{\circ}$ C and the agitation speed was 170 rpm. The samples were collected at the end of the experiments, before being filtrated on 1.2 µm pore size filter membranes. The residual concentration in the solution (C_{eq} , mmol L $^{-1}$) was analyzed using ICP-AES (inductively coupled plasma atomic emission spectrometer, ICPS-7510 Shimadzu, Tokyo, Japan). For kinetic study, samples were collected at fixed contact times, and analyzed (after filtration) for plotting the concentration decay curves. The sorption capacity (q_{eq} , mmol Sr g $^{-1}$) was determined by the mass balance equation:

$$q_{eq} = \frac{(C_0 - C_{eq}) \times V}{m} \tag{1}$$

For sorption isotherms, the sorbent dosage was set to $0.4\,\mathrm{g\,L^{-1}}$, the initial concentration was varied between 10 and 600 mg Sr L $^{-1}$ while the initial pH of the solution was fixed at pH 8.0. The suspension was stirred for 48 h. The pH was not controlled during the sorption but the final pH value was monitored together with the residual metal concentration. The experiments have been duplicated and the comparison of the duplicates appears as Run #1 and Run #2 in the Figures. The experimental conditions have been systematically checked regarding metal precipitation and sorption artifacts on experimental setup.

For desorption studies, the sorbents collected during the sorption kinetics were mixed with 0.5 M HCl solutions and samples were regularly collected and analyzed for evaluating the released Sr(II) and for plotting the desorption kinetics. For the study of sorbent recycling, the sorption and desorption procedures were used and a rinsing step was processed using demineralized water between each sorption and desorption step. The efficiency of sorption and desorption were compared for five successive cycles.

The presence of other metals may interfere with the removal of Sr (II). In order to evaluate the impact of alkali and alkali-earth elements (abundant in seawater and/or having similar physicochemical behavior in solution) on Sr(II) uptake, a series of experiments were performed with equimolar multi-component solutions (i.e., $1.3 \, \text{mmol L}^{-1}$) at different pH values (in the range 3–8). The sorbent dosage was set at $0.125 \, \text{g L}^{-1}$; the suspension was maintained under agitation for 48 h.

After pH measurement and filtration, the residual concentrations were analyzed by ICP-AES and the sorption capacities were calculated by the mass-balance equation. The distribution ratio (L $\rm g^{-1}$) was calculated by:

$$D = \frac{q_{eq}}{C_{eq}} \tag{2}$$

And the selectivity coefficient (SC_{Me1/Me2}) was deduced by:

$$SC_{Me1/Me2} = \frac{D_{Me1}}{D_{Me2}} = \frac{q_{eq,Me1} \times C_{eq,Me2}}{C_{eq,Me1} \times q_{eq,Me2}}$$
(3)

Sorption tests were also performed in fixed-bed columns. The sorbent was immobilized in a column (diameter: 5 mm; height: 47 mm). The ratio column diameter/particle size being much higher than 12, the wall effect can be neglected. The column was fed up flow (at flow rates: 0.2, 0.4 and 1 mL min $^{-1}$, corresponding to superficial flow velocities of 0.61, 1.22 and 3.06 m h $^{-1}$); 10 mL-samples were collected in series for analysis of outlet concentrations.

The models used for fitting uptake kinetics (i.e., pseudo-first order and pseudo-second order rate equations, PFORE and PSORE; and the Crank equation for resistance to intraparticle diffusion) and sorption isotherms (i.e., Langmuir, Freundlich and Sips equations) are summarized in Tables AM1 and AM2. Non-linear regression analysis was used for the determination of model parameters, while the determination coefficient was determined comparing experimental and simulated data (linear regression analysis). The Akeike information criterion was also applied for comparing the fits of experimental data with selected equations [53]. For the modeling of desorption kinetics (PFORE and PSORE, [54]) and breakthrough curves (Thomas equation [55]), the model equations are reported in the footnotes of relevant tables (see below).

2.5. Sr(II) recovery from seawater

Sorption tests on seawater samples were also performed in batch, using the same experimental procedure as with synthetic solutions: 500 mL of water (which pH was controlled to pH 4, 8 and 12, using either HCl or NaOH solutions) were mixed with 0.1 g of sorbent for 48 h (samples were collected at intermediary times for plotting kinetic curves). Semi-quantitative EDX analysis on loaded samples was used for determining the metals to investigate (i.e., Na(I), K(I), Mg(II), Ca(II), Sr (II), B(III), Ru(III) and Mo(VI)).

3. Results and discussion

3.1. Characterization of sorbents

In the preliminary part of the work, different formulations of bifunctionalized sorbents were compared to the sorption properties of mono-functionalized supports for selecting the best compromise (regarding Sr(II) uptake); results are summarized in the Section 3.2.1. Therefore, the characterization of the material was mainly focused on this specific "optimized" sorbent (i.e., PAO₂₀MMA₈₀/SiO₂).

3.1.1. Morphology and textural properties

Sp herical silica-based resins have a particle size in the range 119–132 µm (Fig. AM1). The surface is coated by a polymer layer that gives to the sorbent an irregular surface topography. Fig. AM2 shows the BET analysis of raw SiO2, PMAA/SiO2 (mono-functionalized) and PAO₂₀MAA₈₀/SiO₂ (bi-functionalized) beads. While the chemical grafting of silica microparticles usually contributes to strongly decrease the specific surface area (SSA), the pore size and the pore volume [56], in the case of mono- and bi-functionalized silica beads, the textural properties shows contradictory trends. The SSA hardly changes with the successive modifications: f rom 69.1 m 2 g $^{-1}$ (SiO₂) t o 62.4 m 2 g $^{-1}$ (PMAA/SiO₂) and 64.7 m² g⁻¹ (bi-functionalized sorbent). Surprisingly, the post-modification (amidoxime grafting) slightly improves the SSA of the sorbent. However, the differences are not marked. The average pore size is of the same order of magnitude for the functionalized materials (around 300 Å) and little higher than the pore size of raw silica beads (around 230 Å). The raw support can be considered a mesoporous silica based on the average size of the pores (Fig. AM3). The coating of the silica beads with the polymer layers contribute to fill in the smallest p ores meaning that the average (global) p ore size slightly increases. Unexpectedly, the pore volume also increases with the p rogressive modification of t he s up p ortf rom 0.299 c m^3 g $^{-1}$, to 0.376 cm³ g⁻¹ for methacrylic acid-grafted silica, and up to 0.399 cm³ g⁻¹ for the bi-functionalized material.

The average pore size largely exceeds the ionic radius of hydrated Sr (II) species (i.e., $1.26\,\text{Å}$ [57]), meaning that the diffusion of strontium ions will not be strongly hindered by the p orous p rop erties of the composites.

3.1.2. Elemental analysis

Table 1 reports the elemental analysis of different sorbents (homo- and heteropolymers supported on SiO₂: PAN, PMAA, PAO, at different ratios of substitution). The residue (to 100%, w/w) is probably associated with SiO₂ core (the residue being Si element): the average value is 31.26% (with a standard deviation close to 0.85%), representing about 53.52 \pm 1.45% (in weight fraction).

The comparison of the N content for PAO vs. PAN derivatives clearly demonstrates that the amidoximation of nitrile groups is quantitative (the N content is roughly doubled, consistently with expected reaction (Scheme 1b). The amidoximation is also quantitative for bi-functionalized materials: the N content (mmol N g $^{-1}$) is doubled while converting PAN $_x$ MMA $_{100-x}$ /SiO $_2$ into PAO $_x$ MMA $_{100-x}$ /SiO $_2$.

Table AM3 (a-d) shows the semi-quantitative EDX analysis of the surface of composite materials at different stages of the synthesis of sorbents (also including after Sr(II) sorption). Obviously, these semiquantitative analyses must be considered as an indication or a confirmation of the chemical modifications of the materials along the synthesis procedure for the different compounds. In the SEM analysis, all materials keep the spherical properties of the silica particles with surface coating due to grafting of the hydrocarbon chains. For monofunctional grafted composites, the trends are consistent with suggested synthesis routes. The amidoximation substantially increases the N atomic fraction (NAF, almost doubled when converting PAN/SiO2 to PAO/SiO₂: from 8.06 to 15.37%). The rate of substitution is close to 95%. This is associated with a relative increase of O atomic fraction due to the conversion of CN into C(=NOH)NH2. On the other side, for PMAA/SiO₂, the grafting of acrylic acid allows drastically increasing O content (up to 55.52%); this is directly associated with the potential grafting of two oxygen atoms on carboxylic groups per monomer unit deposited on SiO₂ micro-particles. For bi-functional sorbents, using N_{AF} as the marker of substitution allows characterizing the quasi-linear increase in substitution rate (AO_{Subst,Ratio}) for PAN_xMAA_{100-x}/SiO₂ sorbents $(N_{AF} = 0.07 \times AO_{Subst.Ratio} + 1.49)$ and PAO_xMAA_{100-x}/SiO_2 sorbents ($N_{AF} = 0.11 \times AO_{Subst.Ratio} R + 2.83$). It is noteworthy that

the comparison of NAF for nitrilated intermediary materials and amidoximated sorbents demonstrates that the amidoximation rate varies between 83% and 91% for bi-functional materials. As expected, when the rate of amidoximation increases the atomic fraction of O decreases (from 50.5 and 52.9% to 30.8 and 40.2% for PAN_xMAA_{100-x}/SiO_2 and PAO_xMAA_{100-x}/SiO₂, respectively). The conversion of nitrilated monofunctional composite into amidoximated resin is almost quantitative (1.91 times the content of N in the PAN/SiO2 material, close to the theoretical factor; i.e., 2). In the case of bi-functional composites, the comparison of PAN_xMAA_{100-x}/SiO_2 with PAO_xMAA_{100-x}/SiO_2 shows a lower degree of substitution (NAF increases after amidoximation by a ratio lower than 1.81; (i.e., between 1.61 and 1.81)). The presence of metacrylic acid moieties limits the efficiency of the amidoximation step. This can be explained by two main reasons: (a) the limitations due to steric hindrance, and/or (b) the inhibiting effect associated with the presence of polar carboxylic groups.

