Highly selective cesium(I) capture under acidic conditions by a layered sulfide

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Radiocesium remediation is desirable for ecological protection, human health and sustainable development of nuclear energy. Effective capture of Cs\(^+\) from acidic solutions is still challenging, mainly due to the low stability of the adsorbing materials and the competitive adsorption of protons. Herein, the rapid and highly selective capture of Cs\(^+\) from strongly acidic solutions is achieved by a robust K\(^+\)-directed layered metal sulfide KInSnS\(_4\) (InSnS-1) that exhibits excellent acid and radiation resistance. InSnS-1 possesses high adsorption capacity for Cs\(^+\) and can serve as the stationary phase in ion exchange columns to effectively remove Cs\(^+\) from neutral and acidic solutions. The adsorption of Cs\(^+\) and H\(_3\)O\(^+\) is monitored by single-crystal structure analysis, and thus the underlying mechanism of selective Cs\(^+\) capture from acidic solutions is elucidated at the molecular level.

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With the rapid development of nuclear energy, growing concerns about its safety have been raised. $^{137}$Cs, as the fission product of $^{235}$U, is one of the main sources of radioactivity in spent fuel ($1230$ g/t) $^{1}$. $^{137}$Cs with a long half-life (12.3 years) can emit gamma-rays. It is highly migratory in the environment, causing cell damage, cancer, and even death in humans. $^{2}$ Once radioactive cesium is released into the environment, it will be extremely polluting and cause harm to the entire ecosystem. $^{3}$ For example, the Fukushima nuclear accident released a large amount of radioactive Cs$^{+}$ ions into the environment, causing total radioactivity levels in some fish to remain above the limit (100 Bq/kg) until now. $^{4}$ Therefore, the disposal of $^{137}$Cs has received much attention due to its potential threat to the environment.

The strongly acidic high-level-liquid-wastes (HLLWs) resulting from the recovery of uranium and plutonium from spent fuel by the PUREX process contain $^{137}$Cs in ionic form. $^{5-7}$ The key for the disposal of HLLWs is the removal of strongly radioactive ions (such as $^{137}$Cs) to reduce their radioactivity level. $^{6}$ The separated and purified $^{137}$Cs can also be prepared as isotope sources and reused in agriculture and medical fields. $^{3}$ However, HLLWs are extremely complex, containing not only Cs$^{+}$ but also Sr$^{2+}$, La$^{3+}$, Na$^{+}$ and so on. $^{8,9}$ Such complex components would pose a huge challenge for the selective separation of Cs$^{+}$ from HLLWs. Methods such as solvent extraction, chemical precipitation, and ion exchange/sorption have been used for the enrichment and separation of radiocesium. $^{9}$ These methods, however, have some disadvantages in terms of operation, cost, or selectivity, such as the expensive and toxic extractants used in solvent extraction and the great volume of radioactive sludge produced by precipitation. $^{10}$ Although ion exchange is considered an ideal method for controlling radioactive contamination due to its simplicity of operation, high efficiency, and lack of secondary contamination, the development of stable and highly selective ion exchangers for the efficient capture of Cs$^{+}$ in acidic solutions still remains a great challenge.

Due to the instability or poor selectivity of materials for Cs$^{+}$ capture under acidic conditions, most of the current studies are restricted to that under neutral or weak acidic conditions. $^{9,11,12}$ In contrast, materials that can effectively remove Cs$^{+}$ ions under extremely acidic condition are still very limited, exemplified mainly by ammonium phosphomolybdate and its complexes or cupric aromatic crown ether-modified silyl compounds. $^{13-16}$ In recent years, metal sulfides have been a very promising class of radioactive ion exchangers, which display excellent removal performance for Cs$^{+}$. However, the capture of Cs$^{+}$ by metal sulfides under strongly acidic conditions (nitric acid concentrations $>1$ mol/L) is challenging, and in particular, the adsorption mechanism has been not clearly identified. Therefore, it is of vital significance to develop acid-tolerant ion exchangers that can selectively capture Cs$^{+}$ from strongly acidic solutions for radioactive liquid waste treatment and to clarify the mechanism of Cs$^{+}$ removal for revealing the structure–function relationship.

Herein, the strategy to improve stability of materials by introducing Sn$^{4+}$ and In$^{3+}$ with high valency and large radii into sulfides has been applied. Such an approach leads to a robust K$^{+}$-directed layered metal sulfide KInSnS$_4$ (denoted as InSnS-1) with excellent acid and irradiation resistances. Specifically, it can maintain its $[\text{InSnS}_4]$ layers even in 1-4 mol/L HNO$_3$ solutions. InSnS-1 has high Cs$^{+}$ ion-exchange capacity ($q^\text{m}=316.0$ mg/g in neutral solutions; $q^\text{m}=272.6$ mg/g in 1 mol/L HNO$_3$ solutions). Under the coexistence of high-concentration competing ions such as Na$^+$, Sr$^{2+}$, and La$^{3+}$, InSnS-1 exhibits high selectivity for Cs$^{+}$ in acidic solutions, which endows InSnS-1 with excellent Cs-Sr or Cs-La separation in the acidic environment. Moreover, ion-exchange columns loaded with InSnS-1 can effectively treat neutral and acidic solutions containing high concentrations of Cs$^{+}$ (190.13 and 84.25 mg/L, respectively) with treatment efficiencies up to 1300 and 650 bed volumes, respectively. The successful determination of the single-crystal structures of InSnS-1 and its three ion-exchange products help visualize the ion exchange between Cs$^{+}$ or hydrated protons (H$_3$O$^+$) and interlayered K$^+$ ions of InSnS-1. This is a systematic study of the selective Cs$^{+}$ capture under strongly acidic condition with an emphasis on revealing the capture mechanism at the molecular level.

