Insights into transport velocity of colloid-associated plutonium relative to tritium in porous media

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Although faster transport velocities of colloid-associated actinides, bacteria, and virus than nonreactive solutes have been observed in laboratory and field experiments, some questions still need to be answered. To accurately determine the relative velocity \( \frac{U_{Pu}}{U_T} \) of \(^{239}\text{Pu} \) and tritium representative of the bulk water, a conceptual model of electrostatic interactions coupled with the parabolic water velocity profile in pore channels is developed. Based on the expression for \( \frac{U_{Pu}}{U_T} \) derived from this model, we study the effects of water flow rates and ionic strengths on the \( \frac{U_{Pu}}{U_T} \). Also, the velocity relationship between Pu, tritium and \( \text{Sr}^{2+} \) is explored. The results show that \( \frac{U_{Pu}}{U_T} \) increased fairly linearly with decreasing water flow rates; \( \frac{U_{Pu}}{U_T} \) declined approximately exponentially with increasing \( \text{Na}^+ \) concentrations; the charge properties of colloid-associated Pu (negative), tritium (neutral) and \( \text{Sr}^{2+} \) (positive) had a close association with their transport velocities as \( U_{Pu} : U_T : U_{Sr}^{2+} = 1.41 : 1 : 0.579 \).

Because of long half-life and high radiotoxicity of \(^{239}\text{Pu} \), its mobility is a key concern involved in both performance assessments of high-level radioactive waste repositories\(^{1,2} \) and remediation of contaminated fields such as nuclear test sites\(^{3,4} \). Transport velocity of Pu (\( U_{Pu} \)) is one necessary parameter for both evaluating its mobility and then predicting its potential risks to human health. Organic and inorganic colloids with the diameter of 1 nm to 1 \( \mu \text{m} \) are ubiquitous in underground and vadose zone water\(^{5-8} \). The Pu(IV) having great sorption affinity for the colloids could thus exist in a stable species by strong association with functional groups on the colloidal surfaces\(^{9-13} \). Accordingly, \( U_{Pu} \) is a direct reflection of mobilization of the colloid-associated Pu in the geosphere.

More recently, colloid-associated Pu, Am, and Np were observed to have transport velocities 1–5% faster than tritium (\( U_T \)) in porous media\(^{14-18} \). The tritium (tritiated water) is a neutral tracer and its transport may be representative of bulk water velocities. In field and column experiments, model colloids such as bacteria\(^{19-21} \), virus\(^{22-24} \), and nanoparticles\(^{25,26} \) were also found to travel much faster than anionic ions (e.g. \( \text{Cl}^- \) and \( \text{Br}^- \)). This velocity enhancement was commonly explained by size exclusion\(^{27} \) and matrix diffusion\(^{28} \) mechanisms. In brief, colloid-associated actinides and/or colloids have relatively low diffusion in pore channels because of colloid-sized effects, and hence are excluded from much less mobile water domains near the pore walls where the water velocities are lower. The pore channels include both the pore spaces formed adjacent to grain–grain junctions and the intra-grain pores larger than the colloid sizes.

These velocity enhancement phenomena were widely reported. There are, however, still some questions that remain unanswered. One is how to determine velocity relationship between Pu (or other actinides) and tritium, i.e. the velocity enhancement factor \( \frac{U_{Pu}}{U_T} \), since the previously observed relationships were considered as rough estimates\(^{14-18} \). Typically, the ratio of two locations with the highest effluent peaks in breakthrough curves for colloidal contaminants and solute tracers was generally specified as the factors\(^{14,19,20,29,30} \). Such locations were usually plotted by time or effluent volumes vs. concentrations. The velocity relationship was thus established according to the only two single samples collected successively from the effluents\(^{14-20} \), and then might have large deviations from the true factors. The locations of the highest peaks are, in fact, difficult to be targeted sometimes. For example, in the case of slightly faster velocities of contaminants than solutes\(^{19,20} \), the two specified locations most often met at the almost same time. For long-pulse experiments, the effluent concentrations in the steady limbs of the breakthrough curves exhibited nearly identical peak heights\(^{20,30,31} \). In such cases, it is hard to judge which two single data should be used for indicating their relative velocity relationship. In this study, a simple
conceptual model of electrostatic interactions coupled with the parabolic water velocity profile is developed to more accurately determine the $U_{Pu}/U_{T}$ by using all breakthrough data instead of the two single.