3.1.3. Thermal degradation

Fig. AM4 shows the thermogravimetric analysis of PAN₂₀MAA₈₀/ SiO₂ and its amidoximated derivative (i.e., PAO₂₀MAA₈₀/SiO₂). The comparison of TGA profiles shows relatively close behaviors. Below 269.1-266.5 °C, superficial water is released from the materials with a weight loss limited to 1.41% and 2.78% for PAN20MAA80/SiO2 and PAO₂₀MAA₈₀/SiO₂, respectively. A second weight loss step (representing 5.55% and 8.26% for the nitrile and amidoximated materials, respectively) is observed between 269.1-266.5 °C and 564.9-493.7 °C. This weight loss may be associated with the degradation of polyacrylonitrile [58,59], amidoxime substituents [60], poly methacrylic acid compound [61], and to the depolymerization of the silica coating. The amidoximation decreases the stability of the composite (highest weight loss and lower temperature of degradation) [60]. Actually, this second step combines different degradation mechanisms associated with the diversity of polymer and functional groups that coat silica particles: several weak shoulders/waves are observed, especially on the TGA profile of nitrile-composite (this is less apparent for amidoximated material). The third step in the thermal degradation corresponds to the degradation of the char, above 493.7 °C for PAO₂₀MAA₈₀/ SiO₂ and above 564.86 °C for PAN₂₀MAA₈₀/SiO₂; the total weight losses correspond to 13.63% and 7.63%, respectively. This means that the polymer layer coating silica beads is relatively limited: less than 8% in the case of the intermediary compound (nitrile-composite) and less than 14% for the amidoximated bi-functionalized composite.

Fig. AM5 compares the DTG (derivative thermogravimetry) profiles for the thermal degradation of the two composites. The nitrile-composite shows a smoothed (poorly resolved) curve with a very broad exothermic band (in the range 100–400 °C) without any specific peak. On the opposite hand, after amidoximation the DTG curve shows a little shorter range of temperature (in the range 100–394 °C) for the poorly resolved exothermic band, which is completed by twin peaks at 265.14 °C and 272.55 °C; another small exothermic peak is also detected at 586.48 °C. The cyclization of nitrile was reported at 584 °C [62]. In the case of silica/PAN hollow nanoparticles and silica/poly

Table 1 Elemental analysis of materials (C, N, H and O elements, wt% and mmol N g^{-1}).

Materials	C (%)	N (%)	N (mmol g ⁻¹)	H (%)	O (%)
PAN	21.4	6.9	4.93	3.6	36.28
PAO	22.02	14.18	10.12	4.1	29.2
PMAA	21.8	0.2	0.14	4.62	40.08
PAN ₂₀ MAA ₈₀ /SiO ₂	17.6	2.18	1.56	3.71	46.31
PAN ₅₀ MAA ₅₀ /SiO ₂	15.2	4.57	3.26	3.8	45.33
$\mathrm{PAN}_{80}\mathrm{MAA}_{20}/\mathrm{SiO}_2$	13.2	5.65	4.03	3.86	45.89
PAO ₂₀ MAA ₈₀ /SiO ₂	17.9	4.71	3.36	5.01	41.18
PAO ₅₀ MAA ₅₀ /SiO ₂	15.8	9.82	7.01	5.13	38.35
PAO ₈₀ MAA ₂₀ /SiO ₂	14.3	10.93	7.80	5.21	38.66

acryclic acid hollow nanogels, the maximum exotherm peak was observed close to 440 °C for the two composites (indifferently of the grafted polymer) [63].

3.1.4. FTIR analysis

FTIR spectroscopy was used for characterizing SiO_2 , PAN/SiO_2 (Fig. AM6a), $PMAA/SiO_2$ (Fig. AM6b), PAN_xMAA_{100-x}/SiO_2 , and PAO_xMAA_{100-x}/SiO_2 (at different c op olymerr atios (Fig. AM7a,b,c)). The spectra were also characterized for $PAO_{20}MAA_{80}/SiO_2$ after Sr(II) sorption, after metal desorption and after five cycles of sorption desorption. Tables AM4–AM7 report the main characteristics peaks (wavenumbers) and their assignments.

Raw silica shows peaks at $1110\,\mathrm{cm}^{-1}$, $806\,\mathrm{cm}^{-1}$, and $474\,\mathrm{cm}^{-1}$, which are assigned to asymmetric stretching and bending vibrations of Si–O–Si bonds [64–66]. The first two peaks are overlapped with C–N and C–C respectively in polymer-coated SiO₂ [64,67,68]. The broad peaks at $3442\,\mathrm{cm}^{-1}$ and $1632\,\mathrm{cm}^{-1}$ are assigned to OH groups in SiO₂, associated with water binding at the surface of the solid [65].

After acrylonitrile grafting (in both homo- and hetero-polymer), some specific peaks appear including nitrile groups 2280–2240 cm [69,70]. Carboxylate groups associated to the incorporation of MAA (both in homo- and hetero-polymers) are identified at 1750–1725 cm⁻ [70–72]. Other peaks attributed to carboxylate ester groups are also identified at 1420-1300 cm⁻¹ for O-C=O group stretching vibrations, which overlapped with C-N stretching of amide and -OH bending of amidoxime groups in copolymeric sorbents [70,73]. These peaks may overlap with those of silicate and eventually with amide groups of amidoxime [74] (occurring at the second step of the modification in the co-polymer grafting). After the conversion of nitrile to amidoxime, the peak at $2280-2240\,\mathrm{cm}^{-1}$ fully disappears, showing the effective and complete amidoximation; this is consistent with the conclusions raised from elemental analysis. In addition, the intensity of the broad peak at around 1600-1690 cm⁻¹ increases due to the grafting of the amide groups of amidoxime [75]. The peak at 1050–990 cm⁻¹ is attributed to N-O stretching band [66]. In some cases (especially for SiO2 composites), this band may overlap with the broad band of Si-O-Si and/or with C-N band that appears at 1090-1020 cm⁻¹ [66].

The sorption of Sr(II) is characterized by some shifts or broadening of bands. For example, the broad band around $3400-3600~\rm cm^{-1}$ tends to enlarge after metal binding; this is probably associated with the interaction of strontium cation with nitrogen and hydroxyl groups (direct and/or changes in their chemical environment). In the case of monofunctionalized PMAA/SiO₂, the most significant changes are observed in the range $1500-1700~\rm cm^{-1}$ with the decrease in the intensities of the peaks at $1635~\rm cm^{-1}$ and $1680~\rm cm^{-1}$, replaced by a peak at $1641~\rm cm^{-1}$.

This is directly associated with the interaction of Sr(II) with carboxylate groups. For PAO/SiO₂, the intensities of the band at $943\,\mathrm{cm}^{-1}$ (N–O stretching) and the shoulder at $569\,\mathrm{cm}^{-1}$ (assigned to –OH and C–H bending vibrations) decrease with metal binding. This may be explained by the interaction of Sr(II) with amidoxime moiety. The low sorption capacities obtained with mono-functionalized sorbents (see below) may explain the difficulty to identify large changes in the FTIR spectra. Bi-functionalized sorbents (especially for PAO₂₀MAA₈₀/SiO₂) have higher affinity for Sr(II); therefore, the spectra are also more influenced by the sorption of the metal ions. More specifically, the most representative modifications are reported at:

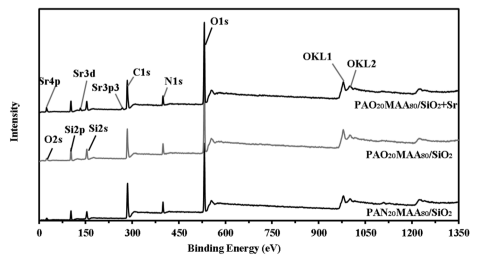
- (a) 2426 cm⁻¹ and satellite peaks around 2360 cm⁻¹: new small peaks, which could be associated with carbon dioxide at the surface of the sorbent.
- (b) $1763\,\mathrm{cm^{-1}}$ peak was shifted to $1718\,\mathrm{cm^{-1}}$; associated with carboxylic groups (C = O stretching) for $PAO_{20}MAA_{80}/SiO_2$ (not detectable for other PAO_xMAA_{100-x}/SiO_2 sorbents that have much lower acrylic acid fractions).
- (c) 1360–1430 cm⁻¹: peak widening; the sharp peak at 1383 cm⁻¹ (C-N stretching and OH bending) is replaced with a broad band of increasing widening when the fraction of amidoxime decreases; in addition, a peak at 1398 cm⁻¹ and two shoulders at 1429 cm⁻¹ and 1358 cm⁻¹) also appear after functionalization.
- (d) 825 cm⁻¹: new sharp peak appearing only for bi-functionalized materials (sometimes associated with nitrate compounds [70]).