### Results

#### Synthesis

Among the reported metal sulfide ion exchangers, K$_2$MnSn$_3$-S$_6$ ($x = 0.5$–0.95, KMS-1) and KInSnS$_4$ (KMS-5) have shown excellent acid resistance (retaining the parent structure at pH < 1). $^{1,7,11}$ A common feature of both compounds is the high coordination number of metal ions in the framework (i.e., metal ions are coordinated with five S atoms to form octahedral geometries), which is the highest among the reported metal sulfide ion exchangers. $^{13,17}$ The high coordination of metal ions may be beneficial to the structural stability of metal sulfides. In addition, it has been shown that the introduction of high-valency metal ions in the framework helps to resist the exchange of competing protons. $^{8}$ Therefore, we used the strategy of introducing high-valency, large-radius metal ions to synthesize metal sulfides with highly coordinated metal ions aiming to improve the acid stability of sulfide ion exchangers and their ability to selectively capture target ions. Thus, a robust K$^{+}$-directed layered metal sulfide InSnS-1 has been synthesized by a simple solid-phase method according to the following reaction formula (Eq. (1)):

$$\text{KInS}_2 + \text{Sn} + 2 \text{S} \xrightarrow{750 \text{C}, 4 \text{days}} \text{KInSnS}_4$$

Although KInSnS$_4$ with the same stoichiometric ratio has been reported, the previous synthesis methods yielded mixed phase of KInSnS$_4$ with two distinct structures (α-KInSnS$_4$ and β-KInSnS$_4$). $^{23}$ Moreover, the previous synthesis methods reported are operationally complex, requiring the use of Cs$_2$ vapor stream or K$_2$CO$_3$. Notably, when K$_2$CO$_3$ is used, the reaction mixture usually needs to be heated to a specific temperature to release carbon dioxide firstly before sealing the quartz tube. By contrast, KInS$_2$ as a precursor was utilized here, which allows the synthesis of single-phase KInSnS$_4$ (InSnS-1) on a large scale easily. Moreover, though the structure of InSnS-1 is similar with α-KInSnS$_4$, InSnS-1 shows different K$^+$ positions. The detail can be found in the following crystal structures part.

#### Crystal structures and insight into the Cs$^{+}$ removal mechanism under neutral and acidic conditions

The reported α-KInSnS$_4$ and β-KInSnS$_4$ both present two-dimensional (2D) anionic layer of [InSnS]$_n$-octahedra interconnected via edge sharing. $^{23}$ The difference is that in α-KInSnS$_4$, the S atoms in the [InSnS]$_n$ octahedron are disordered with half-occupancy resulting in two alternative orientations for [InSnS]$_n$-layers (Supplementary Fig. 3). $^{23}$ Accordingly, the interlayer disordered K$^+$ ions are hexa-coordinated with S atoms to form KS$_6$, with both triangular antiprismatic and prismatic orientations in α-KInSnS$_4$, while only triangular antiprismatic KS$_6$ are formed in β-KInSnS$_4$. Although it seems that InSnS-1 has similar structure as α-KInSnS$_4$ (Supplementary Fig. 3), structural refinements of InSnS-1 show that the K$^+$ ions reside in two different sites of K1 (1b) and K1B (2d), rather than only one site (1b) in α-KInSnS$_4$ (Supplementary Fig. 4a, b).

The ion-exchange properties of α-KInSnS$_4$ and β-KInSnS$_4$, especially for the selective capture of Cs$^{+}$ under strongly acidic condition have not been systematically investigated. In order to
Supplementary Tables 1

...investigate the capture mechanism of Cs\(^+\) under neutral and acidic conditions, the single-crystal structures of ion-exchange products (InSnS-1-Cs, InSnS-1-Cs/H, InSnS-1-H) were analyzed and compared with that of the pristine InSnS-1. This allows us to visualize the mechanism of selective Cs\(^+\) capture by InSnS-1 and effects of protons on the process of Cs\(^+\) capture. Combining single-crystal structure analysis and EDS results, the ion-exchange reaction can be represented by the following equations (Eqs. (2)–(4)):

\[
\text{KInSn}_4 + \text{Cs}^+ \rightarrow \text{CsInSn}_4(\text{InSnS}-1-\text{Cs}) + \text{K}^+ \tag{2}
\]

\[
\text{KInSn}_4 + \frac{1}{3}\text{Cs}^+ + \frac{2}{3}\text{H}_2\text{O}^+ \rightarrow \text{Cs}_3(\text{H}_3\text{O})_2\text{InSn}_4(\text{InSnS}-1-\text{Cs}/\text{H}) + \text{K}^+ \tag{3}
\]

\[
\text{KInSn}_4 + \text{H}_2\text{O}^+ + \text{H}_2\text{O} \rightarrow (\text{H}_3\text{O})\text{InSn}_4 \cdot \text{H}_2\text{O}(\text{InSnS}-1-\text{H}) + \text{K}^+ \tag{4}
\]

Detailed structural information of compounds is shown in Supplementary Tables 1–6. Compared with InSnS-1, both InSnS-1-Cs and InSnS-1-Cs/H exhibit an increase in the interlayer spacing (from 8.431 Å to 8.690 Å and 8.939 Å, respectively; Fig. 1a–c), which is attributed to the substitution of K\(^+\) (138 pm) by Cs\(^+\) with larger radius (170 pm). Although the radius of H\(_2\)O\(^+\) (112 pm) is smaller than that of K\(^+\), the interlayer spacing of InSnS-1-H still increases to 8.631 Å because of the entry of one lattice water molecule per formula. These results indicate that the interlayer spacing of InSnS-1 is tunable upon ion exchange and its layer structure shows flexibility even under acidic conditions, which contributes to the effective capture of Cs\(^+\).

