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Uptake of anionic radionuclides onto degraded cement pastes and competing effect of organic ligands

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Cement / Degradation / Nuclear waste / Radionuclides / Organic / Uptake

Summary. Hardened cement pastes (HCP) present a high affinity with a lot of radionuclides (RN) and can be used as waste confining materials in radioactive waste repository. Indeed, in cementitious media, RN can be removed from solution via (co)precipitation reactions or via sorption/diffusion mechanisms. In this study, the affinity of anionic RN (Cl⁻, I⁻, SeO₃²⁻ and CO₃²⁻ chemical forms) with a CEM-I HCP has been studied vs. the degradation of the HCP particles. These RN are considered as mobile in repository media and it is important to have a set of distribution ratio (R_d) in cement environment. The R_d values have been measured in batch experiments as a function of the pH, used as the degraded state parameter of the HCP suspensions. The $R_{\rm d}$ values increase in all cases, from the unaltered state (pH 13.3) to the altered state of HCP, i.e. until all portlandite is dissolved, corresponding to pH 12.6. Then, R_d values decrease until degraded states (pH 12.0), corresponding to the decalcification of the calcium silicate hydrate (C-S-H) phases. The behaviour of anionic RN seems to be correlated to the evolution of calcium concentration and is opposed to the evolution of sulphate concentration in solution which could have a competing effect. Comparison is done with the behaviour of caesium and uranium(VI), which is a cationic RN but has a major negative hydrolysed species at high pH. As awaited, the uranium(VI) behaviour is very different from purely anionic RN one in accord with spectroscopic analyses from literature works. The $R_{\rm d}$ values have also been measured for the organic ligands isosaccharinate (ISA) and EDTA. The uptake of ISA can be important and competing effect with the sorption of SeO₃² has been evidenced in HCP suspensions as a function of the ISA concentration.

1. Introduction

In the French design of a deep geological radioactive waste repository, clay media seem favourable host rocks for repositories as they have interesting properties for confinement (low permeability, high radioactive waste retention capacity ...). Mortar and concrete could be used as container or buffer, and backfill materials of engineered barrier systems between the wastes and the geological medium, at least for B-wastes, which are long-lived intermediate level wastes mainly generated by the facilities associated with the nuclear-fuel cycle and by research establishments. Cement materials will contribute to the retardation of dissolved radionuclides (RN) by the combination of physical transport and chemical interactions. The chemical interactions may retard mobile RN and lower transport rates. Then, it is important to quantify the sorption of RN with hardened cement pastes (HCP).

One of the alkaline cement materials drawbacks is their high leaching rates, which lead to the alteration of the material [1]. In deep clay geological formations, as the permeability of the rock is very low, cement pastes would be much less degraded as the underground water removal will be very slow. Nevertheless, more important leaching phenomena could occur in some cases as in case of borehole crossing accidentally the repository or as in sub-surface radioactive waste repository concepts. Then, it is necessary to quantify the uptake rates of RN not only in the case of unaltered HCP but also for degraded HCP.

The uptake of actinides, lanthanides, and cationic fission and activation products is high in cement environments and is well documented in literature [2, 3, and references therein]. Anionic RN behaviour in cement environments has been less investigated as these RN are more mobile in clay and cement materials than cationic RN. Then, their confinement concept is mostly based on the anionic exclusion properties during the diffusion through clay media. But as ³⁶Cl, ¹²⁹I and ⁷⁹Se are the main contributors to the final dose in performance assessment calculations [2], it seems important to evaluate the possible contribution of cement materials to the retardation of these RN in radioactive waste repository. 14C was also studied in this study (as the carbonate form ¹⁴CO₃²⁻) because mortar and concrete materials could also be used in sub-surface graphite waste storage, issued from natural uranium graphite gas nuclear power plants.

In a previous study, we have measured the evolution of the zeta potential (ζ) of different HCP vs. their degradation

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states [4]. The pH of the equilibrium solution was considered as the degradation parameter. Two pH_{IEP} (pH of the isoelectric point) were observed with 12.9 and 11.7 average values. Then, it could be of interest to study the affinity of anionic mobile RN vs. the degradation of HCP and its correlation with the evolution of ζ .