The changes in the FTIR spectra after Sr(II) sorption confirm the simultaneous contributions of carboxylic groups (from MAA), amine and hydroxyl groups (from amidoxime moiety) in the binding of strontium.

3.1.5. XPS analysis

The XPS survey spectra for $PAN_{20}MAA_{80}/SiO_2$, $PAO_{20}MAA_{80}/SiO_2$ and $PAO_{20}MAA_{80}/SiO_2 + Sr$ are reported on Fig. 1. The XPS spectra confirm the presence of silica core (Si 2p and Si 2s), and polymer markers (C 1s, N 1s and O 1s). The sorption of Sr(II) is confirmed by the appearance of the typical peaks of Sr element (Sr 4p, Sr 3d, Sr 3p and Sr 3s).

Table AM8 shows the high resolution spectra (HRES) of the C 1s, O 1s, N 1s and Si 2p signals. Table AM9 reports the HRES of Sr 3d, Sr 4p, Sr 3p and Sr 3s for PAO $_{20}$ MAA $_{8}$ 0/SiO $_{2}$ after metal sorption. These tables summarize the assignments, atomic fractions and full-width at half maximum (FWHM) for the deconvolution of the different signals. These analyses contribute to confirm the successful chemical modification of



 $\textbf{Fig. 1. XPS} \ analysis \ (survey) \ of \ functionalized \ materials \ (PAN_{20}MMA_{80}/SiO_2, \ PAO_{20}MMA_{80}/SiO_2 = sorbent, \ sorbent \ + \ Sr(II)).$

the support and the interpretation of binding mechanisms.

The composition of the composite bi-functionalized materials offers a wide diversity of signals associated to the different bonds and chemical environments of the C, O and N elements in the different p olymer-coated sorbents. For C 1s signal, the deconvolution of the signal shows the presence of many different bonds, a ssociated with methacrylic, acrylonitrile and/or amidoxime moieties: C-C and C-H (or adventitious C) around $284.3 \pm 0.3 \,\text{eV}$; C-O or C=N at $286.2 \pm 0.4 \,\text{eV}$ [76]; O-C=O (carboxylic groups) at $287.3 \pm 0.2 \,\text{eV}$; C=N at 288.0 \pm 0.2 eV [77-80]. The composite materials (supported on SiO₂) contain MAA with either nitrile or amidoxime ending groups; this is confirmed by the disappearance of C≡N signal (at 288.13 eV) when PAN₂₀MAA₈₀/SiO₂ is amidoximated (together with the appearance of a band corresponding to C=NOH, at 286.2 eV). The sorption of Sr(II) involves a small decrease in the binding energies but also an increase in the atomic fractions of some C 1s bands, including both C (carboxyl) and C (amidoxime) groups (from 10.26 to 12.6% and from 6.13 to 17.97%, respectively).

The chemical modification of the PAN₂₀MAA₈₀/SiO₂ is also characterized by the changes observed on N 1s signal. Ma et al. [76] reported the presence of two bands for N 1s on polyacrylonitrile fibers at 398.63 eV and 400.60 eV (nitrile group, C≡N, and shake-up line, respectively). After amidoximation, the N 1s bands appeared at 398.37 eV (C=N-O, part of amidoxime) and at 398.82 eV (C-NH₂). Li et al. [81] investigated the sorp tion of uranium on MOF functionalized with amidoxime groups and reported three bands at 399.2 eV, 400.4 eV, and 401.45 eV for C-NH₂ or C=N, C-NOH and C-NH₂⁺, respectively. This is consistent with the group s identified on r educed g rap heneoxide grafted with amidoxime groups [82]. With PAO₂₀MAA₈₀/SiO₂, the two p eaks corresponding to N=C-OH and NH2 are detected at around 398.9 eV and 400.65 eV, respectively; the differences in BEs are not very marked compared with the nitrile and shake-up lines of precursor. After strontium binding, the BEs of the signals are weakly shifted by 0.15 eV toward higher BE for N=C-OH (proving the donation of lone pair of electrons from N through the coordination of Sr(II) with N [83]) and lower energy for NH2. Wang et al. [78] investigated uranyl sorption on hydrolyzed amidoximated-PAN fibers and observed that the signal corresponding to -NH2 was slightly shifted after U(VI) binding; while -NOH bond did not appear to be affected by metal binding and a new peak assigned to N-U complexation (around BE = 404.68 eV) appeared. Here also, consistently with Wang et al. [78], a new weak band appears at 405.3 eV, which is associated with the direct interaction of the metal with N (from amino groups).

For O 1s, Wang et al. [78] identified three peaks, on amidoximated-PAN fibers, at 531.18 eV, 530.34 eV and 529.54 eV corresponding to -OH, -NO and -CO groups, respectively. For amidoximated -MOF, Li et al. [81] showed substantially higher binding energies for -OH, C=O and C-O groups at 531.4 eV, 532.1 eV and 532.9 eV, respectively. Chen et al. [82] synthesized amidoxime-functionalized reduced graphene oxide and they assigned the bands at 531.48 eV, 532.60 eV and 533.70 eV to $-COO^-$, >C=O and -C-OH groups, respectively. In the composite sorbents, the presence of SiO2 core introduces a complementary band at around 532.1 \pm 0.2 eV. For PAN₂₀MAA₈₀/SiO₂, O element only appears in the silica compartment (532.27 eV; atomic fraction, AF, close to 95%) and in the methacrylic acid (530.65 eV): however, traces of O-Na bond are also identified at 529.04 eV (residue from synthesis procedure). After amidoximation, the fraction of O associated with SiO₂ core decreases due to the appearance of -C(NH₂)= N-OH moieties (from 95 to 92%), not compensated by the fraction of C=NOH that appeared, but by the increasing proportion of C=O. This may be explained by the conversion of some nitrile groups to carboxylic acid groups by the effect of temperature (and activation technique) and reagents (pH, alkaline conditions) [84]. The sorption of Sr(II) is followed by the appearance of a new peak at BE = 528.2 eV (weak signal) and the relative increase in the AF of -C=O. This means that Sr(II) binding may involve some carboxylic groups as mono-dentate ligand and/or combined with amine groups to form multi-dentate complex.

The sorption of Sr(II) is confirmed on the survey spectrum by signals at binding energies close to 18.42 eV (Sr 4p), 134 eV (Sr 3d) and 277 eV (Sr 3p). The deconvolution of Sr 3d HRES spectrum shows the presence of 2 doublet-peaks (spin-orbital components, corresponding to the Sr $3d_{5/2}$ and Sr $3d_{3/2}$ spin-orbital splits) for two Sr chemical environments. The Sr-O bond is the most intense (largest atomic fraction), which is associated with Sr $3d_{5/2}$ at $132.48\,\text{eV}$ and Sr $3d_{3/2}$ at $134.09\,\text{eV}$. Gregurec et al. [85] reported the Sr $3d_{5/2}$ at BE close to 133.5 eV in the case of strontium binding by carboxylic groups of poly(acrylic) brushes. Similarly, Wang et al. [27] confirmed the binding of Sr(II) on carboxylic groups of methylacrylic acid/silica composite by the presence of two peaks at 133.55 eV (Sr $3d_{5/2}$) and 135.09 eV (Sr $3d_{3/2}$). In the case of algal/PEI beads functionalized with amidoxime groups, Wei et al. [41] reported Sr(II) sorption on amidoxime reactive groups with Sr 3d_{5/2} and Sr 3d_{3/2} bands at lower BEs (i.e., 132.91 eV and 134.59 eV, respectively). The coexistence of the four peaks (with very low AF for the peaks at 133.87 and 135.55 eV) means that amidoxime groups constitute the most active binding groups although carboxylate groups contribute marginally for strontium chelation or stabilization. Other

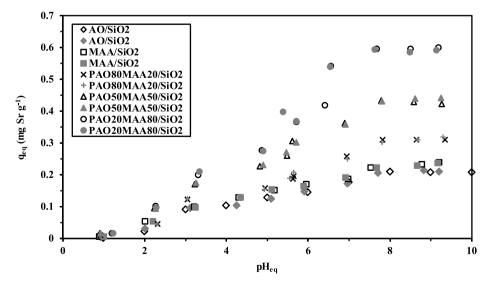


Fig. 2. Effect of pH on Sr(II) sorption using mono-functional and bi-functional polymers coated silica microbeads (Sorbent dosage, SD: 0.4 g L^{-1} ; C_0 : 50 mg $L^{-1} = 0.59 \text{ mmol Sr L}^{-1}$; agitation time: 48 h; agitation speed, 170 rpm; T: 22 ± 2 °C; duplicated experiments: Run #1:opened symbols, Run #2: closed symbols).

signals for Sr are observed: Sr 4p (at 18.42 eV), Sr 3s (poorly resolved peak, at 355.69 eV), and Sr 3p (two different spin-orbital splits: Sr $3p_{3/2}$ at 268.3 eV, and Sr $3p_{1/2}$ at 278.65 eV). These observations confirm the multi-binding groups of Sr(II) onto the sorbent through two different types of bonds involving probably amine and carboxylate groups from amidoxime and acrylic moieties, respectively, and hydroxyl groups from amidoxime and/or acrylic moieties.