It is noteworthy that each Cs\(^+\) ion is coordinated by six S atoms to form CsS\(_6\) with triangular antiprismatic and prismatic geometries in both InSnS-1-Cs and InSnS-1-Cs/H with Cs–S distances of 3.606 Å and 3.675 Å, respectively (Supplementary Fig. 5), which indicate strong interactions between S\(^2-\) and Cs\(^+\). Based on Pearson’s Hard-Soft-Acid-Base (HSAB) theory, K\(^+\) is a hard acid whereas Cs\(^+\) is relatively softer. Thus, the soft basic S\(^2-\) from [InSnS\(_4\)]\(^{2-}\) layers has a stronger affinity for relatively soft acidic Cs\(^+\) than K\(^+\). Therefore, the flexible framework and the adjustable layer spacing of InSnS-1 as well as the strong interaction of Cs\(^+\) with S\(^2-\) play important roles for the rapid and highly selective capture of Cs\(^+\).

In addition, K\(^+\) ions can be completely exchanged by H\(_3\)O\(^+\) ions under acidic conditions without Cs\(^+\) ions. H\(_2\)O\(^+\) ions or water molecules become more disordered between the layers in InSnS-1-H (Fig. 1d). Although there have been studies on the capture of Cs\(^+\), the clear structural revelation is still scarce. Especially, the microstructural illumination for the selective Cs\(^+\) capture under acidic conditions has not been reported. The current deep insight into the structural analyses facilitates the understanding of the mechanism of selective Cs\(^+\) capture by ion exchange and provides a reference for the design and synthesis of novel ion exchangers with the high selectivity for target radionuclides.

To further investigate the relationship between structures and properties of materials, a number of K\(^+\)-directed layered or three-dimensional (3D) microporous sulfides including 3D-K\(_2\)Cu\(_2\)Ge\(_2\)S\(_6\)\(^{30}\), 2D-KCu\(_2\)S\(_3\)\(^{31}\), 2D-KYS\(_2\)\(^{32}\) have been synthesized. It is a pity that our preliminary batch absorption experiments show that they have no Cs\(^+\) ion-exchange properties. To comparatively analyze the structure-function relationship, the reported structures including 2D-KMS-1\(^{17,22}\), 2D-KMS-2\(^{18}\), 2D-KTS-3\(^{19}\), and their ion-exchange properties have been also summarized. The shortest distances from K\(^+\) to anionic frameworks (noted as D\(_{\text{min,K-S}}\)) and the coordination numbers of K\(^+\) have been summarized (Supplementary Table 7).

In the case of 2D layered sulfides, 2D-KYS\(_2\) features an anionic layer of [YS\(_2\)]\(^{2-}\) built up by edge-sharing [YS\(_4\)] octahedra\(^{32}\), which is similar to KMS ion exchangers\(^{8,17,18}\). However, 2D-KYS\(_2\) does not have ion-exchange properties for Rb\(^+\), Cs\(^+\), and Sr\(^2+\), even though 2D-RbYS\(_2\) with the same layer structure as KYS\(_2\) has been reported. It is found that D\(_{\text{min,K-S}}\) in 2D-KYS\(_2\) (3.174 Å) is much shorter than those in 2D-KMS-1 (3.353 Å), 2D-KMS-2 (3.494 Å), and 2D-KMS-5 (3.439 Å). 2D-β-K\(_2\)CdS\(_2\)S\(_6\)\(^{33}\) with short D\(_{\text{min,K-S}}\) (3.157 Å) also cannot undergo ion-exchange...
reactions. Thus, the strong attraction between $\text{S}_2^{2-}$ of anionic layers and K$^+$ makes it difficult for K$^+$ to escape. In the case of 3D microporous sulfides, 3D-$\text{K}_2\text{SnS}_4$ has a large $D_{\text{min}}^\text{K-S}$ (3.281 Å) but cannot accommodate other ions such as Cs$^+$ due to small pore size excluding K$^+$ (3.7 Å $\times$ 19.4 Å)\textsuperscript{34}. Conversely, 3D-$\text{K}_2\text{SnS}_4$ has three different cavities, namely K1, K2, K3 cavities. The K$^+$ ions (K3: 4 coordination, $D_{\text{min}}^\text{K-S} = 3.253$ Å) in K3 cavity (diameter: $\sim$4.1 Å) can be easily exchanged by Cs$^+$ with strong selectivity. The K$^+$ ions in the smaller K1 and K2 cavities ($\sim$3.0 Å; K1: 8 coordination, $D_{\text{min}}^\text{K-S} = 3.446$ Å; K2: 6 coordination, $D_{\text{min}}^\text{K-S} = 3.144$ Å), however, cannot be exchanged by Cs$^+$ due to the limitation of the pore size.

Based on these facts, it can be inferred that $D_{\text{min}}^\text{K-S}$ is one of the main key factors in determining whether ion exchange occurs for layered sulfides. The smaller the value of $D_{\text{min}}^\text{K-S}$ is, the more attractive the anionic sulfide framework is for K$^+$, and the less likely ion exchange will occur. Based on the currently available data, we can tentatively assume that ion exchange can be realized when $D_{\text{min}}^\text{K-S}$ is longer than 3.30 Å, and otherwise ion-exchange reaction difficulty occurs for layered sulfides. $D_{\text{min}}^\text{K-S}$ of 2D-InSnS-1 is 3.503 Å here, and thus the current compound exhibits excellent ion-exchange property. However, for 3D microporous compounds, the pore size and the coordination number of K$^+$ are also necessary factors in addition to the $D_{\text{min}}^\text{K-S}$. If necessary, elevated reaction conditions will favor the promotion of ion-exchange reactions\textsuperscript{35}. Of course, the universal conclusion remains to be analyzed in more depth with more examples.