A comparison can be done with the behaviour of cationic RN in order to consolidate the observation for the anionic RN. Caesium is used as a pure cationic RN, as it is only present in cement water under the Cs⁺ form and because ¹³⁷Cs is a relevant RN for performance assessment [2]. Uranium(VI) is also studied as it is a cationic RN, as UO₂²⁺, but is supposed to have major negative hydrolysed species at high pH. For UO₂²⁺, the objective is to complete previous results [5] in order to check whether its anionic hydrolysed species have similar behaviour than pure anionic one in cement environment because few points measured on three degraded states in [5] suggest that another comportment occurs.

Organic products are also present in B-wastes and could have an important effect on the immobilization of RN in HCP media, because of their ability to form complexes with cationic RN, enhancing their solubility or mobility in the cement pore water. They also could be sorbed onto the HCP surface with a competing effect on anionic RN uptake. The influence of organics such as isosaccharinic acid (ISA), the most important degradation product of cellulose, on the behaviour of cationic fission products and actinides has already been studied in cementitious solution and materials [6–11]. The sorption of ISA on undegraded HCP surface has already been observed and measured [12], and its competing effect on anionic species sorption onto HCP has already been observed [13]. In the present study, this effect has been investigated through batch sorption experiments with selenite (SeO₃²⁻) in HCP suspensions as a function of ISA and EDTA concentration. Henceforth, the behaviour of anionic organic species with the degradation of HCP has also been measured.

2. Materials and methods

2.1 Cementitious suspensions preparation

CEM-I, an Ordinary Portland Cement, purchased from "Ciment d'Origny" manufacturer, was used. The experimental details of the clinker composition, the cure of the HCP (water/cement ratio of 0.38) and the degradation protocol of the paste are detailed in [4]. Briefly, a four years old HCP, setting in saturated portlandite water, has been crushed and different masses of the sieved powder have been put in contact with milliQ water and/or an artificial fresh cement water. The artificial fresh cement water was synthesized dissolving NaOH, KOH and Ca(OH)2 in demineralised degassed water. The pH of this water is about 13.3, and its composition is close to calculated pore water composition of undegraded cement pastes [1]. These equilibrations allow getting about twelve degradation states between pH 11.5 and 13.3. The batches were carried out in 30 mL polysulfone (PSF) centrifuge tubes (with polypropylene screw closure). All preparations (weighting, batch preparation, and sampling) were performed in a glove

box under nitrogen atmosphere to prevent carbonation of the suspensions ($p_{\rm CO_2} < 1$ ppm). After two weeks of equilibration under tri-dimensional agitation, the tubes were ultracentrifuged (50 000 g for 1 h) and 1 mL of solution was sampled and acidified with an Ultrapur HNO₃ solution. Then, total concentrations of calcium, chloride and sulphate have been measured by ionic chromatography (Dionex DX-120). The pH values were measured in the supernatant solution of the batch with a combined microelectrode (Mettler Inlab, electrolyte filling KCl 3 M/AgCl saturated) calibrated at pH 7.01 (phosphate buffer as KH₂PO₄/Na₂PO₄) and 12.65 (saturated solution of portlandite) at 20 °C. The uncertainty of pH measurement was estimated to be 0.05.

2.2 Organic ligands

EDTA salt, ethylenediaminetetraacetic acid disodium dihydrate – $C_{10}H_{14}N_2O_8\cdot 2H_2O$, was purchased from Fluka (ref. 03679) with purity higher than 99%. Ca(ISA)₂ was synthesised by lactose alkaline degradation in saturated lime water, as described in [14]. The NaISA form was obtained by treating a solution of dissolved Ca(ISA)₂ (20 g of solid in 2 L of Milli-Q water) through a column of cation exchange resin (CHELEX 100, converted to the Na⁺ form with 0.2 M NaOH solution). Characterisation data of Ca(ISA)₂ solid and dissolved NaISA are available in [13].