3.1.6. Titration – pH_{PZC}

The pH_{PZC} of the optimized bi-functionalized sorbent is determined by the pH-drift method (Fig. AM8). The average value is close to 6.47 (\pm 0.02), showing the limiting effect of the concentration of the background salt (0.1 M vs. 1 M NaCl solution). The co-existence of methacrylic and amidoxime groups brings reactive groups with specific proton exchange properties. For example, the pKa values of amidoxime-based compounds strongly depend on the substituents but ranges between 3 and 5.3 in most cases [86]. The p Ka of methacrylic acid is rep orted close to 4.65 [87]; the p olymerization and the chemical modification of polymethacrylic chain strongly modulate the acid-base properties of functionalized material [88].

Selected functionalized sorbent (i.e., $PAO_{20}MMA_{80}/SiO_2$) is protonated in acidic solutions; meaning that the sorbent can electrostatically attract anionic species at pH lower than 6, and repulse cationic species in acidic conditions (pH lower than 3–4). This protonation also opens possibilities for ion-exchange mechanisms at intermediary acidic pH values (moderate pH around pH 5) and metal complexation at neutral pH.

3.2. Sorption studies

3.2.1. Effect of pH and sorbent selection

Fig. 2 reports the comparison of Sr(II) sorption capacities for monoand bi-functionalized silica composites (including variable levels of amidoxime grafting). The experimental series were duplicated and the overlapping of the curves clearly demonstrates the reproducibility of sorption performances. At pH 1, the sorption properties for strontium are negligible for the different materials. The strong protonation of reactive groups limits the capacity of the sorbent to bind strontium, which is present as Sr²⁺ (about 84.7%) and SrNO₃⁺ (15.3%). The acid used for pH control leads to the presence of nitrate anions that may bind to protonated reactive groups; the negligible sorption capacity means that Sr(II) could not be sorbed as ion pair at the surface of the sorbent. When the pH increases, the sorption capacity linearly increases up to pH 7.5-8. The competition of protons decreases and the support progressively deprotonates making possible the binding of strontium cations. Sr(II) is present as free strontium species at pH above 4 (with progressive decrease of the fraction of strontium nitrate cation). Above pH8, the sorption stabilizes; similar observations were reported by Asgari et al. [37] for Sr(II) sorption on MOF. It is noteworthy that the pH-profiles overlap for mono-functional silica beads (both AO and MAA modified beads, with sorption capacity below 0.24 mmol Sr g $^{-1}$ (under selected experimental conditions). The substitution of AO on MAA/SiO₂ backbone (i.e., bi-functional sorbents) increases the sorption of Sr(II) (compared with mono-functional beads). This increase depends on the yield of substitution; actually, highest sorption is obtained at the lowest (tested) rate of substitution (i.e., 20/80 ratio between AO and MAA moieties). The maximum sorp tion cap acities stabilize around $0.31 \text{ mmol Sr g}^{-1}$ for $PAO_{80}MAA_{20}/SiO_2$, $0.44 \text{ mmol Sr g}^{-1}$ for $\rm PAO_{50}MAA_{50}/SiO_2,$ and up to 0.59 mmol Sr $\rm g^{-1}$ for $\rm PAO_{20}MAA_{80}/SiO_2.$ Fig. AM9 shows the correlation between the rate of AO substitution and the maximum sorption capacity (obtained on the stabilization plateau reached above pH 7.5).

Table AM3 (a-d) compares the semi-quantitative EDX surface analysis of the different sorbents after Sr(II) sorption. This data confirm the observations from Fig. 2: the sorption of Sr(II) decreases with the excessive amidoximation of intermediary PAN_xMAA_{100-x}/SiO_2 composite.

However, compared with PMAA/SiO₂, the presence of amidoxime groups increases the binding of Sr(II): Sr_{AF} increases from 1.59 (for PMAA/SiO₂) to 2.19 (for PAO₂₀MAA₈₀/SiO₂), while a larger substitution rate decreases metal uptake. These observations are consistent with Fig. AM9.

Figure AM10 shows the \log_{10} -plots of the distribution ratio (D = q_{eq}/C_{eq} , L g^{-1}) vs pH_{eq}. The insert focuses on the linear section of these curves (registered between pH 2 and 8). The slopes vary between 0.109 and 0.176. In ion-exchange reactions, the slope is frequently associated with the stoichiometric ratio for proton exchange at the surface of the sorbent and metal ions. In present cases, the slope analysis cannot be correlated to appropriate ion-exchange ratio.

The pH frequently changes during the sorption process and it is important correlating sorption properties with these pH variations. In the case of mono- and bi-functionalized silica sorbents, the pH variation is relatively limited (by less than 0.5 pH unit) (Fig. AM11): the pH slightly increases at pH 4 and 5 and tends to slightly decreases at pH above 6. This is consistent with the study of pH $_{\rm PZC}$ (around 6.5, for PAO $_{\rm 20}$ MAA $_{\rm 80}$ /SiO $_{\rm 2}$), although the pH variations are much less marked in the presence of strontium.

Based on these results, further experiments are performed at pH₀ 8 (corresponding to pH_{eq} around 7.4-7.6). The optimum sorption being obtained with the lowest rate of AO substitution, further experiments are also carried out with this specific optimized sorbent (i.e., PAO₂₀MAA₈₀/SiO₂). Alexandratos et al. [46] discussed the interest of bi-functionalization of polyacrylonitrile fibers (using amidoxime and amine groups) for improving the sorption of U(VI). They recognized several effects for this double functionalization: (a) the amine groups (preferentially diethylenetriamine, DETA) introduce additional sorption sites for U(VI) binding, and (b) the interaction of DETA with AO (though hydrogen bonding) contributes to improve the dissociation of the acidic N-O-H group. In the case of bi-functional (carbonyl and phosphoryl groups) resins, Alexandratos and Smith reported the counter-current effects of the bi-functionalization through intraligand cooperation and intraligand hydrogen bonding mechanisms. In the case of uranyl sorption (as carbonate anionic species), the synergistic effect of positive charge and amidoxime coordination have been demonstrated to explain the high efficiency of bi-functional resins [47]. Piechowicz et al. [39] explained the remarkable sorption of uranyl from seawater onto a bi-functional polymer by the simultaneous weak coordination of uranyl with chelating carboxylates and monodentate binding by amidoxime groups. Bi-functionalization (amidoxime and sulfur compounds) was also discussed by Qu et al. [48] for improving the selective removal of Hg(II) from multi-component solutions. The literature is much less abundant concerning the enhancement of Sr(II) sorption properties using bi-functionalized sorbents. El-Naggar et al. [89] described the enhancement of sorption properties for divalent cations (including Sr(II)) while modifying a resin by simultaneously grafting different reactive groups (P-based ligands). The multi-group cooperation, called supported ligand synergistic interaction, depends on the target metal and the acidity of the solution. Recently, Younis et al. [43] reported a substantial increase in Sr(II) (and Ca(II)) sorption by simultaneous grafting of sulfonic and carboxylic groups on a polymer backbone (styrene butadiene/acrylonitrile resin) for metal recovery from highly saline oilfield wastewater.

3.2.2. Uptake kinetics

The uptake kinetics can be controlled by the resistance to bulk diffusion, to film diffusion and to intraparticle diffusion, in addition to the proper reaction rate of sorption. Providing a sufficient agitation to prevent particle settling and heterogeneous distribution of the solute in the solution and to minimize the resistance to film diffusion allows neglecting the first two diffusion criteria. For film diffusion, the resistance to mass transfer is usually limited to the first minutes of contact. Therefore, the resistance to intraparticle diffusion (RIDE, the so-called Crank equation) is more appropriate for the modeling of kinetic

profiles on a wider time range (Table AM1, see Additional Material Section). The sorption process can be modeled using, by analogy with homogeneous reactions, the pseudo-first order rate equation (PFORE) and/or the pseudo-second order rate equation (PSORE) (Table AM1). Fig. 3 compares the fits of experimental data for Sr(II) uptake at three different initial metal concentrations using PAO20MAA80/SiO2 and the three models. The solid lines corresp ond to fitted curves with the parameters summarized in Table 2. Fig. AM12 shows an example of reproducibility test for uptake kinetics. The sorption of Sr(II) is relatively fast: 25-60 min of contact are necessary for reaching equilibrium, depending on metal concentration. More than 90% of total sorption occurs within the first 25 min of contact. The small size of the sorbent (around 125 µm), the large pore size of the composites and the thin polymer layers (which controlled the diffusion thickness to be passed through) may explain the relatively fast sorption kinetics. The hydrated ionic size of strontium is close to 1.26 Å [57]; this means two orders of magnitude lower than the average pore size of the sorbent.