Characterizations of ion-exchange products. Single-crystal X-ray diffraction directly confirms that InSnS-1-Cs, InSnS-1-Cs/H, and InSnS-1-H can be obtained by ion exchange of InSnS-1 under different conditions. The successful ion exchange of K$^+$ by Cs$^+$ or H$_3$O$^+$ can be further corroborated by EDS, XPS, ICP. The X-ray powder diffraction patterns of InSnS-1 and its ion-exchange products were in high agreement with the corresponding single-crystal simulated ones (Supplementary Fig. 7). The elemental distribution mapping and EDS analysis results of ion-exchange products show that Cs$^+$ ions are evenly distributed in the samples (Supplementary Figs. 8 and 9). As seen in the XPS spectra (Supplementary Fig. 10), two peaks of K 2p at 292.1 eV (K 2p$_{3/2}$) and 294.8 eV (K 2p$_{1/2}$) disappeared after Cs$^+$ or H$_3$O$^+$ ion exchange, which confirms that K$^+$ ions were completely exchanged. Conversely, the appearance of Cs 3d$_{5/2}$ and Cs 3d$_{3/2}$ peaks at 723.3 eV and 737.2 eV was observed in the XPS spectra of InSnS-1-Cs and InSnS-1-Cs/H\textsuperscript{36}.

Compared to InSnS-1, InSnS-1-Cs has a significantly darker yellow color, while InSnS-1-Cs/H and InSnS-1-H display lighter yellow colors (Fig. 1), which correlates with the redshift and blueshift of their optical absorption edges, respectively. Compared to InSnS-1 (2.211 eV), the optical absorption edge of InSnS-1-Cs is slightly red-shifted to 2.147 eV, while InSnS-1-Cs/H and InSnS-1-H are blue-shifted to 2.255 and 2.220 eV, respectively (Supplementary Fig. 11). TG analysis (Supplementary Fig. 12) shows that InSnS-1 has good thermal stability. In addition, it can be seen that the exchanged products have obvious delamination compared with the pristine from the SEM images (Supplementary Figs. 13 and 14).

Acid stability studies. A scavenger for Cs$^+$ should be acid-stable as HLLWs are typically highly acidic\textsuperscript{6}. The InSnS-1 crystals retained the crystallinity and the layered network after being immersed in highly concentrated (0.1–4 mol/L) acidic solutions for 10 h (Fig. 2a, b). The leaching rates of In$^{3+}$ and Sn$^{4+}$ in solution were less than 2.79% and 0.06%, respectively, which further confirms the high acid resistance of InSnS-1 (Fig. 2c). As the acidity increased further to 5 mol/L HNO$_3$, the substance began to break down.

Kinetic studies of Cs$^+$ ion removal. It is essential that the ion exchanger is able to capture the target ions quickly. Therefore, kinetic experiments for the Cs$^+$ capture by InSnS-1 in neutral and strongly acidic solutions were carried out. In neutral solution, the removal rate R (%) of Cs$^+$ by InSnS-1 rapidly reached over 90% within 5 min (Fig. 3a). In 1 mol/L HNO$_3$ solution, R$_\text{Cs}$ also rapidly reached over 80% within 20 min (Fig. 3b). Such a fast and efficient removal of Cs$^+$ under acidic conditions is rarely reported because ion-exchange competition with protons is a severe issue\textsuperscript{8,20}. Compared to the pseudo-first-order kinetics model\textsuperscript{37}, the kinetic data of Cs$^+$ capture by InSnS-1 in neutral and 1 mol/L HNO$_3$ solutions were better fitted with the pseudo-second-order kinetics model\textsuperscript{37} with high correlation coefficients R$^2$ (>0.9999) (Supplementary Fig. 15), which proves that the adsorption process is chemical sorption\textsuperscript{38}.

Cs$^+$ ion adsorption isotherm studies. The exchange capacities of InSnS-1 for Cs$^+$ in neutral and 1 mol/L HNO$_3$ solutions at room temperature (RT) have been measured by static batch experiments. The ion-exchange capacity can be determined by fitting the equilibrium data with the Langmuir and the Freundlich isotherm models\textsuperscript{39}. The isotherm curves (Fig. 3c, d) were better fitted with the Langmuir–Freundlich isotherm model, with the correlation coefficients (R$^2$) of 0.99 and 0.98 in neutral and 1 mol/L HNO$_3$ solutions, respectively. The maximum exchange capacity ($q_{\text{max}}^{\text{Cs}}$) of InSnS-1 for Cs$^+$ reached 316 mg/g in neutral solutions, which is higher than that of many common Cs$^+$ adsorbents\textsuperscript{9}, including bentonite (Turkish samples: 300.35 mg/g\textsuperscript{40}), zeolite (Turkish samples: 89.18 mg/g\textsuperscript{40}), ferri-cyanide (Prussian blue granules: 241 mg/g\textsuperscript{41}, carbon-based materials (GO: 76.9 mg/g\textsuperscript{42}), metal sulfides (KMS-117: 226 mg/g, KTS-319: 300.35 mg/g\textsuperscript{40}). It is noteworthy that InSnS-1 can still hold an exchange capacity for Cs$^+$ of 98.6 mg/g in strongly acidic solutions (1 mol/L HNO$_3$), which is even higher than that of KMS-1 at pH = 0.8 (~90 mg/g\textsuperscript{17}). Noteworthy is that
InSnS-1 under a due to the effects of protonation8,20. Some previously reported R ion-exchange materials such as KMS-218, KTS-319, FJSM-GAS-147, 1.6 × 103 mL/g (Equilibrium data for Cs carried out. The distribution coefficient has an effect on the adsorption/ion exchange20,44,45. The majority pH-dependent experiments describe the affinity of the material for the target ions.