The other question is whether environmentally relevant conditions, typically including the variations in water flow rate and ionic strength, influence the $U_{Pu}/U_{T}$. The pore water flow rates functioned as a physical mechanism may dominate colloid-associated Pu and tritium dispersion$^{23-34}$ that is directly related to their transport pathways in porous media systems$^{25-28,32,35}$. The lower dispersion of colloidal contaminants than that of the solutes$^{23,27,30}$ can lead to an inconsistent diversity of their pathways. In addition, the variation in this physical condition may give rise to the changes in mobile and immobile water contents in pore water$^{41-43}$. Accordingly, the various $U_{Pu}/U_{T}$ is expected to arise. On the other hand, the ionic strength as a chemical mechanism may control the electric double layer repulsion$^{44-45}$ between natural colloids as carriers of Pu and pore walls.

The natural colloids and media, such as soils and granites, usually have negatively charged surfaces under natural aquatic conditions. This repulsive interaction may hence alter the transport domain of colloid-associated Pu in pore channels but not the neutral tritium. It is well known that the water velocity has the parabolic distribution across the section of the channels. Therefore, the change in $U_{Pu}/U_{T}$ with ionic strengths would possibly occur. Until now, enough attention has not been paid to the dependence of $U_{Pu}/U_{T}$ on these two environmentally relevant conditions.

Moreover, since the colloidal forces have the potentials to alter the transport domain of Pu, the electrostatic attraction between positively charged ions (e.g. Sr$^{2+}$) and media surfaces may regulate the transport domain of Sr$^{2+}$ and then the $U_{Pu}/U_{T}$. Strontium-90 as one fission production from nuclear activities has been large-scale released into subsurface environments$^{46-47}$. It is necessary to determine accurately its relative transport velocity by our new model rather than the two single data method.

The aims of this work are to develop one conceptual model for establishing the velocity relationship of $U_{Pu}/U_{T}$ to investigate the role of water flow rates and ionic strengths in $U_{Pu}/U_{T}$, and to obtain the relative velocity of $U_{Pu}: U_{Pu}/U_{T}$, $U_{T}$. The mineral colloids with an average diameter ($d_{a}$) of 460 ± 78 nm were isolated from sandy soils collected at the vadose zone of Lop Nor and used for preparing the colloid-associated Pu. The sandy soil grains (300 to 700 μm) are used as the porous media packed into the columns.

Results

Electrokinetic properties of colloids and media. The ζ-potentials shown in Fig. S1 were determined according to the measured electrophoretic mobilities. Negative values of the ζ-potentials are an indication of negatively charged surfaces on the soil colloids (460 ± 78 nm shown in Fig. S2) and media (elemental composition shown in Table S1). The negative surface charge emanates from acidic functional groups$^{46}$, dominating the mineral surfaces under environmentally relevant pH 8.5 conditions. The charge properties of the mineral surfaces became less negative with increasing Na$^{+}$ concentrations. This suggests that the amount of variable negative charge decreased while introducing the cationic Na$^{+}$ to the suspensions. Large differences in ζ-potentials of the colloid and media surfaces were not observed, due to the fact that they were derived from the same in situ sandy soils.

Relative transport velocity of colloid-associated Pu and tritium as affected by water flow rates and ionic strengths. The results of Pu oxidation state distribution as determined via solvent extraction (Fig. S4) show that Pu(IV), both in solution and on the surfaces, was the predominant oxidation state (>95%) in the suspensions, and that minor changes of the distribution were not observed in the effluents. The high Pu(IV) percentage was presumably due to the reduction of initial Pu(VI) in the stock solution to Pu(IV) by the redox-sensitive Fe and Mn on the colloid surfaces, as also observed by many researchers$^{40,46}$. The relative transport of Pu and tritium in the porous media as affected by the water flow rates (Q) is shown in Fig. 1.