The superimposition of the curves with the three models shows the difficulty to di scriminate be tween the di fferent models, at least at higher metal concentrations. The curve obtained at the lowest Sr(II) concentration (i.e., C₀: 0.604 mmol Sr L⁻¹) shows a greater inaccuracy in the time range 15-30 min (corresponding to the highest curvature). In this case, the best fit was obtained with the PFORE; this is confirmed by both the determination coefficient (i.e., R²) and the value of the sorption capacity at equilibrium $(q_{eq,1}, calculated value from PFORE)$ closer to the experimental value $(q_{\rm eq,exp})$ than $q_{\rm eq,2}$ (calculated from PSORE). The PFORE is frequently associated with physical sorption while the PSORE counts on chemical interactions. The indistinct fitting trends cannot help in discriminating between the two types of interaction modes. The complementary analysis of fitting parameters with the Akaike information criterion (AIC) shows that at low metal concentration the curve is better fitted by the PFORE while at higher concentration the RIDE better simulates the kinetic profiles. The values of apparent rate coefficients (k₁ and k₂) slightly vary with metal concentration: in the range $8.9-16.7 \times 10^{-2} \text{ min}^{-1}$ for PFORE and between 19.3 and $12.2 \times 10^{-2} \text{ L mmol}^{-1}$ min⁻¹ for PSORE. At higher concentrations (i.e., 2.37 and 3.57 mmol Sr g⁻¹), the RIDE fits well the kinetic p rofiles. The effective diffusivity coefficient (Deff) varies between $6.8 \times 10^{-11} \text{ m}^2 \text{ min}^{-1}$ and $11.7 \times 10^{-11} \text{ m}^2 \text{ min}^{-1}$; These values are about three orders of magnitude lower than the self-diffusivity of Sr(II) in water (i.e., $4.75 \times 10^{-8} \text{ m}^2 \text{ min}^{-1}$, [90]). This is a confirmation that the resistance to intraparticle diffusion plays a role in the control of the up take kinetics. These values are about one order of magnitude lower than the values reported for diffusivity of Sr(II) in sorbent based on titanium phosphate (i.e., 6×10^{-10} m² min⁻¹) [91] but several orders of magnitude higher than the effective diffusivity of Sr(II) into straw-derived char $(5.4 \times 10^{-14} \text{ m}^2 \text{ min}^{-1})$ [92]. For macroporous resin, Sr(II) diffusivity was evaluated close to 1.2×10^{-8} m² min⁻¹ [93]. In the case of Dowex resins, Boyd and Soldano [94] identified the contribution of crosslinking ratio to the control of diffusion properties (in the range: $1.37-2.03 \times 10^{-9}$ m² min⁻¹); similar value was reported for Dowex 50 W-X8 resin (i.e., 1.17×10^{-9} m² min⁻¹) [95]. These different d ata c onfirm th at the mass transfer properties of the bi-functionalized SiO₂ composite are roughly comparable to those reported for conventional resins.

3.2.3. Sorption isotherms

The distribution of the solute between the liquid and solid phase for increasing concentrations is described by sorp tion isotherms. The sorption isotherms can be fitted by different models such as Langmuir equation (mechanistic model) and Freundlich or Sips equations (empirical models). Fig. 4 compares the modeling of Sr(II) sorption isotherm at pH $_0$ 8 for PAO $_{20}$ MAA $_{80}$ /SiO $_{2}$ sorbent. The superimposition of the two duplicated series confirm the reproducibility of sorption performance on selected sorbent. The general shape of the curve is characterized by a progressive increase of sorption capacity followed by an

asymptotic plateau (for equilibrium concentration above 4 mmol Sr g⁻¹). This asymptotic trend is not consistent with the power-type function represented by the Freundlich equation: this is consistent with Table 3 that summarizes the fitted parameters. The initial slope of the curve is representative of the affinity of PAO20MAA80/SiO2 for Sr(II). Actually, this initial slope is not steep; showing that appreciable sorption capacities require high metal concentration: the removal of Sr(II) traces will require high sorbent dosage for achieving complete metal recovery. The initial slope corresponds to the coefficient $q_{m,L}\times b_L$ (L g⁻¹); in the present case, this term, analogous of distribution ratio, tends to 2.14 and 2.29 L g⁻¹ for series #1 and #2, respectively. The comparison of the fits with experimental data show that the Freundlich and the Sips equations tend to overestimate the sorption capacities at the saturation of the sorbent, contrary to the Langmuir equation. The insert shows the initial section of the curve where the Langmuir equation underestimates sorption capacities while the Freundlich underestimates experimental data; the Sips equation fits better the profile

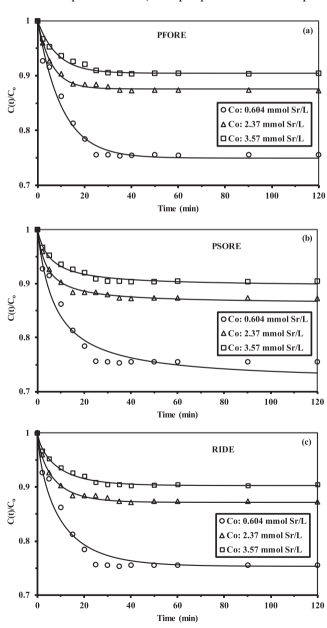


Fig. 3. Sr(II) uptake kinetics using $PAO_{20}MAA_{80}/SiO_2$ for different initial concentrations – Modeling with the PFORE(a), the PSORE (b) and the RIDE (c) (pH₀: 8; pH_{eq}: 7.59–7.51; SD: 0.25 g L⁻¹; agitation speed: 170 rpm; T: 22 ± 2 °C).

Table 2 Parameters of kinetic models for Sr(II) sorption using PAO₂₀MAA₈₀/SiO₂.

		C_0 (mmol Sr L^{-1})			
		0.6		2.37	3.57
Model	Parameter	Run #1	Run #2		
Experimental	q _{eq} (mmol U g ⁻¹)	0.591	0.592	1.21	1.36
PFORE	$q_{eq,1}$ (mmol U g^{-1})	0.608	0.604	1.18	1.36
	$k_1 \times 10^2 (min^{-1})$	8.88	9.59	16.7	12.1
	R^2	0.982	0.981	0.993	0.979
	AIC*	-127.6	-132.3	-159.6	-157.8
PSORE	$q_{eq,2}$ (mmol U g^{-1})	0.703	0.683	1.30	1.50
	$k_2 \times 10^2$ (L mmol ⁻¹	15.5	19.3	18.9	12.2
	min ⁻¹)				
	R^2	0.947	0.958	0.992	0.983
	AIC*	-110.4	-114.4	-154.3	-156.2
RIDE	$D_e \times 10^{11} \ (m^2 \ min^{-1})$	6.79	7.30	11.69	9.30
	\mathbb{R}^2	0.958	0.971	0.996	0.991
	AIC*	-119.5	-123.0	-167.2	-164.8

AIC (calculated on C(t)/C0 values). D0 (strontium): $4.75 \times 10^{-8} \text{ m}^2 \text{ min}^{-1}$.

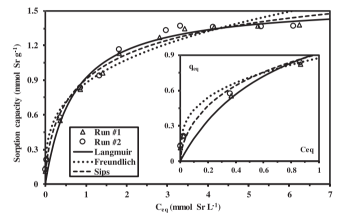


Fig. 4. Sr(II) sorption isotherm using $PAO_{20}MAA_{80}/SiO_2$ – Modeling with Langmuir, Freundlich and Sips equations (C_0 : 4–600 mg Sr L $^{-1}$ = 7.71–7.49; SD: 0.4 g L $^{-1}$; agitation time: 48 h; agitation speed: 170 rpm; T: 22 ± 2 °C).

Table 3Parameters of models for Sr(II) sorption isotherms using PAO₂₀MAA₈₀/SiO₂.

			0 20	30, 2
Model	Parameter	Run #1	Run #2	Combined
Experimental	q _m (mmol U g ⁻¹)	1.379	1.371	1.379
Langmuir	$q_{m,L}$ (mmol U g^{-1})	1.575	1.573	1.574
-	b_L (L mmol ⁻¹)	1.356	1.454	1.405
	R^2	0.990	0.983	0.988
	AIC	-53.6	-50.3	-53.4
Freundlich	$k_{\rm F}$	0.869	0.881	0.875
	n_{F}	3.349	3.361	0.875
	R^2	0.971	0.959	0.968
	AIC	-50.9	-47.0	-50.8
Sips	$q_{m,S}$ (mmol U g^{-1})	1.981	2.060	2.021
•	b _s (L mmol ⁻¹)	0.829	0.807	0.817
	n_S	1.556	1.675	1.617
	R^2	0.988	0.977	0.984
	AIC	-53.0	-48.0	-52.8

AIC (calculated on q_{eq} values).

at low residual metal concentration. Table 3 confirms that the Langmuir equation fits slightly better experimental curve than the alternative models. However, the calculated sorption capacity at saturation of the monolayer (i.e., $q_{m,L}$) significantly overestimates the maximum sorption capacity: 1.57 mmol Sr g $^{-1}$ νs . 1.38 mmol Sr g $^{-1}$. The affinity coefficient (i.e., b_L) stands around 1.41 L mmol $^{-1}$. The AIC calculation normalizes the fitting of experimental data for the different models in

Table 4Comparison of Sr(II) sorption properties of PAO₂₀MAA₈₀/SiO₂ with alternative sorbents (optimum pH, equilibrium time, Langmuir parameters).