pH-dependent experiments. In general, the pH value of solution has an effect on the adsorption/ion exchange20,44,45. The majority of reported materials are unstable under strongly acidic conditions or their ion-exchange properties are significantly reduced due to the effects of protonation8,20. Some previously reported good-performance Cs+ ion-exchange metal sulfides have shown a significant decrease in their ion-exchange performance when the concentration of HNO3 just approaches 1 mol/L18,19,46. For example, KMS-218, KTS-319, and FJSM-SnS46 have reduced concentration of HNO3 around 103 mL/g at pH 62.1%) even in 2 mol/L HNO3 solution. KdCs and KdSr values all above 104 mL/g, respectively (Supplementary Fig. 17). The separation factor $K_{des}$ of InSnS-1 in 1 mol/L HNO3 solutions exceeded 100 when the Na/Cs molar ratios were 191, 387, and 983 ($K_{des}$ of 172.3, 109.6, and 166.2, respectively). SF is related to the selectivity of the material which is used to measure whether the two ions can be well separated18. Excellent selectivity can be considered when the SF value is higher than 10018. Such excellent selectivity of InSnS-1 for Cs+ is attributed to that Na+ is hard acid, while Cs+ is softer acid compared to Na+. According to HSAB theory25, the soft base S2− has a stronger affinity for Cs+ than Na+.

The effect of some other ions in HLLWs (represented by Sr2+ and La3+) on the selective capture of Cs+ cannot be neglected; thus, extra experiments were performed8. In the individual competing ion system, the influence of solution acidity and competing ion concentrations on the performance of InSnS-1 in removing Cs+ is similar. Under neutral conditions, when the concentrations of Cs+ and competing Sr2+ or La3+ were low (i.e., the Sr/Cs molar ratios were 1.44 and 14.9 or the La/Cs molar ratios were 2.09 and 21.8), InSnS-1 possessed high Kd and R (above 103 mL/g and 80%) for Cs+, Sr2+, La3+, respectively (Fig. 5a, d). The maximum values of KdCs, KdSr, KdLa were 4.74 × 104 mL/g, 8.57 × 103 mL/g, 4.89 × 105 mL/g, respectively, and the maximum removal rates of these ions were 97.93%, 98.85%, and 99.80%, respectively. In 1 mol/L HNO3 solution, KdCs are still above 103 mL/g with $R_{dCs}$ around 80% even under very high concentrations of competing ions. By contrast, the removal irradiation stability studies. Irradiation stability is one of the necessary properties for radioactive ion scavengers. Therefore, the Cs+ ion-exchange capacities of InSnS-1 samples after 100 kGy and 200 kGy γ-irradiation were investigated. The results demonstrate that InSnS-1 has excellent irradiation stability, maintaining stable structure and ion-exchange properties ($R_{dCs}>91%$, $K_{dCs}>10^4$ mL/g) after intense irradiation (Fig. 3f, Supplementary Fig. 17).

Competitive experiments in the presence of interfering ions. The selective extraction of Cs+ from HLLWs containing high concentrations of Na+ has been a major challenge because Na+ and Cs+ have similar hydration radii (219 pm for Na+ and 218 pm for Cs+)8,24. Therefore, the performances of InSnS-1 capturing Cs+ in neutral and acidic solutions with different Na/Cs molar ratios were investigated in detail. In neutral solutions, $K_{dCs}$ could still amount to 103 mL/g when the Na/Cs molar ratio was 380 (Fig. 4a). In 1 mol/L HNO3 solutions, $K_{dCs}$ could still reach 2.16 × 103 mL/g ($R_{dCs}=68.35$%) even when the Na/Cs molar ratio was 1.92 × 105 (Fig. 4b). Importantly, $R_{dCs}$ and $K_{dCs}$ of InSnS-1 in 1 mol/L HNO3 solutions were higher than those in neutral solutions when the Na/Cs molar ratios were in the range from 190 to 3.07 × 103, and the highest values of $K_{dCs}$ and $R_{dCs}$ were 4.0 × 103 mL/g and 79.71%, respectively (Supplementary Fig. 18). Even in 3 mol/L HNO3 solutions with Na/Cs molar ratios of about 2.9 × 103 and 5.8 × 103, $R_{dCs}$ could still reach 37.11% and 31.06%, respectively, which were higher than $R_{dCs}$ values in neutral or 1 mol/L HNO3 solutions under the same conditions (Fig. 4c, Supplementary Fig. 18).

The adsorption capacity of sulfide materials for Cs+ under such strongly acidic condition has not been found before.
performed InSnS-1 for Sr$_2$$^+$ or La$_3$$^+$ in 1 mol/L HNO$_3$ were significantly lower, with $K_{d}^{M}$ of 7.49–54.1 mL/g and $R^{M}$ of 0.74%–5.13% (M = Sr or La) (Fig. 5b, e). Even in 3 mol/L HNO$_3$ solutions containing high concentrations of competing Sr$_2$$^+$ or La$_3$$^+$, InSnS-1 still held $R^{Cs}$ of 32.78%–47.74%, while $R^{M}$ (M = Sr or La) were only in the range of 0.42%–6.15% (Fig. 5c, f).