Normalized effluent concentrations of Pu (C/C$_{0}$) were increased with the Q, suggesting that hydrodynamic shear acting on the colloid-associated Pu was an important process to promote colloid-facilitated transport of Pu. Notably, Pu concentration fronts appeared ahead of tritium. Accordingly, Pu(IV) was transported in the colloidal species but not the dissolved species through the porous media. This was further confirmed by the colloidal fraction of Pu (> 0.995) in the effluents of the peak positions. This fraction is nearly identical to the C$_{p}$/C$_{0}$ = 0.996 of the initial Pu suspensions (see Fig. S5 at Na$^{+}$ = 0.002 mol/L).

Since Pu eluted earlier than the tritium, the consistent collection percentages (Cp) of Pu and tritium could not meet at the same locations with the effluent volumes (V) but at the two separate V$_{p}$ and V$_{Pu}$ with V$_{p}$ > V$_{Pu}$. The collection percentage is defined as, $C_{p}$ = (cumulative effluent mass at one effluent volume/the total effluent mass) × 100%. In the other word, the time for Pu arriving at one specified Cp was earlier than that for tritium. The difference in their effluent volumes, i.e. $V_{s} = V_{p} - V_{Pu}$ designated as the separation volumes, may be experimentally determined by using the breakthrough data in Fig. 1.

Because of high recovery percentages (Rp ≈ 100%) for Cl$^{-}$ and tritium as inert tracers fed into the columns, the difference in their V at two separate locations where their Rp(s) were overall 50% was used to determine their $V_{s}$ by Krupp et al.$^{11}$. In contrast to the inert tracers, Pu had significantly lower C/C$_{0}$ (<< 1) and thus extremely small Rp (e.g. the maximal Rp = 0.324% at Q = 4.56 mL/min). Hence we determine the $V_{s}$ between Pu and tritium at the two locations with the effluent volumes of $V_{T}(C_{p}=50\%)$ and $V_{Pu}(C_{p}=50\%)$ but not with the $V_{T}(R_{p}=50\%)$ and $V_{Pu}(R_{p}=50\%)$:

$$V_{s} = V_{T}(C_{p}=50\%) - V_{Pu}(C_{p}=50\%)$$

(1)

The dash lines in Fig. 1 show the $V_{T}(C_{p}=50\%)$ and $V_{Pu}(C_{p}=50\%)$. For the inert tracers, their total mass ($m_{I}$) injected into the columns are almost equal to their total effluent mass $m_{b}$ because of Rp ≈ 100% so that the specified V with the Cp = 50% can yield also Rp = 50% resulted from Rp = ($m_{b} \times 50\%$)/$m_{I}$ = 50%. Therefore, Eq. (1) is considered as a common expression suitable for the reactive as well as the inert tracers.

Plutonium exclusion volumes $V_{ex-Pu}$, i.e. the separation volumes per unit mass of the media, are determined as

$$V_{ex-Pu} = \frac{V_{s}}{m}$$

(2)

where m (45.0 g) is the mass of porous media packed into one column.

The water content in the exclusion domain (R - r$_{1}$ domain in Fig. S6), i.e. the exclusion water content $\theta_{ex-Pu}$, is calculated as

$$\theta_{ex-Pu} = \frac{V_{ex-Pu}}{\pi \times a^{2} \times H}$$

(3)

where a and H are the radius of the column (1.0 cm) and the height of the media (10.0 cm).

The velocity relationship between colloid-associated Pu and tritium can be derived from a conceptual model of electrostatic interactions coupled with the parabolic water velocity profile (Fig. S6), and thus is found to exist as

$$U_{Pu}/U_{T} = 1 + \frac{\theta_{ex-Pu}}{\theta}$$

(4)

where $U_{Pu}$ and $U_{T}$ are the average transport velocity of Pu and tritium; $\theta$(cm$^{2}$/cm$^{3}$) is the measured water content in the column.
The derivation of Eq. (4) from the conceptual model developed in this study is presented in the Supplementary Information.

The experimental conditions and the determined parameters of \( V_S \), \( V_{ex-Pu} \), \( h_{ex-Pu} \), and \( U_{Pu}/U_T \) are reported in Table 1.