Sorbent	pН	Time	$q_{m,L}$	b_{L}	Ref.
Clay	3	300	0.034	294	[99]
Almond green hull	6	3	1.33	0.421	[100]
Oxidized multiwall carbon nanotubes	7	1440	0.076	53.3	[10]
Moss	6	240	0.149	5	[16]
Magnetic chitosan beads	8.2	360	0.132	-	[101]
Zr-Sb oxide/PAN composite	4.74	270	0.498	73.6	[102]
Mixed Ni-K ferrocyanide/hydrated	7.8	180	1.598	0.366	[31]
TiO ₂					
Co-aminophosphonic chitosan	6	60	0.0389	78.0	[103]
Saccharomyces cerevisiae	3-4	2000	0.427	3.08	[14]
Nd-BTC-MOF	8	60-90	0.667	0.876	[37]
Melamine-styrene resin	8	240	1.63	3.33	[104]
Magnetic zeolite nanocomposite	6	20-30	1.01	1.31	[5]
T-46 (sulfonic)/A-33 (quaternary) mixed resins	7	30	0.109	8.39	[105]
Graphene oxide	6	1440	1.573	0.876	[106]
Amidoximated algal/PEI beads	6	60-90	2.36	2.01	[107]
Carboxyl-functionalized SiO ₂	10	30	0.913	1822	[27]
Zeolite	7.9	10	0.272	10.9	[108]
CHA-type zeolite	7	60	0.132	_	[109]
PAO ₂₀ MAA ₈₀ /SiO ₂	8	40–60	1.57	1.41	This work

Time: min; $q_{m,L}$: mmol Sr L^{-1} ; b_L : L mmol $^{-1}$.

function of the number of adjustable parameters. The Sips model that includes a third-adjustable parameter (compared with the two other models) could improve the quality of the fit; this is not the case, as shown by both the determination coefficients and the AIC values.

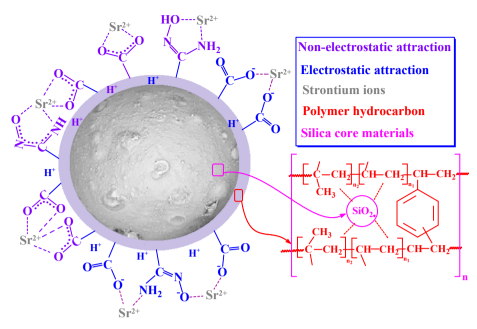
Table 4 compares the sorption properties of selected composite with alternative sorbents reported in literature. Combining the kinetic and equilibrium criteria, the bi-functionalized SiO_2 sorbent appears among the most efficient sorbents for $\mathrm{Sr}(\mathrm{II})$. It is noteworthy that, based on TGA analysis, the polymer content represents about 13.6% of the composite; this means that the functionalized polymer layer is highly efficient: sorption capacity represents about 10.15 mmol $\mathrm{Sr}~\mathrm{g}^{-1}$ functionalized polymer (not taking into account silica contribution to sorbent weight).

3.2.4. Sorption mechanism

The combination of data collected from the characterization sections and the sorption performances (pH effect and sorption isotherms) allows suggesting the main interactions involved in Sr(II) binding on PAO₂₀MAA₈₀/SiO₂ (Scheme 5). The main reactive groups on the sorbent consist of carboxylic acid and amidoxime moieties. At the optimum pH (close to neutrality), the pH_{PZC} value confirms the deprotonation of reactive groups. The slope analysis for pH effect (log plot of distribution coefficient vs. pH) does not allow identifying ion-exchange mechanism and strontium is supposed to be bound through chelation. FTIR analysis shows the contribution of carboxylic and amidoxime (through -NH, -NH₂, C=O and -OH functions). The molar ratio between carboxyl and amidoxime groups in $PAO_{20}MAA_{80}/SiO_2$ may explain the diversity of interaction modes, which is also confirmed by XPS analysis (identification of -N-Sr and -O-Sr bonds). Interactions may be of electrostatic and non-electrostatic nature due to deprotonation or electron delocalization of O- and N-based functional groups (Scheme 5).

3.2.5. Selectivity study

The selectivity of the sorbent for Sr(II) against other alkali and alkali-earth metal ions is shown in Fig. AM13. The fraction of strontium on the sorbent at equilibrium varies between 41% and 59% while varying the pH. The highest fractions are obtained at pH higher than 7. This is a first evidence that the sorbent has a marked preference for Sr (II). Indeed, the fraction in the solid is significantly enriched compared with the initial fraction in the solution (i.e., 20%). The cumulative



Scheme 5. Suggested mechanisms of Sr(II) sorption on bi-functionalized composite sorbent.

sorption capacity increases with up to pH 5.39 (1.39 mmol metal g^{-1}) and progressively decreases (down to 1.16 mmol metal g^{-1}) at pH 7.62.

On the other hand, consistently with the results obtained with single-component solutions, Sr(II) sorption continuously increases with pH and reaches (under selected experimental conditions) a sorption capacity close to 0.69 mmol Sr g⁻¹ (in equilibrium with a residual concentration close to 0.513 mmol Sr g⁻¹). This is also consistent with the sorption capacity reached on the sorption isotherm (single-component) for the same residual concentration (i.e., $\approx 0.7 \text{ mmol Sr g}^{-1}$ calculated from Freundlich and Sips equations, which provide the best fits of experimental isotherm in this concentration range). This means that the presence of co-ions in large excess does not impact the sorption capacity of Sr(II); although these co-ions can be bound to the sorbent (probably on other reactive groups). The cumulative sorption capacity reaches 1.16 mmol metal g^{-1} for a residual (cumulative) concentration close to 5.16 mmol metal L^{-1} . This concentration corresponds to the saturation plateau of the Sr(II) single-component isotherm at pH 8; the sorption capacity is found close to 1.36 mmol Sr g⁻¹ (1.38 mmol Sr g⁻¹ for simulated value, using the Langmuir equation). The multi-component solutions decreases by 16% the total sorption capacity compared with the corresponding residual concentration in mono-component solution. The sorbent is remarkably stable in sorption performance: Sr (II) sorption is hardly affected by the presence of competitor or co-ions and the total sorption capacity is of the same order of magnitude (though slightly lower than for Sr(II)-single solutions). It is noteworthy that at pH 5.39, Ca(II) sorption is relatively higher than the sorption of Ba(II), K(I) and Na(I).

Table AM10 shows the semi-quantitative analysis of $PAO_{20}MAA_{80}/SiO_2$ sorbent (surface) after being mixed with multi-component solutions at different pH values. These results confirm the relative selectivity of the sorbent for Sr(II). The atomic fraction at the surface of the sorbent for co-ions (i.e., Na(I), K(I), Ca(II) and Ba(II)) remains systematically below 0.08% while for Sr(II) the content increases with pH from 3 (i.e., 0.15%) up to 0.86–0.84% at pH 6–7, and increases again at pH 10 (up to 1.07%). The cumulative atomic fraction for co-ions ions does not exceed 0.22% at pH 6 to pH 8. In multi-component solutions, the sorbent shows a pH-response consistent with the results obtained with mono-component solutions.

Fig. 5 shows the selectivity coefficient ($SC_{Sr/metal} = D_{Sc}/D_{metal}$) of Sr (II) against Ba(II), Ca(II), K(I) and Na(I), at selected pH values. Regardless of the pH the SC is higher than 3; this confirms the preference

of the sorbent for Sr(II) over selected metal cations. However, the SC is strongly affected by the pH. Indeed, above pH 5 the SC tends to increase, especially between 7 and 7.6. Against Ba(II), Ca(II), K(I) and Na (I), the SC reaches a maximum at pH 7.62 with values ranging between 10.7 against Na(I) up to 17.6 against K(I). PAO $_{20}$ MAA $_{80}$ /SiO $_{2}$ preference for Sr(II) against other alkali-earth metal ions (such as Ca(II) or Ba(II) is confirmed by the SC ranging between 12 and 14.2 at pH $_{0}$ = 8 (pH $_{eq}$ = 7.62). At pH 6, where the differences are more marked between the different systems (SC $_{\rm Sr/metal}$), the selectivity increases according: Ca(II) < Na(I) < Ba(II) < K(I). This ranking was compared with the relative values of different physicochemical properties of these elements (summarized in Table AM11). Actually, a relatively good correlation can be found between the SC $_{\rm Sr/metal}$ parameters and the relative Pauling electronegativity ratios ($\chi_{\rm metal}/\chi_{\rm Sr}$), as shown in Fig. AM14.