2.3 Batch sorption experiments

After sampling for calcium, chloride and sulphate analyses, the batches were spiked with radioactive solutions. Series of batch sorption experiments were performed for chloride (Cl⁻), iodide (I⁻), selenite (SeO₃²⁻) and carbonate (CO₃²⁻). Series were also performed with two cationic RN, caesium (Cs⁺) and uranium (UO₂²⁺). Lastly, a series was done with ISA and two batches were also performed to measure the uptake of EDTA. After equilibration under tri-dimensional agitation for one month (Cl, I, Se, and C), or for two weeks (Cs, U, ISA and EDTA), the suspensions were ultracentrifuged at 50 000 g for 1 h. The supernatants were analysed by liquid scintillation counting in order to determine the residual amount of radioactive tracer or by total organic carbon (TOC-5000, Shimadzu) analysis for ISA and EDTA. Results are expressed as distribution ratio R_d values, which correspond to:

$$\begin{split} R_{\rm d}({\rm mL/g}) &= \frac{[{\rm RN}]_{\rm sorbed}({\rm mol/g})}{[{\rm RN}]_{\rm solution}({\rm mol/mL})} \\ &= \frac{[{\rm RN}]_{\rm initial} - [{\rm RN}]_{\rm solution}}{[{\rm RN}]_{\rm solution}} \times \frac{{\rm solution~volume~(mL)}}{{\rm solid~mass~(g)}} \,. \end{split}$$

The mass of solid used for the R_d calculation is the dry mass determined by ignition at 1000 °C. The same method is applied for the R_d calculation of organic ligand.

Solutions of Na³⁶Cl, Na¹²⁵I, Na₂⁷⁵SeO₃ and Na₂¹⁴CO₃ were used in batch experiments. Average total concentrations of elements before sorption, taking into account carriers, are equal to: $[Cl^-] = 7.6 \times 10^{-8} \text{ M}$, $[I^-] = 1.4 \times 10^{-8} \text{ M}$, $[SeO_3^{2-}] = 2.2 \times 10^{-8} \text{ M}$, and $[CO_3^{2-}] = 10^{-7} \text{ M}$. The initial concentration of CO_3^{2-} is lower than the calculated solubility of carbonate in a saturated portlandite water

(about 6×10^{-6} M). The calculation has been performed with PHREEQC [15].

²³²U (UO₂Cl₂ in HCl 2 M) is used as tracer. In order to avoid the interferences related to the formation of ²²⁸Th and its daughters, the experimental time is limited to 3 weeks for U(VI) sorption experiments. The stock solution of ²³²U is purified with a BIORAD AG-1X 8 200–400 mesh resin.

A 137 CsCl solution was used and added in the batch experiments with an average concentration before sorption equal to 10^{-9} mol L^{-1} .

The initial concentrations of the organic ligands in the batches were [ISA $^-$] = 8.5×10^{-4} M and [EDTA $^-$] = 1.1×10^{-3} M.

3. Results and discussions

3.1 Cementitious materials characterisations

The cementitious minerals quantifications of the undegraded CEM-I HCP were performed in [4] by differential scanning calorimetry coupled to thermogravimetry (DSC/TG) analysis and by Rietveld's calculations of X-Ray diffraction (XRD) diffractogramms. The analyses shown that portlandite (Ca(OH) $_2$, 22% 1), ettringite ([Ca $_3$ Al(OH) $_6$ ·12H $_2$ O] $_2$ (SO $_4$) $_3$ ·2H $_2$ O, 4.7% 1) and calcite (CaCO $_3$, 4.3% 1) are the major crystallised cementitious phases. C-S-H phases were also detected (mass estimated to 50%), but due to their nano-crystalline structure, only large peaks between 3.05 and 2.83 Å were observed. The evolution of the cement pastes mineralogy with leaching was not analysed in this study and the degradation states were only characterised through the chemical composition of the equilibrium solutions.

3.2 Evolution of the chemical compositions of the cement waters vs. degradation

The calcium concentration has been measured as it has a strong influence on the uptake of anionic species, allowing the formation of a global positive charge near the surface of cementitious grains when its concentration is sufficiently high. We have already measured and calculated the ζ evolution of the same CEM-I HCP as a function of its degradation state [4]; two pH_{IEP} have been identified (see gray line in Fig. 1). Indeed, ζ increased from -17 to +20 mV for pH decreasing from 13.2 to 12.6 (undegraded HCP states), corresponding to the increase of calcium concentration from 1.1×10^{-3} to 2×10^{-2} M. Then, ζ decreases from +20 to -8 mV for pH 12.6 to 11.0 (degraded HCP states), corresponding to the decrease of calcium concentration from 2×10^{-2} to 4×10^{-4} M. Hence, ζ is highly dependent on calcium concentration and for both pH_{IEP}, the calcium concentration value is about $2-4 \times 10^{-3}$ M. The evolution of the calcium concentration measured in the present study is reported in Fig. 1. One can see that for this sample the first pH_{IEP} was also attempted to be about 12.9 and the second about 11.7.