Obviously, the enhancement of acidity has resulted in a greater attenuation in the removal efficiencies of Sr$_2$$^+$ and La$_3$$^+$ rather than Cs$. In 1 mol/L HNO$_3$ solutions, $SF_{CSr}$ (183.45, 630.87, and 110.70) exceeded 100 at three different Sr/Cs molar ratios, while $SF_{CSr/La}$ also reached 184.31 and 133.93 when La/Cs molar ratios were 2.30 and 21.5, respectively. Even under 3 mol/L HNO$_3$ condition, $SF_{CSr}$ remained 170.00 and 189.97 at the Sr/Cs molar ratios of 1.48 and 14.8, respectively. The above results demonstrate that InSnS-1 can effectively achieve Cs-Sr (or Cs-La) separation in acidic solutions even though the concentration of Sr$_2$$^+$ (or La$_3$$^+$) is much higher than that of Cs$^-$. Sulfide ion exchanger AgSnSe-1 can achieve Cs-Sr separation ($SF_{CSr/La}$ = 121.4) in neutral solutions with low concentration of Cs$^-$ and Sr$_2$$^+$ (both around 6 mg/L). However, to our best of our knowledge, no example of $SF_{CSr/La}$ value that exceeds 100 in strongly acidic solution has been reported, where the Sr$_2$$^+$ (or La$_3$$^+$) ion concentration is much higher than the Cs$^-$ ion concentration. Clearly, the current InSnS-1 is the first case for the excellent Cs-Sr or Cs-La separation in an acidic environment, which demonstrates again its excellent selectivity for Cs$^-$ capture.

We also investigated the selective capture ability of InSnS-1 under the coexistence of Sr$_2$$^+$, La$_3$$^+$, and Cs$. In neutral solutions, their $K_{d}$ (6.5 $\times$ 10$^5$ mL/g, 5.3 $\times$ 10$^4$ mL/g, 1.96 $\times$ 10$^4$ mL/g, respectively) and $R$ values (99.85%, 98.15%, 95.14%, respectively) were all very high when the Sr/La/Cs molar ratio was 1.38:1.27:1 (Fig. 5g). Nevertheless, in 1 mol/L HNO$_3$ solution, $K_{d}^{Cs}$ could still reach 2.66 $\times$ 10$^5$ mL/g with $R^{Cs}$ of 72.71% when the Sr/La/Cs molar ratio was 13.6:8:45.1, whereas $K_{d}^{Sr}$ and $K_{d}^{La}$ were only 4.37 mL/g and 50.7 mL/g with low $R^{Sr}$ and $R^{La}$ of 0.43% and 4.83%, respectively (Fig. 5h). And when the Sr/La/Cs molar ratios were 1.47:0.925:1 and 13.6:8:45.1, $SF_{CSr/La}$ retained 249.89 and 610.01, respectively. Even in the 3 mol/L HNO$_3$ solution with high concentrations of La$_3$$^+$ and Sr$_2$$^+$ (the Sr/La/Cs molar ratio is 123:73.7:1), $R^{Cs}$ could reach 19.1% which is significantly higher than Cs$^-$.

![Fig. 4 Selective capture of Cs$^-$ by InSnS-1 in the coexistence of Na$^+$ and Cs$^-$](image)

**Fig. 4 Selective capture of Cs$^-$ by InSnS-1 in the coexistence of Na$^+$ and Cs$^-$, $K_{d}^{Cs}$ and $R^{Cs}$ values of InSnS-1 in a neutral, b 1 mol/L HNO$_3$, and c 3 mol/L HNO$_3$ solutions with different Na/Cs molar ratios. Detailed experimental data can be found in Supplementary Table 15.**

![Fig. 5 Selective capture of Cs$^-$ by InSnS-1 in the presence of Sr$_2$$^+$ and/or La$_3$$^+$](image)

**Fig. 5 Selective capture of Cs$^-$ by InSnS-1 in the presence of Sr$_2$$^+$ and/or La$_3$$^+$, $K_{d}$ and $R$ of Cs$^-$ and Sr$_2$$^+$ ions removed by InSnS-1 in a neutral, b 1 mol/L HNO$_3$, and c 3 mol/L HNO$_3$ solutions with different Sr/Cs molar ratios. $K_{d}$ and $R$ of Cs$^-$ and La$_3$$^+$ ions removed by InSnS-1 in d neutral, e 1 mol/L HNO$_3$, and f 3 mol/L HNO$_3$ solutions with different La/Cs molar ratios. $K_{d}$ and $R$ of Cs$^-$, Sr$_2$$^+$, and La$_3$$^+$ ions removed by InSnS-1 in g neutral, h 1 mol/L HNO$_3$, and i 3 mol/L HNO$_3$ solutions with different Sr/La/Cs molar ratios (A = 1.38:1.27:1, B = 13.6:8:45.1, C = 123:73.7:1, D = 1.47:0.925:1, E = 13.6:8:45.1, F = 122:73.7:1, G = 1.45:0.880:1, H = 13.5:8.36:1, I = 123:73.7:1). Detailed experimental data can be found in Supplementary Tables 16–18.
than R\(^{\text{Sr}}\) and R\(^{\text{La}}\) (5.17% and 0.47%) (Fig. 5i). The above results indicate that the selectivity of InSnS-1 for Cs\(^+\) is higher in acidic solution compared to in neutral solution.

Clearly, in neutral solutions with low ion concentrations InSnS-1 retains high removal efficiencies for all of Cs\(^+\), Sr\(^{2+}\), and La\(^{3+}\) in both individual competing ion system and mixed Sr\(^{2+}\), La\(^{3+}\), and Cs\(^+\) system (Fig. 5a, d, g). However, in acidic solutions, K\(_d\)\(^{\text{Cs}}\) and R\(_d\)\(^{\text{Cs}}\) remain high and far exceed those of La\(^{3+}\) and Sr\(^{2+}\) ions even when the concentrations of La\(^{3+}\) and Sr\(^{2+}\) ions are much higher than that of Cs\(^+\). Besides, the increase in Na\(^+\), Sr\(^{2+}\) or La\(^{3+}\) concentration has less effect on Cs\(^+\) capture by InSnS-1 in strongly acidic solutions (1 and 3 mol/L HNO\(_3\)) solutions than in neutral solutions. Similarly, we obtained similar results in Cs\(^+\) selectivity experiments with different K/Cs or Ca/Cs molar ratios (Supplementary Fig. 19, Supplementary Tables 19 and 20) as well as in actual water sample experiments (Supplementary Note 3).