Fig. AM15 provides complementary information on the comparison of sorption properties for selected metals. The \log_{10} plots of the distribution ratios is plotted against equilibrium pH. The superimposition of the experimental points for Sr(II) sorption in multi-component

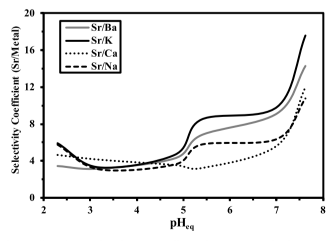


Fig. 5. Effect of pH on the selectivity coefficient for separation of Sr from monoand divalent cations (multi-component equimolar concentrations; C_0 (individual): 1.3 mmol metal L^{-1} ; SD: 0.125 g L^{-1} ; agitation time: 48 h; agitation speed: 170 rpm; T: 22 \pm 2 °C).

solutions with the curve obtained from mono-comp onent solutions (filled symbols) confirms that the sorption of Sr(II) is hardly affected by the presence of these co-ions. In addition, the sorption levels (D_{Sc}) are significantly higher than those observed for the other co-ions. In the case of Sr(II), the curve is linear between pH 2 and 8 (and stabilizes above), while for the other metals a slope break is observed at the maximum between pH 5 and pH 6: this is consistent with the enhanced selectivity of the sorbent for Sr(II) at pH higher than 6. The ranking in sorption for alkali and alkali-earth metal ions at the optimum pH follows the series: Ca(II) > Na(I) > Ba(II) > K(I). This ranking cannot be directly correlated to the charge of metal ions. Table AM11 reports the main physicochemical characteristics of selected metal ions. Actually, the ranking does not follow the ionic radius of hydrated species (Shannon data) [57]: $Na(H_2O)_6^+$ (1.02 Å) $< Ca(H_2O)_8^+$ (1.12 Å) < Ba $(H_2O)_8^+$ (1.42 Å) < $K(H_2O)_8^+$ (1.51 Å). A better correlation is reported with Pauling electronegativity Ca(II)(1) > Na(I) (0.93) > Ba(II) (0.89) > K(I) (0.82). This confirms the influence of co-ion electronegativity on the selectivity of Sr(II) sorption using PAO₂₀MAA₈₀/SiO₂.

3.2.6. Metal desorption and sorbent recycling

Metal desorption is a critical step in the design of a sorption process since the recovery of the metal and the recycling or the sorbent are imperative for the competitiveness of the treatment. Fig. AM16 shows the p erfect rep roducibility of desorp tion kinetics. Fig. 6 shows the desorption kinetics for Sr(II) elution from loaded $PAO_{20}MAA_{80}/SiO_2$ using 0.5 M HCl solutions. The desorption profiles are very close for the different levels of Sr(II) loadings: the desorption is little slower with the low metal loading (i.e., 0.59 mmol Sr g⁻¹) while the profiles are superimposed for the higher loadings (i.e., 1.21 and 1.36 mmol Sr g^{-1}). The sorption kinetic profiles were not superimposed (Fig. 2). Complete metal desorption occurs within 20-30 min of contact; this is shorter than the equilibrium time required for metal sorption (40-60 min). The PFORE better fits desorption kinetics than the PSORE (especially considering the complete desorption for PFORE while PSORE modeling does not achieve complete metal elution); this is another significant difference with the sorption step, where the differences between the models were less significant. The apparent rate coefficient (for PFORE) are little higher than the values obtained during sorption step (Tables 2 and 5).

Table 6 compares the performances of sorption and desorption for Sr(II) removal using PAO $_{20}$ MAA $_{80}$ /SiO $_{2}$ for five successive cycles. The desorption of Sr(II) from metal-loaded resin is very efficient and stable for 5 cycles using 0.5 M HCl solution as the eluent: metal desorption is completely achieved for, at least, the first five cycles. The sorption efficiency progressively decreases (from 58.3 \pm 0.5% to 54.5 \pm 0.1%). This 6.5%-loss in sorption efficiency (despite the complete desorption of the metal), is probably due to a partial degradation of the sorbent. This is an important parameter for higher number of recycling cycles. However, the combined sorption/desorption performances are globally very attractive for the selective recovery of Sr(II) from complex solutions.

3.2.7. Dynamic sorption - fixed-bed column

The resins were applied in fixed-bed column for dynamic sorption (Fig. 7 and Fig. AM17). The breakthrough curves are correctly fitted by the Thomas equation [55] (Table 7). The rep roducibility of breakthrough curves is clearly demonstrated by the superposition of experimental profiles. The comparison of linear (LR) and non-linear (NLR) regression analysis shows that the non-linear regression analysis gives a better mathematical fit. Increasing the flow rate (and superficial velocity) improves the quality of the fit. The calculated sorption capacity is consistent with the experimental values (little overestimation, by less than 4% for LR); in the case of NLR, the sorption capacity is more overestimated (despite the better global fit of experimental profiles). Surprisingly, the sorption capacity linearly decreases with superficial velocity (while using the same amount of sorbent and the same initial

metal concentration); this means that the saturation of the sorbent is not achieved. The expected sorption capacity at equilibrium is deduced from the sorption isotherms (close to 0.74 mmol Sr g⁻¹). The sorption capacities obtained in dynamic system are systematically much lower than the expected value, from 0.247-0.270 to 0.535-0.544 mmol Sr g⁻¹, depending on the flow rate. This is another confirmation that the sorbent is not fully saturated. Ang et al. [96] observed close values for the saturation capacities in the case of the sorption of anesthetic gaseous agent on activated carbon when varying the flow rate. The rate coefficient k_T (L mmol⁻¹h⁻¹) linearly increases with the superficial velocity (between 0.32 and 2.390 L mmol⁻¹h⁻¹ for LR; between 0.246 and 2.09 mmol⁻¹h⁻¹ for NLR); consistently with the conclusions raised by Ang et al. [96]. In the case of base metals using melamine-functionalized SBA-15 mesoporous silica, Shabazi et al. [97] reported that a maximum sorption capacity was obtained at intermediary flow rate; the relevant sorption capacities were also lower than the corresponding sorption capacities obtained in batch mode. Araneda et al. [98] investigated the dynamic sorption of copper on extractant-immobilized microcapsules; they also observed a significant decrease of the sorption capacity calculated by the Thomas model with increasing the flow rate. The rate coefficient also increased with the flow rate.

3.2.8. Sr(II) sorption from seawater

The semi-quantitative EDX analysis of seawater treatment by sorption onto $PAO_{20}MAA_{80}/SiO_2$ is reported in Table AM12 for different pH values. The SiO_2 core is appearing through the two major peaks of EDX spectra for Si and O elements that represent between 69 and 77% of the sorbent (atomic fractions, AF). The polymer coating is associated to C content, which increases from 9 to 12% (AF) when the pH increases from 4 to 8–10: this C increases may be explained by the formation of carbonate complexes or the accumulation of organic materials

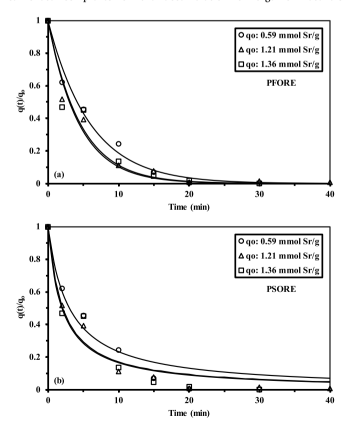


Fig. 6. Sr(II) desorption kinetics from metal-loaded $PAO_{20}MAA_{80}/SiO_2$ – Modeling with the PFORE (a) and the PSORE (b) (sorbent collected from uptake kinetic experiments; desorption conditions: 0.5 M HCl as eluent; SD: 1 g L⁻¹; agitation speed: 170 rpm; T: 22 \pm 2 °C).

Table 5 Parameters of kinetic models for Sr(II) desorp tion from metal-loaded $PAO_{20}MAA_{80}/SiO_2$ using the PFORE and PSORE fits [54].

Loaded conc. (mmol Sr g^{-1})	Model	PFORE		PSORE		
(IIIIIOI SF g)	Parameter	k _{D1} (min ⁻¹)	\mathbb{R}^2	β_2	$k_{\rm D2}~({\rm min}^{-1})$	R ²
Run #1 - 0.590 Run #2 - 0.591 1.21 1.36		0.171 0.163 0.228 0.218	0.985 0.970 0.979 0.949	0.989 0.990 0.992 1.000	0.319 0.327 0.492 0.484	0.970 0.964 0.973 0.951

PFORE: $\frac{q(t)}{q_0}=e^{-k_{D1}t}$ with: k_{D1} the apparent rate coefficient for desorption (min⁻¹) PSORE: $\frac{q(t)}{q_0}=\frac{1}{\beta_2+k_{D2}t}$ with: k_{D2} the apparent rate coefficient for desorption (min⁻¹) and β_2 (dimensionless) the constant for PSORE (in desorption).

Table 6 Sr(II) sorption and desorption efficiencies for $PAO_{20}MAA_{80}/SiO_2$ sorbent recycling.

Cycle #	Sorption efficiency (%) Average S.D.		Desorption efficiency (%)		
			Average	S.D.	
1	58.3	0.5	100.1	0.4	
2	57.0	0.4	100.3	0.6	
3	56.4	0.2	100.3	0.5	
4	54.8	0.2	100.0	0.6	
5	54.5	0.0	100.2	0.3	

Experimental conditions – Sorption: C_0 : 51 mg Sr $L^{-1}=0.582$ mmol Sr L^{-1} ; pH: 8.05; SD: 0.61 g L^{-1} ; time: 24h; T: 22 \pm 2 °C; agitation speed: 170 rpm/Desorption: eluent: 1 M HCl; SD: 1.5 g L^{-1} ; time: 2 h T: 22 \pm 2 °C; agitation speed: 170 rpm.