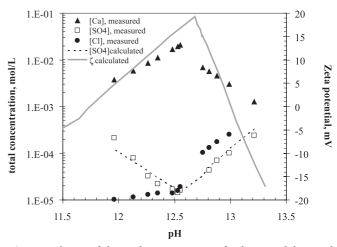


Fig. 1. Evolution of the total concentrations of calcium, sulphate and chloride as a function of pH in the suspensions of CEM-I HCP powder in contact with demineralised and decarbonated water and/or artificial fresh cement solution. The error bars of the measured concentrations are contained in the size of the symbols. The evolution of $[SO_4^{2-}]$ resulting from the modelling is also reported. ζ values are issued from [4].

The concentrations of sulphate have been measured as it could have a competing effect on anionic RN uptake. Indeed, the sorption of ${\rm SO_4}^{2-}$ on the surface of C-S-H phases has been measured in [16]. The evolution of sulphate monitored in Fig. 1 is well represented by the solubility of ettringite calculated with the PHREEQC code, using the mean solubility product of:

$$\begin{split} & \text{Ca}_{6}[\text{Al}(\text{OH})_{6}]_{2}(\text{SO}_{4})_{3} \cdot 26\text{H}_{2}\text{O} \rightleftharpoons \\ & 6\text{Ca}^{2+} + 2\text{Al}(\text{OH})_{4}^{-} + 3\text{SO}_{4}^{2-} + 4\text{OH}^{-}, \log K = -44.9 \,, \end{split}$$

determined in [17].

From pH 13.2 to 12.6, the evolution of the sulphate concentration would be in the favour of the uptake of anionic RN whereas from pH 12.6 to 12.0, the evolution of sulphate concentration might be unfavourable to the sorption of anionic RN.

Chloride concentration has also been measured in order to determine the total chloride concentration in $^{36}\text{Cl}^-$ batch sorption experiments. The behaviour of chloride is different from sulphate one as the concentration decreases regularly with the alteration of the HCP. The solubility of the chloride is probably controlled by the Friedel's salt (Ca₂Al(OH)₆(Cl, OH) · 2H₂O) as reported in [18]. Nevertheless, chloride has a high solubility and it is difficult to estimate the initial amount of chloride that is present in the HCP as the cement pastes have previously been stored in a saturated portlandite solution where the chloride may have been dissolved and diffused.

3.3 Behaviour of anionic RN in cement pastes suspensions vs. degradation

The sorption kinetics of $^{75}\text{SeO}_3^{2-}$, monitored by R_d evolution vs. time, are reported in Fig. 2. For the three degradation states (corresponding to pH 13.2, 12.4 and 11.9), long times are required to reach the equilibrium state, almost 30-50 days. This result suggests that the uptake of Se(IV)

¹ The quantity represents the percent of mass of the mineral in HCP.

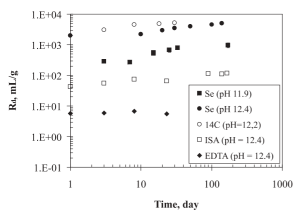


Fig. 2. The uptake kinetic of $^{75}SeO_3^{2-}$, $^{14}CO_3^{2-}$, ISA and EDTA by HCP CEM-I is represented at different degradation states. Kinetic R_d values of $^{75}SeO_3^{2-}$ reported at the degraded state pH 12.4 are issued from [25].

is not only a sorption mechanism on the cement minerals surface but, in addition, this phenomenon could be followed by a diffusion mechanism in the cement minerals structure. 30 days has been selected as the contact time used in the following batch sorption experiments.