The separation volumes \( V_S \) may further express the exclusion domain that was totally inaccessible to the colloid-associated Pu, ascribed to the repulsive interaction mechanism. In contrast, tritium could diffuse into the entire domain in pore channels and therefore its transport velocity \( (U_T) \) is indicative of bulk pore water velocities. As shown in Table 1, the \( V_{ex-Pu} \) increases with decreasing \( Q \), so that the colloid-associated Pu was forced into smaller spaces surrounding the center axis where the pore water velocity was highest. This might cause a fast transport velocity of Pu relative to the bulk pore water, i.e. \( U_{Pu}/U_T \) > 1. Furthermore, the relative velocity of \( U_{Pu}/U_T \) is increased approximately linearly from 1.35 to 1.52 with the \( Q \) from 4.56 to 0.472 cm\(^3\)/min, as shown in Fig. 2a. The higher \( U_{Pu}/U_T \) can be attributed to the continually compressed transport domain closer to the center axis.

The \( \theta_{ex-Pu} \) was related to immobile water (layer), since the exclusion domain was inaccessible to the colloid-associated Pu. The immobile water content for the tritium \( (h_{im-T}) \) as often determined by the Two-region Model) should be \( \theta_{im-T} < \theta_{ex-Pu} \) due to its molecule diffusion faster than the Brownian diffusion of colloid-associated Pu into the immobile water domain. Gvirtzman.

Table 1 | The experimental conditions and their effects on separation volumes \( (V_S) \), exclusion volumes \( (V_{ex-Pu}) \), exclusion water content \( (h_{ex-Pu}) \), and relative transport velocity of Pu and tritium \( (U_{Pu}/U_T) \)

| Flow rate, \( Q \) [cm\(^3\)/min] | Na\(^+\) [mol/L] | Water content, \( \theta \) [cm\(^2\)/cm\(^3\)] | \( V_S \) [mL] | \( V_{ex-Pu} \) [cm\(^3\)/g] | \( h_{ex-Pu} \) [cm\(^2\)/cm\(^3\)] | \( U_{Pu}/U_T \) |
|-----------------|-----------------|-----------------|-------|-----------------|-----------------|-----------------|
| 4.56            | 0.002\(^a\)     | 0.372           | 4.06  | 0.0903          | 0.129           | 1.35            |
| 3.37            | 0.002           | 0.345           | 4.16  | 0.0925          | 0.133           | 1.38            |
| 2.28            | 0.002           | 0.306           | 4.42  | 0.0983          | 0.141           | 1.46            |
| 0.472           | 0.002           | 0.280           | 4.56  | 0.101           | 0.145           | 1.52            |
| 2.28            | 0.01            | 0.306           | 3.86  | 0.0857          | 0.123           | 1.43            |
| 2.28            | 0.17            | 0.306           | 1.92  | 0.0428          | 0.0613          | 1.21            |
| 2.28            | 0.96            | 0.306           | 1.36  | 0.0302          | 0.0432          | 1.14            |

\(^a\)This concentration was produced while preparing Pu suspensions by the introduction of NaOH to adjust the pH to 8.5.

The dash lines indicate the effluent volumes where the collection percentages \( (C_p) \) of Pu (black) and tritium (red) are overall equal to 50%. The difference in their effluent volumes (blue line areas), i.e. \( V_S = V_T(C_p=50\%) - V_{Pu}(C_p=50\%) \), is the separation volumes \( (V_S) \). The initial tracer-level concentrations of Pu \( (C_0) \) = 10\(^-9\) mol/L. The error bars resulted from the standard deviations of \( C_0 \) (three samples) and from the measured relative standard deviations (RSD) of \( C \).
indicated the existence of $\theta_{\text{an}} < \theta_{\text{ex-noon}}$ for the anionic ions such as $\text{SO}_4^{2-}$ and $\text{Cl}^-$, resulting from their exchange model. Evidently, $\theta_{\text{an}} < \theta_{\text{ex-Pu}}$ was an indication that Pu was forced into the more centre streamlines, so that $U_{\text{Pu}}/U_T > 1$ was observed. This is consistent with the results determined via our conceptual model. In general, the immobile water content $\theta_{\text{im}}$ tends to increase with the decreases in $Q$ and/or $\theta^{41-43,53}$, as followed by the $\theta_{\text{ex-Pu}}$ in Table 1. In comparison, the increasing trend in $\theta_{\text{ex-Pu}}$ potentially prevailed in terms of the slower Brownian diffusion. This thus led to rising $U_{\text{Pu}}/U_T$ with decreasing $Q$, as shown in Fig. 2a.