Such results demonstrate that the presence of H\(_3\)O\(^+\) is very favorable for enhancing the selective trapping ability of InSnS-1 for Cs\(^+\). This could also be attributed to the repulsive effect of H\(_2\)O\(^+\) compared to Na\(^+\), Sr\(^{2+}\), and La\(^{3+}\). H\(_2\)O\(^+\) has a small radius and is positively charged. Sr\(^{2+}\) and La\(^{3+}\) also have relatively small radii\(^{24}\) and highly positive charges compared to Cs\(^+\). Na\(^+\) has a smaller radius than Cs\(^+\), although it has the same positive charge as Cs\(^+\). Therefore, H\(_2\)O\(^+\) has a relatively stronger repulsive effect on Na\(^+\), Sr\(^{2+}\), and La\(^{3+}\) which prevents them from entering the structure. Previous studies on the Cs\(^+\) ion exchangers with high acid resistance mainly focused on the exchange capacity of materials. By contrast, this work provides the first systematic study of the effect of divalent and trivalent ions on the selective capture of Cs\(^+\) under acidic conditions.

**Desorption and cycling experiments.** To investigate how well the material can be recycled, desorption and recycling experiments were conducted. The 0.2 mol/L KCl solution or 1 mol/L HNO\(_3\) solution can effectively desorb Cs\(^+\) from InSnS-1-Cs (Supplementary Figs. 21 and 22). Due to the large adsorption capacity of InSnS-1 for Cs\(^+\) in 1 mol/L HNO\(_3\) solution, a small number of Cs\(^+\) were retained in the sample, but this problem could be solved by replacing the fresh HNO\(_3\) solution several times. However, to ensure the accuracy of the eluent concentration in the cyclic experiments, the HNO\(_3\) solution was not renewed during the desorption process. The experimental results show that the R\(^{\text{Cs}}\) can be maintained after three cycles, and the desorption rates are all about 100% and the layered structure of the material was maintained (Supplementary Figs. 23 and 24). The experimental results show that although part of Cs\(^+\) retaining in the structure affects the desorption, the newly adsorbed Cs\(^+\) can be completely desorbed in each cycle. The results also indirectly suggest that the acidified material (i.e., the material after desorption in nitric acid solution) still possesses the removal capacity for Cs\(^+\).

**Column experiments.** The breakthrough curves of Cs\(^+\) in the ion-exchange columns filled with InSnS-1 have been carried out under neutral and acidic conditions to evaluate the potential of InSnS-1 for industrial applications. The color of the adsorbent (InSnS-1 crystals) was darkened after treatment under neutral condition while it was lightened under acidic condition (Fig. 6a). The breakthrough curves were fitted to the Thomas model\(^{12}\) (correlation coefficient R\(^2\)>0.99) with a maximum adsorption capacity of 216.06 mg/g under neutral condition and 50.50 mg/g (Supplementary Table 28) under acidic condition (1 mol/L HNO\(_3\)). Usually, a value of C\(_t\)/C\(_0\) equal to 0.05 is used as a penetration point in industrial applications, taking into account the volume of wastewater treated\(^{25}\). For neutral and 1 mol/L HNO\(_3\) solutions with high concentrations of Cs\(^+\) (190.13 and 84.25 mg/L, respectively), the treatment volumes to reach the breakthrough points are approximately 1300 and 650 bed volumes, respectively (Fig. 6b, c).

InSnS-1 exhibits high dynamic adsorption capacity and large effective treatment volume for neutral solution containing high concentration of Cs\(^+\). Such excellent treatment capacity exceeds that of most reported Cs\(^+\) ion exchangers, e.g., MCC-g-AMP, M/SZMs, PEI/ZnFC\(^{54–63}\). Notably, hitherto studies on the column separation of Cs\(^+\) from strongly acidic solutions containing high concentration of Cs\(^+\) are still rare\(^{16,64}\). The current ion-exchange column provides a practical example that shows the effective removal of Cs\(^+\) from the highly acidic solution with high treatment amount. The above results demonstrate that InSnS-1 has great potential for practical application as a column packing in the field of radioesium remediation.

**Discussion**

In this work, the rapid and highly selective separation of Cs\(^+\) ions is achieved by the layered InSnS-1, especially for efficient capture of Cs\(^+\) even under highly acidic conditions. InSnS-1 as a Cs\(^+\) scavenger has the following main advantages: (i) excellent acid and irradiation resistances; (ii) fast adsorption kinetics and high adsorption capacities under neutral and acidic conditions; (iii) wide pH active range; (iv) effective separation of Cs\(^+\)/M\(^{\text{Sr}}\) (M\(^{\text{Sr}}\)=Na\(^+\), Sr\(^{2+}\), and La\(^{3+}\)) in acidic solutions; (v) easy desorption and reuse; (vi) effective dynamic separation of Cs\(^+\) as stationary phase in ion-exchange columns. The uptake mechanism of Cs\(^+\) and H\(_2\)O\(^+\) is directly visualized through single-crystal structural analyses. All promising features of InSnS-1 for the Cs\(^+\) capture originate from the extremely strong interactions between soft S\(^2–\) of sulfide layers and relatively soft Cs\(^+\), the adjustable interlayer spacing, and structural flexibility of InSnS-1 even in acidic condition. In addition, our study elucidates the non-negligible facilitating role of H\(_2\)O\(^+\) in the selective capture of Cs\(^+\) under acidic conditions. The important influence of the shortest K–S distance of K\(^+\)-directed metal sulfides on the ion-exchange performance is also revealed. This work highlights metal sulfides for the selective capture of Cs\(^+\) ion from strongly acidic nuclear wastewaters and sheds light on the design of novel acid-tolerant materials.
sulfide-based ion-exchange materials for radioisotope decontamination with practical applications.