Bonhoure *et al.* [19] have also observed long equilibration times for the uptake of iodide in unaltered HCP suspensions. The authors have measured that, even after 180 days, the R_d values were still increasing: the R_d values were increased by a factor 2 between 7 days and 180 days. No kinetic study of $^{36}\text{Cl}^-$ sorption has been performed so far in the literature, and the same contact time of 30 days has been applied for the batch sorption experiments.

For $^{14}\text{CO}_3^{2-}$ the longest studied time is only 30 days (see Fig. 2) but the major part of the tracer seems sorbed. This experimental time is lower than the one reported in [20], where an increase of the R_d values until 100 days is reported. The uptake mechanism of $^{14}\text{CO}_3^{2-}$ is still puzzling. Bayliss *et al.* suggested that the uptake mechanism was probably an isotopic exchange between ^{14}C and stable carbon of calcite initially present in the materials [20], whereas Noshita *et al.* [21] have studied the sorption of $^{14}\text{CO}_3^{2-}$ on cement minerals and have suggested that it can also sorb on >SiOCa $^+$ site on C-S-H surface.

It could be noticed that the main goals of this study was to obtain operational experimental $R_{\rm d}$ values for a large panel of altered and degraded HCP states. Then, even if the contact time (30 days) is sometimes lower than the required time of equilibrium state, the $R_{\rm d}$ values can still be used as conservative values for performance assessment calculations.

The measured R_d values of $^{36}\text{Cl}^-$, $^{125}\text{I}^-$, $^{75}\text{SeO}_3^{2-}$, and $^{14}\text{CO}_3^{2-}$ are reported in Fig. 3 as a function of the degradation of the HCP. R_d values measured in literature are also reported [19, 22–24]. The evolution trends of R_d values vs. pH are similar for all the RN and two major features can be discriminated during the degradation of the HCP.

Between pH 13.2 and 12.6, when portlandite is still present, $R_{\rm d}$ values increase and, globally, the affinity order is $^{14}{\rm CO_3}^{2-} > ^{75}{\rm SeO_3}^{2-} > ^{125}{\rm I}^{-} > ^{36}{\rm Cl}^{-}$. For the degraded states under pH 12.6 (total leaching of portlandite), $R_{\rm d}$ values slightly decrease. The affinity order of RN for the highly degraded HCP is the same than for the first part of the degradation. It is worthy to notice that the $R_{\rm d}$ values measured

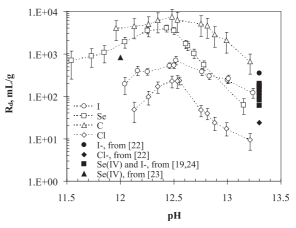


Fig. 3. Evolution of the R_d values of $^{125}I^-$, $^{14}CO_3{}^{2-}$, $^{75}SeO_3{}^{2-}$ and $^{36}Cl^-$ in degraded CEM-I HCP suspensions as a function of pH.

for the more degraded state (pH 12) are always higher than the $R_{\rm d}$ values measured for the unaltered state (pH 13.2), which could be used as a conservative value in performance assessment calculations.

This behaviour is well correlated to the evolution of calcium concentration. It seems that the development of positive charge at/near the negative surface charge of HCP allows the uptake of anionic RN. Furthermore, the R_d evolution of anionic RN agrees with the evolution of the concentration of the competing SO_4^{2-} specie, as R_d and $[SO_4^{2-}]$ are inversely correlated.

Few spectroscopic studies have been performed in literature in order to identify the chemical environment of the sorbed anionic RN in cement materials. Bonhoure et al. [24] have studied by X-ray absorption fine spectroscopy (XAFS) the uptake of Se(IV) oxy-anion by unaltered CEM-I HCP and cement minerals (C-S-H, ettringite ...). They suggested that the coordination sphere of the SeO₃²⁻ entity was maintained upon immobilisation by HCP and non-specific interactions (outer-sphere surface complex) dominate at the given selenite loadings. Macé [25] has measured the R_d values of Se(IV) on different cement minerals (C-S-H, ettringite and hydrogarnet) and has observed that the uptake of Se(IV) in HCP suspension is controlled by aluminate phases (hydrogarnet and ettringite) rather than C-S-H phases that are minor contributors in the immobilization of Se(IV). Toyohara et al. [26, 27] have identified that a hydrogarnet phase (4CaO·Al₂O₃·13H₂O) controls the uptake of iodide in cement. Bonhoure et al. [19] only obtained extremely weak EXAFS oscillations from I⁻ treated unaltered CEM-I HCP and CSH samples, which has limited the interpretation of the EXAFS data. These results suggest that the uptake of ³⁶Cl⁻, ¹²⁵I⁻ and ⁷⁵SeO₃²⁻ are the result of electrostatic interactions with the HCP surface, and that no specific chemical adsorption occurs.