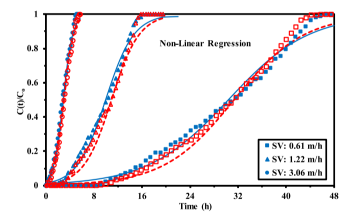


Fig. 7. Breakthrough curves for the sorption of Sr(II) using PAO $_{20}$ MAA $_{80}$ /SiO $_{2}$ – effect of superficial velocity (sorbent weight 400 mg; pH $_{0}$:8); modeling of experimental breakthrough curves with the Thomas equation; Run #1 (closed symbols and solid lines) C $_{0}$: 53 mg Sr L $^{-1}$ = 0.605 mmol Sr L $^{-1}$; Run #2 (open symbols and dashed lines) C $_{0}$: 54 mg Sr L $^{-1}$ = 0.615 mmol Sr L $^{-1}$; parameters determined by non-linear regression analysis).

contained in seawater that can bound to the sorbent active sites. Similarly, for S element: the content increases in neutral or alkaline solutions, probably associated to the binding of sulfate, hydrogen sulfate or metal sulfate. The amount of N element (in the range 4.64–6.34 %, AF) is due to amidoxime moiety. Strontium content increases from 0.04% at pH 4 to 0.54–0.46% at pH 8–10. This is consistent with the previous results obtained with synthetic solutions. Consistently with the trends observed for multi-component synthetic solutions, Ca(II) sorption hardly varies between 0.22 and 0.37%, K(I) between 0.01 and 0.15%, Mg(II) between 0.16 and 0.36 % (increasing with pH). The case of Na(I) is little different: Na content continuously decreases with pH (from 3.22 to 0.04%). Other elements are identified on EDX spectra:

Table 7 Modeling of breakthrough curves for the sorption of Sr(II) using $PAO_{20}MAA_{80}/SiO_2$ at different flow rates (Q, mL min⁻¹; Superficial flow velocity, SV, m h⁻¹) – Parameters of the Thomas model [55].

Q	sv	Run	$q_{exp.}$ (mmol Sr g^{-1})	Regr.	q _{model} (mmol Sr g ⁻¹)	k _T (L mmol ⁻¹ h ⁻¹)	R ²
0.2	0.61	#1	0.535	L	0.533	0.320	0.975
				NL	0.551	0.246	0.989
		#2	0.544	L	0.527	0.357	0.978
				NL	0.562	0.279	0.994
0.4	1.22	#1	0.346	L	0.357	0.829	0.984
				NL	0.359	0.702	0.993
		#2	0.372	L	0.381	0.843	0.983
				NL	0.403	0.705	0.992
1.0	3.06	#1	0.247	L	0.258	2.328	0.993
				NL	0.256	2.090	0.996
		#2	0.270	L	0.278	2.390	0.994
				NL	0.279	2.057	0.997

 $q_{th.} = 0.707 \, \text{mmol Sr g}^{-1}; \; \text{L: linear regression, NL: non-linear regression analysis.}$

Thomas equation:
$$\ln\left(\frac{C_0}{C(t)} - 1\right) = \frac{k_T q_{eq} m}{Q} - k_T C_0 t$$
.

Where k_T is the Thomas rate constant (L mg $^{-1}h^{-1}$), q_{eq} the sorption capacity at equilibrium (mmol Sr g $^{-1}$), m the amount of sorbent (g), Q the flow rate (L h^{-1}); t the time (h), C_0 and C(t): the inlet and outlet concentrations, respectively.

such as B(III), Mo(VI) and Ru(III). A special attention has been paid to the sorption kinetics of this series of metals or elements at different pH values. Fig. AM18 compares the kinetic profiles for traces removal: for B (III) and Sr(II) the residual concentration progressively decreases up to 48 h; the sorption of Mo(VI) is much faster (a pseudo-equilibrium, corresponding to 90% of total sorption, is reached within 6–10 h). For the other (major) elements (not shown), at much higher concentration, the variation of the concentration in the solution is relatively weak (less than 3%, and even less than 1% for Na(I) and K(I)); the equilibrium is reached within 12–24 h.

The sorption capacities are directly controlled by their metal concentration in seawater (Tables AM13 and AM14) and the pH of the solution: sorption capacities increase with pH (consistently with experiments on synthetic solutions), essentially between pH 4 and pH 8. The cumulative sorption capacity increases up to 8.84 mmol metal g $^{-1}$ at pH 10. The Sr(II) sorption capacity increases from 0.071 to 0.103 mmol Sr g $^{-1}$ when the pH increases from 8 to 10. Compared to the sorption isotherm obtained with synthetic pure solution, for a residual concentration close to 0.0104 mmol Sr L $^{-1}$ at pH 8, according the Sips equation (which fits the most closely the isotherm at low metal concentration), the expected sorption capacity should be close to 0.093 mmol Sr g $^{-1}$. This means that the loss in sorption capacity in such a complex solution does not exceed 25%.

The most significant enrichments (compared with initial metal concentrations: $q_{\rm eq}/C_0$, L g^{-1}) are observed for trace elements such as Sr(II) (up to 3.32 L g^{-1}), Mo(VI) (up to 3.16 L g^{-1}) and B(III) (up to 1.37 L g^{-1}) at pH $_0$ 10, while for the other metals the enrichment factor does not exceed 0.175 L g^{-1} . The values of distribution ratio (D, L g^{-1}) are also summarized in Table AM14: highest D values are observed at pH 10 for Sr(II) > Mo(VI) and B(III). The selectivity coefficients for Sr over co-ions are reported in Fig. AM19. The SC increases with time and pH. The systems are difficultly comparable due to the large excess in the concentration of the major elements; however, these data offer an easy comparison tool for highlighting the preference of the sorbent for Sr(II) over the co-ions. Highest SCs are observed according the sequence: Na (I) $(8\,4\,6) >$ K(I) $(2\,3\,8) >$ Mg(II) $(1\,9\,2) >$ Ca(II) $(1\,0\,3) >$ Ru(III) (62) > B(III) (5). On the opposite hand, the sorbent has a little preference for Mo(VI) against Sr(II) (SC_{Sr/Mo} < 1) at pH 4 and 8; at pH 10, the sorbent does not show selectivity between the two metal ions.

These results show that in such a complex solution as seawater, the

sorbent maintains:

- (a) good sorption properties for Sr(II),
- (b) high affinity for other trace metal ions (B(III) and more specifically Mo(VI)), and
- (c) strong enrichment factor and high relative distribution ratios for these trace metal ions (despite huge excess of Na(I), Mg(II), L(I) and Ca(II)).

4. Conclusion

The combined grafting of both polyacrylic and amidoxime groups on silica microspheres (size: around 120 µm) allows enhancing the sorption properties of the composite for Sr(II), compared with monofunctional sorbents (bearing solely PMAA or PAO moieties). However, the comparison of sorption performances shows that maximum sorption efficiency for Sr(II) is reached for specific substitution grade (i.e., corresponding to PAO₂₀MAA₈₀/SiO₂ composite sorbent). The sorbents are characterized by elemental analysis (3.36-7.8 mmol N g⁻¹), semiquantitative EDX analysis, textural properties (SSA: 62-69 m² g⁻¹), TGA (about 13.6% weight loss at 875 °C, representing the hydrocarbon coating fraction), pH_{PZC} (6.42-6.51 for PAO₂₀MAA₈₀/SiO₂ composite sorbent). FTIR and XPS analyses confirm the appearance of functional groups on the polymer coating layer of composite materials, consistently with suggested synthesis procedures. In addition, the changes in spectra confirm the contribution of amine, hydroxyl/carboxyl groups in the binding of Sr(II) on the composite sorbent.

Optimum sorption occurs at pH 8. The uptake is relatively fast: equilibrium is reached within 40-60 min of contact. The sorption is fitted by the pseudo-first order rate equation: apparent rate coefficient varies in the range $0.09-10.17 \,\mathrm{min}^{-1}$. The maximum sorption capacity is closed to $1.38 \, \text{mmol Sr g}^{-1}$ (10.15 mmol Sr g $^{-1}$ when reported in function of the effective amount of functionalized polymer in the composite); the isotherms are equally fitted by the Langmuir (capacity at saturation of the monolayer close to 1.57 mmol Sr g⁻¹) and the Sips equations. These sorption capacities are among the most promising compared with available literature data. The recycling of the sorbent is verified over 5 cycles of sorption and desorption: 1 M HCl solution offers complete, fast (equilibrium within 20 min) and stable desorption of Sr(II) while the sorption efficiency slightly decreases (less than 7% at the fifth cycle). In equimolar multi-component solutions, the sorption is selective for Sr(II) over alkali and alkali earth metal ions. This is confirmed by the tests performed on Sr(II) sorption in seawater: high distribution ratio (in the range 1-10 L g⁻¹) are observed for Sr(II) and B (III) or Mo(VI), despite the huge excess of alkali and alkali-earth metal ions. In dynamic systems (i.e., fixed-bed column), the breakthrough curves for Sr(II) sorption are finely fitted by the Thomas model; although the sorption capacity is significantly under evaluated compared with static systems.

Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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Appendix A. Supplementary data

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