Size exclusion (i.e. hydrodynamic chromatography) thoroughly discussed during the past several decades$^{23,27,30,54}$ seems to also account for the velocity enhancement phenomena. This colloid-sized effect, involving preferential exclusion of the colloid-associated Pu from smaller, more tortuous pore channels, could cause straighter transport pathways for the Pu. Therefore, the relatively fast velocity of Pu was a result of the lower extent of its pathway diversity. Nevertheless, size exclusion, emphasizing on the transport pathway diversity but not the transport domain in pore channels, is suitable for examining the relative movement of neutral particles rather than the charged species whose transport domain can be altered by surface forces. In addition, the colloidal particles and media inevitably exhibit the heterogeneous size distributions in natural environments. In such cases, it is difficult to establish the velocity relationship by the size exclusion mechanism. These aspects have been considered with the variable separation volumes in our conceptual model.

The increase in $U_{\text{Pu}}/U_T$ with decreasing $Q$ was a result of the increased $\theta_{\text{ex-Pu}}$. Here, $\theta_{\text{ex-Pu}}$ depends on this physical condition. Then, what will happen to the $\theta_{\text{ex-Pu}}$ if the change in chemical conditions occurs in pore water? Further transport experiments were performed under the constant physical conditions but different Na$^+$ concentrations in the initial Pu suspensions, as shown in Fig. 3. It is noted that the size exclusion mechanism could not explain the dependence of $U_{\text{Pu}}/U_T$ on the variation in Na$^+$ concentrations. In this study, we can determine the $U_{\text{Pu}}/U_T$ by the Eq. (4) and other parameters by the equations from (1) to (3). The results are provided in Table 1.

It is observed in Fig. 3 that the normalized effluent concentrations of Pu still appeared ahead of tritium. This indicates that the relative transport characterized by $U_{\text{Pu}}/U_T > 1$ could also take place in spite of the constant $Q$. The $V_S$ is decreased from 4.42 to 1.36 mL with Na$^+$ concentrations (0.002 to 0.96 mol/L) and the corresponding $V_{\text{ex-Pu}}$ from 0.0983 to 0.0302 cm$^3$/g. It is evident that Na$^+$ concentrations had an effect on the Pu transport domain in pore channels.

The electrostatic interaction between the two negatively charged surfaces of the media and the colloids may be considered as the sum of two contributions, including the van der Waals attraction $\Phi_{\text{vdW}}$ and the electric double layer repulsion $\Phi_{\text{edl}}$. (The calculations are presented in Supplementary Information). As shown in Fig. 4, the $\Phi_{\text{vdW}}$ is independent of the ionic strength in pore water, whereas the repulsion ($\Phi_{\text{edl}}$) markedly decreases with the increase in Na$^+$ concentrations, regardless of the same interaction distances. The decreased repulsion could make the colloid-associated Pu closer to the pore walls, and thus higher Na$^+$ concentrations resulted in lower $U_{\text{Pu}}/U_T$. The $U_{\text{Pu}}/U_T$, which declines approximately exponentially with increasing Na$^+$ concentrations, is presented in Fig. 2b. On the other hand, the relative high repulsion at lower Na$^+$ concentrations could result in transport of more colloid-associated Pu through the systems (see C/C$_0$ in Fig. 3).

No studies have been carried out to determine the quantitative velocity relationship between Pu and tritium under these environmentally relevant conditions (Q and ionic strength). Incorporation of their velocity relationship into the widely used transport model$^{55-57}$ is necessary to evaluate the environmental risk posed by the low-solubility actinides into the subsurface.