3.4 Behaviour of cationic RN in cement pastes suspensions vs. degradation

The kinetic sorption of uranium(VI) has already been studied in [5], with the same cement materials, until 21 days. The measured R_d values between 9 days and 21 days were similar, suggesting that 9 days was the minimum time re-

quired to reach a steady state. In the present study, a contact time of 15 days has been used. The same time has been used for caesium.

The uptake of caesium in cement materials has already been measured in literature [28, and references therein]. Its behaviour is opposed to the behaviour of the anionic RN as $R_{\rm d}$ always increase from unaltered states to highly degraded state, particularly between pH 12.6 and 11.5, see Fig. 4. Between pH 13.2 and 12.6, the R_d values did not decrease as Na⁺ and K⁺ concentrations are very important and compete with Cs^+ sorption, preventing from high R_d values in undegraded HCP states where ζ values are negative. The behaviour of U(VI) was also investigated. The objective of this experiment was to investigate the behaviour of a positive cation which has negative species in cement water. Indeed, at pH 12, UO₂(OH)₃ and UO₂(OH)₄²⁻ are present at 97 and 3% respectively and at pH 13.2 at 51 and 49% respectively, see Fig. 5. The speciation calculations were estimated using PHREEQC (with Davies equation) and the data from [29]. Then, one could think that in cement environment, U(VI) would have the same behaviour than anionic RN. Nevertheless, the R_d evolution trend of U(VI) is close to Cs+ one's, see Fig. 4, and then different from purely anionic RN one. We could observe that during the first stage of the degradation, the $R_{\rm d}$ values are nearly constant (1000–4000 mL/g) and it is only when C-S-H phases control the solubility of calcium in the second part of the degradation that the $R_{\rm d}$ values highly increase (from 4000 to $200\,000\,\mathrm{mL/g}$), as for Cs⁺, whereas $R_{\rm d}$ values of purely anionic RN decrease in this pH range. In [5], batch sorption experiments results showed that R_d measured in C-S-H suspensions were very close to R_d values measured in HCP suspensions with similar Ca/Si ratio in the solids. Then, it can be supposed that C-S-H phases contribute to the major part of the U(VI) uptake in HCP. Harfouche et al. [30] have studied the uptake of U(VI) by C-S-H using XAFS. A split equatorial oxygen shell, neighbouring silicon atoms at short and long distances, and neighbouring Ca atoms were observed. The structural parameters resemble those reported for uranophane (Ca(UO₂)₂(SiO₃OH)₂·5H₂O). One can also remind that the operational solubility limits of U(VI) obtained in [5, 31] were comparable to the uranophane solubility calculated from thermodynamic data determined by [32]. It can be supposed that inner-sphere complex formation of

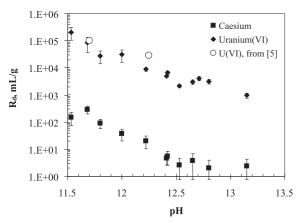


Fig. 4. Evolution of the R_d values of Cs and U(VI) in degraded CEM-I HCP suspensions as a function of pH.

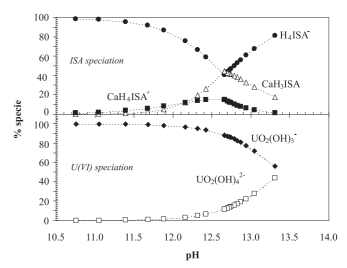


Fig. 5. Speciation of U(VI) and ISA in cement waters as a function of pH and the associated chemical compositions presented in Fig. 1.