Methods

Starting materials. K2CO3 (AR, General-Reagent), Sn (99%, damas-beta), CsCl (99.99%, Shanghai Longjin Metal Materials Co., Ltd.), CaCl2·2H2O (AR, Adamas Reagent Co., Ltd), SrCl2·6H2O (AR, Tianjin Guangfu Reagent Co., Ltd.), MgCl2 (AR, Adamas Reagent Co., Ltd). All the chemicals were used without further purification.

Synthesis of InSnS-1. In2S3 precursor was firstly synthesized by hydrothermal method. A mixture of K2CO3 (9.0 mmol, 1.2439 g), Sn (9.0 mmol, 1.0334 g), CsCl (0.30 mmol, 0.1961 g), and H2O (1.5 mL) was heated in a 28 mL polytetrafluoroethylene (PTFE) lined stainless steel autoclave at 220 °C for 2 days to obtain microcrystalline powder of In2S3. Then a mixture of In2S3 (3.0 mmol, 0.6541 g), Sn (3.0 mmol, 0.3561 g), Cs2CO3 (0.60 mmol, 0.1924 g) was sealed under vacuum in a quartz tube. Then the quartz tube was heated to 750 °C in 4 h at maintenance at 750 °C for 4 days, followed by program-controlled cooling to 550 °C for 3 days before turning off the furnace. After that, the quartz tube was naturally cooled to room temperature in the furnace. The crystals products were washed with deionized water and ethanol, and dried in 122.6 g of pure yellow fluffy crystals of InSnS-1 could be obtained, which are stable in water and air. The higher actual yield than the theoretical value (1.026 g) may be attributed to the water absorption of the compound.

Syntheses of three ion-exchange crystal products. 100 mg of InSnS-1 crystals were mixed with 100 mL neutral solution of 3000 mg/L Cs+ ions, 100 mL 1 mol/L nitric acid solution of 500 mg/L Cs+ ions, and 100 mL 1 mol/L HNO3 solution, respectively, which were shaken for 24 h with room temperature (RT). Then the crystals were washed with deionized water and ethanol, and dried naturally to afford the ion-exchange products formulated as CsInSnS6 (denoted as InSnS-1-Cs), CsIn2S3H (denoted as InSnS-1-Cs/H), and (H2O)InSnS2H2O (denoted as InSnS-1-H), respectively.

Characterizations. Single-crystal diffraction data for InSnS-1, InSnS-1-Cs, InSnS-1-Cs/H, and InSnS-1-H were collected with a SuperNova CCD diffractometer with graphite monochromated MoKα radiation. UV/visible spectra were obtained with a PerkinElmer Lambda 900 at RT. PXRD of solid materials were measured to confirm the phase purity. EDS, scanning electron microscope (SEM), and energy dispersive spectroscopy (EDS), scanning electron microscope (SEM), and elemental distribution mapping analysis were carried out through a JEOl JSM-6700F scanning electron microscope. X-ray photoelectron spectroscopy (XPS) analysis was carried out through a ESCALAB 250Xi spectrometer with AlKα radiation. UV/visible spectra were obtained with a PerkinElmer Lambda 900 at RT. Thermogravimetric analysis (TGA) was performed on a NETZSCH STA 449F3 DTAA–TG analyzer. Ion concentrations in solutions were measured by inductively coupled plasma-mass spectrometry (ICP-MS) or inductively coupled plasma optical emission spectroscopy (ICP-OES). ICP-MS and ICP-OES tests were carried out by Xserise II and Thermo 7400, respectively. InSnS-1 crystal samples were irradiated with γ-rays at a total dose of 100 kGy (1.1 kGy/h for 95 h) and 200 kGy (1.1 kGy/h for 181.8 h) to irradiation dose (2.5 milliCi units) provided by Detection Center of Suzhou CNNC Huadong Radiation Co., Ltd, China.

Acid stability experiments. In acid stability experiments, InSnS-1 crystals were immersed in different concentrations of HNO3 solutions for 10 h. After solid-liquid separation, the concentrations of leaching Sn and In in the solutions were determined by ICP-MS and the leaching percentages of Sn and In from the sample were inferred. PXRD of solid materials were measured to confirm whether the material framework remained stable. The equations and detailed descriptions used for data analysis can be found in Supplementary Note 2.

Batch ion-exchange experiments. A typical ion-exchange experiment of InSnS-1 was carried out in a 20 mL glass bottle containing an aqueous solution (V/m = 1000 mL/L) of CsCl and InSnS-1 powder (obtained by grinding InSnS-1 crystals). The mixture was shaken in a shaker for 4 h at RT and then centrifuged for separation. The supernatant was filtered and diluted for ICP testing and the solid was washed several times with deionized water and anhydrous ethanol before drying. The equations and detailed descriptions used for data analysis can be found in Supplementary Note 2.

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Author contributions
The manuscript was written through contributions of all authors. J.T., M.F., and X.H. conceived the project. J.T. designed and performed most of the experiments and analyzed the data. J.J. offered help in data analyses. W.L., X.Z., W.M., J.L., Y.P., and T.L. offered help in the partial experiments. J.T. prepared the manuscript. M.F. and X.H. revised the manuscript. M.F. supervised the project.

Competing interests
The authors declare no competing interests.

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