Relative transport velocity of colloid-associated Pu, tritium, and Sr$^{2+}$. The results by 10 kD ultrafiltration indicate that the Sr(NO$_3$)$_2$ in the Pu suspensions mainly existed as the dissolved species (~98.2%), i.e. the positively charged Sr$^{2+}$ ions$^{36,59}$. The relative transport of colloid-associated Pu, tritium, and Sr$^{2+}$ in one porous media system are displayed in Fig. 5. As expected, Sr$^{2+}$ appeared behind tritium, revealing that the transport velocity of Sr$^{2+}$ was slower than that of tritium. Then, their transport velocities may be expressed as $U_{\text{Pu}} > U_T > U_{\text{Sr}}$.

Due to the attractive electrostatic interaction between Sr$^{2+}$ and pore walls, Sr$^{2+}$ could be attached onto the walls, and thus be excluded from the center domain in pore channel. This domain (illustrated by $r_2$ in Fig. S6) reflects the separation volumes $V_S$ between Sr$^{2+}$ and tritium, which is opposite to the $V_S$ between colloid-associated Pu and tritium.

$$V_S = V_{\text{Sr}^{2+} (C_p=50\%)} - V_{T (C_p=50\%)}$$

where $V_{\text{Sr}^{2+} (C_p=50\%)}$ and $V_{T (C_p=50\%)}$ are the effluent volumes of Sr$^{2+}$ and tritium.

The exclusion volumes $V_{\text{ex-Sr}^{2+}}$ and exclusion water content $\theta_{\text{ex-Sr}^{2+}}$ also determined by Eq. (2) and (3), respectively, are reported.

Figure 2 | Responses of the relative transport velocity ($U_{\text{Pu}}/U_T$) to (a) water flow rate and (b) Na$^+$ concentration in pore water. The error bars resulted from the measured RSDs of the effluent concentrations ($C$) of Pu and ‘H, and the experimentally determined water content ($\theta$).
in Table 2. The relative transport velocity of $U_{Sr}^{2+}/U_T$ is established as

$$U_{Sr}^{2+}/U_T = 1 - \frac{\theta_{ex-Sr}^{2+}}{\theta}$$

(6)

The derivation of Eq. (6) based on the conceptual transport model is presented in the Supplementary Information.

As a result of the electrostatic attraction, $Sr^{2+}$ ions stayed in the domain near the pore walls, and the exclusion domain surrounding the center axis had the water content of $h_{ex-Sr}^{2+} = 0.129$ cm$^3$/cm$^3$. The transport velocity of $Sr^{2+}$ relative to that of tritium behaved as $U_{Sr}^{2+}/U_T = 0.579$. The $Sr^{2+}$ tailing can be attributed to desorption (detachment) of the $Sr^{2+}$ previously attached onto the pore walls, and then these desorbed $Sr^{2+}$ slowly moved in the domain near the walls. Recently, matrix diffusion of the dissolved species, such as $Cl^-$, $I^-$, $Cs^+$, and tritium in porous fractures was intensively studied, and the consistent results indicated the larger and smaller diffusion for the cations and anions, respectively, as compared with the neutral tritium$^{60-62}$. This carries an implication of slower transport velocity of the cationic $Sr^{2+}$ than tritium, which is further demonstrated by $U_{Sr}^{2+}/U_T = 0.579$ resulted from our conceptual model allowing for the electrostatic attraction.

The electrostatic interactions between contaminants and media surfaces may exhibit as repulsion or attraction, depending on the charge properties of the contaminants. In this study, the charge properties have proven to have a close association with their
transport velocities. The observed relationship between the colloid-associated Pu, tritium and Sr$^{2+}$ is quantitatively determined as $U_{Pu} : U_T : U_{Sr^{2+}} = 1.41 : 1 : 0.579$.

**Discussion**

We developed a new conceptual model of electrostatic interactions coupled with the parabolic water velocity profile in pore channels. This model is much more suitable for determining the transport velocities of the charged contaminants. Some studies were carried out by using the expressions for $U_{Pu}/U_T$ and $U_{Sr^{2+}}/U_T$ derived from this model. The results show that,

1. The relative velocity of colloid-associated Pu and tritium ($U_{Pu}/U_T$) increased fairly linearly with decreasing water flow rates, resulting from the continually compressed transport domain of Pu close to the center streamlines;

2. The $U_{Pu}/U_T$ declined approximately exponentially with increasing Na$^+$ concentrations because of the decreased electrostatic repulsion between the natural colloids as carriers of Pu and the pore walls;