U(VI) onto C-S-H is the predominant uptake mechanism by adsorption in a first time. Other mechanisms as solid-solution formation could occur after sorption, explaining high R_d values measured for U(VI). This uptake mechanism seems then very different from anions and oxy-anions, where literature spectroscopic investigations suggest the formation of outer-sphere complex to explain the uptake of these RN on HCP surface.

3.5 Behaviour of organic ligand in cement pastes suspensions *vs.* degradation

The kinetic of the sorption mechanism is fast and, after three days, the equilibrium state between the solid phase and the liquid phase is reached (see Fig. 2). The evolution of the uptake of ISA is similar to anionic RN one as a function of the degradation of HCP (see Fig. 6). Indeed, $R_{\rm d}$ values increased from 4 to 100 mL/g for pH decreasing from 13.2 to 12.5. Then, $R_{\rm d}$ decreases from 100 to 25 mL/g when pH decreases from 12.5 to 11.8.

Speciation calculations on Fig. 5, using data from [33], show that H_4ISA^- is the major species over the pH range 13.2–12.0; CaH_3ISA is the second species, with the highest contribution when the calcium concentration is highest (at pH = 12.6). The fact that the major species is anionic can explain the similar behaviour of ISA to anionic RN's. EDTA speciation is also dominated by anionic species [33]; 100% of $CaEDTA^{2-}$ in degraded HCP water (pH = 12); $EDTA^{4-}$ accounts for 88% of the speciation in unaltered HCP water (pH = 13.2). The evolution of EDTA affinity seems similar to the ISA one, even if only two R_d values have been measured. R_d values are $0.17^2 \pm 0.04$ and 11 ± 6 mL/g for pH 13.2 and 11.9 respectively, showing a lower affinity than ISA, which has R_d values of 5 ± 2 and 61 ± 21 mL/g, respectively.

As equilibration times are short, shorter than for inorganic anions, uptake of H₄ISA⁻ is probably only controlled

² The $R_{\rm d}$ value of EDTA in unaltered HCP suspension has been measured with a higher solid/liquid ratio than for other $R_{\rm d}$ measurement: s/l = 940 g/L, instead of 1–7 g/L.

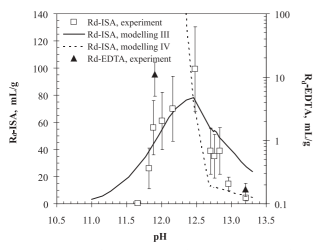


Fig. 6. Evolution of the $R_{\rm d}$ values of ISA and EDTA in fresh (pH = 13.3) and degraded (pH = 11.9) HCP suspensions. Calculated $R_{\rm d}$ values of ISA in HCP suspensions are also represented, using surface equilibria III and IV.

by surface mechanism and modelling attempts have been performed. As in [4], this modelling is based on the Diffuse Layered Modelling (DLM) from [34], a surface complexation modelling (SCM). Two surface equilibria have already been proposed for C-S-H surface chemistry [35, 36] and have also been used for HCP ζ calculation [4]:

 i) the ionization of the surface sites (>SOH), mostly through deprotonation:

$$>$$
SOH $\rightleftharpoons >$ SO $^- +$ H $^+$,
log $K = -12.0$, (equilibrium I)

and

ii) the sorption of Ca²⁺ on the silanol sites, bringing in addition a positive charge through the following equilibrium:

$$>$$
SOH + Ca²⁺ \rightleftharpoons $>$ SOCa⁺ + H⁺,
log $K = -9.2$, (equilibrium II)

The description of the specific adsorption of calcium onto C-S-H can be discussed as Monte Carlo simulations have shown that ordinary electrostatic interactions seem enough to explain the evolution of the effective surface charge of C-S-H particles [37]. Calcium cation overlaps the negative charge of the C-S-H surface by strong physical adsorption resulting from electrostatic interactions. This is why one prefers to talk about global positive charge on/near the surface of HCP and that the SCM proposed in this work has to be considered only as a way for predicting $R_{\rm d}$ values and does not pretend describing the sorption mechanism that occurs between HCP particles and the organic species. In this first modelling attempt, $H_4 \rm ISA^-$ is supposed to sorb on the surface sites > SOH or > SOCa $^+$ and two major equilibria could occur:

i) a surface complexation reaction, forming a mixed surface complex:

$$>$$
SOH + Ca²⁺ + H₄ISA⁻ \Longrightarrow SOCaH₄ISA + H⁺,
log $K = -7.5$, (equilibrium III)

or

ii) a ligand exchange reaction, forming a simple surface complex:

$$>$$
SOH + H₄ISA⁻ + H⁺ \rightleftharpoons
 $>$ SH₄ISA + H₂O . (equilibrium IV)

The thermodynamic calculations have been performed with PHREEQC. Modelling the sorption of H_4ISA^- with Ca^{2+} on a surface site give R_d values in good agreement with experimental R_d values; all the attempts done with a ligand exchange equilibrium were not satisfying (equilibrium IV). Indeed, when the surface equilibrium constant is adjusted in the case of an unaltered state, it is impossible to obtain the observed decrease of the R_d after the total leaching of the portlandite. Then, the uptake rate of ISA can only be quantified considering the formation of a mixed surface complex as the equilibrium III.

3.6 Competing effect of ISA on SeO₃²⁻ uptake in cement pastes suspensions

In this study, the influence of ISA and EDTA has been studied on the uptake of ⁷⁵SeO₃²⁻ by the undegraded HCP at pH 13.2 and degraded state of HCP at pH 12.0. Batch sorption experiments results (see Fig. 7) show that the uptake of ISA has a competing effect on the uptake of ⁷⁵SeO₃²⁻. Indeed, in unaltered HCP at pH 13.2 and degraded HCP at pH 12.0, the R_d values measured without ISA are divided by a factor of two when the ISA concentration is about 2×10^{-3} and 5×10^{-4} M, respectively. The proposed explanations of the $R_{\rm d}$ decrease in presence of ISA are i) a competing effect of the sorption of the negative species H₄ISA⁻ on the HCP surface and/or ii) the complexation of Ca²⁺ by ISA, forming CaH₃ISA and CaH₄ISA⁺ complexes, inducing a more negative charge near the surface of the HCP particles and preventing the formation of a mixed surface complex on HCP. Such a modelling attempt does not provide any argument to discriminate between the two hypotheses.

Data are also reported to show the effect of EDTA. For HCP at pH 13.2, no effect is observed in the limit of the highest studied organic concentration (10^{-3} M). An effect is only observed for HCP at pH 12.0 for the highest concentration and it is not possible to extrapolate the feature of the global influence of EDTA.

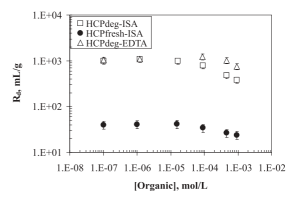


Fig. 7. Evolution of the R_d values of $^{75}\mathrm{SeO_3}^{2-}$ in fresh (pH = 13.3) and degraded (pH = 11.9) HCP suspensions as a function of ISA or EDTA concentrations. The R_d values have been measured here after 90 days of contact.

One can see that organic ligand present in the radioactive waste containers, or produced from this organic matter under alkaline environment as ISA, can have a competing effect on the anionic RN uptake and might be taken into account in the safety performance assessment calculations if their concentration is high.

4. Conclusion

A set of experimental distribution ratio (R_d) has been acquired for Cl-, I-, SeO₃²⁻ and CO₃²⁻ in degraded CEM-I HCP suspensions, as a function of the pH. These data could be used for migration calculation of ³⁶Cl, ¹²⁹I, ⁷⁹Se and 14C in cementitious barriers which could be used in radioactive waste containers and repository. The R_d values increase for the first part of the alteration of the HCP, until the portlandite is buffered (for corresponding pH 13.2 to 12.6). Then, the $R_{\rm d}$ values decrease until pH 12.0. This behaviour is well correlated to the evolution of the calcium concentration that controls the surface potential near the surface of the HCP particles. The behaviour of the anionic RN is specific and can not be extrapolated to the behaviour of cationic RN with anionic species which control their speciation in the pH range of cementitious materials. The organic ligand initially present or produced in the HLW containers, as ISA, can have a competing effect on the anionic RN uptake and might be taken into account in the performance assessment calculations.

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