3. The charge properties of colloid-associated Pu (negative), tritium (neutral) and Sr$^{2+}$ (positive) had a close association with their relative transport velocities, which was determined as $U_{Pu} : U_T : U_{Sr^{2+}} = 1.41 : 1 : 0.579$ under the experimental conditions of water flow rate $Q = 2.27 \text{ mL/min}$, water content $\theta = 0.306 \text{ cm}^3/\text{cm}^3$, Sr$^{2+} = 0.0065 \text{ mol/L}$, and Na$^+ = 0.01 \text{ mol/L}$.

The previously reported transport velocities of the colloid-associated Pu, Am, and Np were ~1–5% faster than tritium$^{14–18}$, as determined by only two single data in breakthrough curves. This rough estimate possibly caused lower velocities than the $U_{Pu}/U_T$ (1.14–1.52, depending on the experimental conditions in Table 1) determined by the newly developed model in this study. Our results do fall into the ranges observed from the field experiments in the aquifers, such as 1.25 times (bacteria/Br$^-$)$^{13}$, 1–2 times (bacteria/groundwater)$^{63}$, 1.5–2.5 times (bacteria/Br$^-$)$^{23,64}$, and 1.25 times (bacteria/groundwater)$^{63}$. Because filed experiments of actinides remain scarce, these bacteria transport may be analogous to the colloid-associated Pu by following considerations. Both bacteria and colloid-associated Pu had the nano-scale sizes and the negatively charged surfaces, and therefore they exhibited similar transport mechanisms such as size exclusion and repulsive interaction.

**Methological methods**

Poros media. Sandy soils collected in northwestern China (Lop Nor) were air-dried, sieved with a stainless steel sieve. The elemental composition of the soils determined by X-ray diffraction (D/MAX-2500) are 50% quartz, 15% amorphous, 11% sericite (clay minerals), 6% orthoclaste, 5% calcite, 4% dolomite, 4% clay minerals, 2% amphibole, 1% gypsum, 1% pyrite and 1% undetected minerals. The elemental composition determined by X-ray fluorescence spectroscopy (Axios) is presented in Table S1. Other characteristics were a cation exchange capacity of 0.0286 mmol/g and pH of 8.5 ± 0.1 (1:1 water-media).

Natural colloids. The natural colloids were isolated from the sandy soils. The sieved soil sample of 150 g (< 0.3 mm in diameter) was added to a glass beaker (4 L pure water), and ultrasonically dispersed for 15 min. The colloid suspensions with Stoke's

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**Table 2** | Separation volume $V_S$, exclusion volume $V_{ex}$, exclusion content $\theta_{ex}$ and relative transport velocity between both Pu/tritium and tritium/Sr$^{2+}$ under the conditions of water flow rate $Q = 2.27 \text{ mL/min}$, water content $\theta = 0.306 \text{ cm}^3/\text{cm}^3$, Sr$^{2+} = 0.0065 \text{ mol/L}$, and Na$^+ = 0.01 \text{ mol/L}$.

| Pu  | Sr$^{2+}$ | Relative transport velocity |
|-----|----------|-----------------------------|
| $V_S$ (mL) | $V_{exPu}$ (mL/g) | $\theta_{exPu}$ (cm$^3$/cm$^3$) | $V_S$ (mL) | $V_{exSr^{2+}}$ (mL/g) | $\theta_{exSr^{2+}}$ (cm$^3$/cm$^3$) | $U_{Pu}/U_T$ | $U_{Sr^{2+}}/U_T$ |
| 3.93 | 0.0873 | 0.125 | 4.05 | 0.0899 | 0.129 | 1.41 | 0.579 |
diameter of < 1 μm were siphoned from the upper suspension into a polypropylene vessel and then stored in the refrigerator (4 °C), used as the colloidal source materials. Because the isolated colloids were derived from the sandy soils, their elemental composition was identical to the reported results of the media (see Table S1).

To determine the mass concentration of the parent suspensions, 25 mL aliquots of the suspensions were transferred to concave Teflon membranes, respectively, and dried at 50 °C by infrared light. The determined concentration was 267 ± 6 mg/L. The parent suspensions could be diluted with pure water to obtain the desired colloidal concentrations.

Characterization of colloids and media. Both Brunauer Emmett Teller (BET) surface areas (A) and pore volumes (V) of the porous media (300–700 μm in diameter) were measured using the N₂ adsorption method at 77 K (ASAP 2020, Micromeritics). Adsorption/desorption curves of N₂ are shown in Fig. S3, and the measured results are given in Table S2.

The porous media of about 5 g were crushed in an agate mortar. The crushed particles larger than 10 μm in diameter were removed using gravity sedimentation.

Electrokinetic potentials (ζ) of the remaining particles < 10 μm and the colloids, as a function of pH, were determined from ζ potential concentration of 0.001 to 1.0 mol/L were determined using Nano ZS (Malvern) under the conditions of pH 8.5, and Na⁺ = 0.002 mol/L. The experimental results are presented in Fig. S1.

The colloidal diameter in the prepared suspensions was measured with the dynamic light scatter (Nanosizer ZS, Malvern) under the conditions of Na⁺ = 0.002 mol/L, pH = 8.5, and salt concentration = 100 mg/L. The result in Fig. S2 shows that the average diameter (dₐ) was 460 ± 78 nm, and that the colloidal suspensions could remain relatively stable.

Pu suspensions. Small amounts of 239Pu stock solution (95 mg/g) in 1.5% HNO₃ were drop by drop added to the stirred colloidal suspensions in Teflon bottles. The atom ratio in the solution was 239Pu²⁺/239Pu⁴⁺ = 0.13, 0.27, 0.54, 1.08, 2.16, and 2.31 for Pu suspensions containing 15, 30, 60, 90, 120, and 140 μg/g Pu respectively. After 0.5 h of the transport experiments, 2.2 mL of 8 mol/L HNO₃, 3 mL of 10 mol/L HCl, and 3 mL of 3 mol/L HNO₃. The purified Pu was in the feeding solution were removed via washing the columns using 5 mL of 8 mol/L HNO₃ concentration was 800 Bq/g) was added to the Pu suspensions for studying the relative transport of Pu, tritium and Sr²⁺. The Na₂SO₄ was added into one aliquot of Pu suspensions into a polypropylene vessel and then stored in the refrigerator (4 °C), used as the colloidal source materials. Because the isolated colloids were derived from the sandy soils, their elemental composition was identical to the reported results of the media (see Table S1).

Analytical methods. Plutonium oxidation state distribution, both in solution and on the colloidal surfaces was determined by a combination of centrifugal ultrafiltration (10 kD, Amicon Ultra-4, Millipore) and solvent extraction (TTA and DHEHP, Sigma-Aldrich Co.) according to the analytic flow diagram in Fig. S4. Plutonium-239 concentrations of all samples were measured by ICP-MS using the isotope dilution method, as follows. Plutonium-242 was used as a spike, and the atom ratio of Pu²⁺/Pu⁴⁺ was determined introducing known amounts of 239Pu in the samples. HNO₃ was added to adjust the solution to 8 mol/L HNO₃, NaNO₃ was used to adjust Pu valence to Pu(V) at 90 °C for 15 min, and then the solution cooled at room temperature. These solutions were fed into Dowex® × 2 resin (Dow Chemical Co.) packed in glass columns with 3 mm × 50 mm length. U and matrix elements in the feeding solution were removed via washing the columns using 5 mL of 8 mol/L HNO₃, 3 mL of 10 mol/L HCl, and 3 mL of 3 mol/L HNO₃. The purified Pu was eluted from the columns with 1.2 mL of 0.01 mol/L HNO₃, 0.01 mol/L HF solution, and determined with the ICP-MS.

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Author contributions
J.C.X. isolated the natural colloids and carried out most of the transport experiments. J.C.L. helped plan the experiments, and participated in the preparation of the manuscript. J.F.L., S.P. and R.M. helped plan the experiments, and participated in the preparation of the manuscript. J.F.L., S.P. and R.M. helped plan the experiments, and participated in the preparation of the manuscript. J.F.L., S.P. and R.M. helped plan the experiments, and participated in the preparation of the manuscript.

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