Separation of Cations from Nitric Acid Solutions Using Commercially Available Ion Exchange Resins

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Abstract

In a previous publication, the authors described a chromatographic process for the selective separation of fission products and minor actinides from uranium and plutonium in nitric acid solution. This paper has evaluated commercially available ion exchange materials for capacity, rate of uptake and selectivity for some fission products (inactive) and cerium (iii) and (iv) as surrogate for Pu and U. The fission products studied where Cs, Sr and Zr as these for various reasons present major challenges for the existing PUREX process and waste management. The commercial resins evaluated were various sulfonic acid, chelate ion exchangers and an undisclosed inorganic material all supplied by Purolite Ltd.

Keywords: Reprocessing; PUREX; Ion chromatographic separation; Cesium; Strontium; Commercial resins

Introduction

The potential for extractive chromatography as a replacement and/or complementary reprocessing option to the PUREX process has been previously described [1]. UCLan’s process involves the separation of fission products and minor actinides from uranium and plutonium isotopes in 1 to 3 M nitric acid. This paper evaluates the potential of commercially available ion exchangers for the separation of Sr, Cs, Co and Zr ions from Ce (iii/iv) ions the surrogates for Pu and uranium [2]. These fission product cations were selected as:

1. Their chemistries and behaviour are different,
2. They account for a significant amount of β/γ activity present in spent fuel dissolver liquor,
3. Zr is responsible for significant challenges in the PUREX process [3].

Cerium ions (iii/iv) have been used as a surrogate for Pu in a variety of studies ranging from reprocessing, fuel fabrication to waste management [4]. The liquid-liquid extraction of cerium ions from nitrate solution using tri butyl phosphate was well established even before the conception of the PUREX process [5]. The Ce (iv) ion forms relatively weak nitrate complexes in nitric acid solution (~ 1 mole/l), with the Ce+4 ion predominating but with Ce(NO3)2+ and Ce(NO3)3+ ions increasing in stronger nitric acid [6].

Xe, Zr, Mo, Nd, Cs and Ru isotopes which are formed in the largest amounts in thermal fission, constitute about 70% of the fission product weight after a cooling time of ten years. Cesium and strontium and daughter isotopes account for about >90% of the total activity (Bq/t U) from fission products for 8-year cooled fuel, with a burn-up of 45 GWe/t (Table 1). At cooling times 10 to 1,000 years the activities of strontium-90 a strong β emitter with a half-life of 28.8 years and caesium-137 with a half-life of 30 years a strong β/γ emitter dominate among the fission products. They are the two most important fission products when considering reagent stability in a reprocessing flowsheet.

Although the only important oxidation number for zirconium is +4, the solution chemistry is complicated as a number of ionic species can be present depending on the solution conditions. Zirconium ions can undergo extensive hydrolysis to produce colloidal species and polymerisation, which are pH and concentration dependent [7].

Although there is an abundance of metal extraction data for the selected commercial ion exchangers most if not all are restricted to slightly acid, neutral to alkaline pH regions far removed from 1 to 3 M acid [8]. The influence of pH (acidity) is known to influence both extraction and separation efficiencies; in addition metal speciation and hydrated ionic radii will be affected [9,10].

UCLan’s chromatography process relies on the sequential separation of individual radionuclides or families of radionuclides using appropriate stationary phases. The envisaged process will initially remove the comparative short-lived radionuclides such as Cs and Sr isotopes (significant β/γ emitters) thus reducing the radiation damage to downstream stationary phases (Figure 1) [11]. Reduction in radiation flux will allow organic exchangers to be considered for the subsequent separation of other radionuclides in later stages.

A major, if not the major separation consideration for the stationary phase will be the ability to achieve decontamination values (D) comparable, if not better with those attained in the PUREX process for example 107 [12]. To achieve this figure would require the stationary phase to reduce say the Cs/Sr concentration from ~ 3 g/Kg U to 0.3 µg/Kg U, based on the data in Tables 1 and 2 [8].

The optioneering investigations reported in this paper are based on batch studies to identify candidate exchangers that will be further evaluated in column experiments. Some of the selected commercial ion exchangers exhibit good extraction efficiencies for copper; this metal was included in this study to provide a comparison between their intended commercial use and their performance in 1 to 3 M nitric acid conditions, i.e., benchmarking exercise [8,13].

Extraction efficiencies, separation factors and rate of cation uptake were calculated from equilibrium batch studies over a range of nitric acid molar strengths.

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and possible limiting factor on the separation of ions and efficiency of

Reagents and instrumentation
Purolite Ltd., provided a variety of resins and adsorbents; their
general properties are displayed in Table 3: 500 ppm stock solutions
were prepared of Cs, Sr, Ce (III), Ce (IV) Co and Cu nitrate salts in 1%
general properties are displayed in Table 3: 500 ppm stock solutions

Materials and Methods
All the reagents were of analar grade quality or equivalent with the
exception of nitric acid, which was of trace metal analytical grade. The
cation concentrations were determined using Thermo X-Series ICP-MS
with Thermo PlasmaLab version 2.5.11.321. The deionised water which
had a conductivity of 18.2 MΩ.cm used in these experiments was sourced
with Thermo Scientific Barnstead NANOPURE ion exchanger.

All equipment was routine standardised and appropriate standards
were used, in particular for ICP-MS determinations. Equilibration of cation
concentrations were determined using Thermo X-Series ICP-MS
- for batch uptake experiments; for the rate of uptake experiments, a

Batch uptake experiments
As the acid molar strength of the feed solution will be a determining
and possible limiting factor on the separation of ions and efficiency of
Table 1: Heat rating factor for some radioisotopes present in spent fuel dissolver liquor*.

<table>
<thead>
<tr>
<th>Radioisotope</th>
<th>Bq/t U</th>
<th>Heat rating factor (W/Bq)</th>
<th>Heat rating contribution (W/tU)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Y-90</td>
<td>3.14E+15</td>
<td>1.50E-13</td>
<td>4.70E+02</td>
</tr>
<tr>
<td>Sr-90</td>
<td>3.14E+15</td>
<td>3.13E-14</td>
<td>9.84E+01</td>
</tr>
<tr>
<td>Rh-106</td>
<td>8.90E+13</td>
<td>2.60E-13</td>
<td>2.31E+01</td>
</tr>
<tr>
<td>Sb-125</td>
<td>4.65E+13</td>
<td>8.54E-14</td>
<td>3.97E+00</td>
</tr>
<tr>
<td>Cs-134</td>
<td>5.02E+14</td>
<td>2.75E-13</td>
<td>1.38E+02</td>
</tr>
<tr>
<td>Cs-137</td>
<td>4.38E+15</td>
<td>2.98E-14</td>
<td>1.31E+02</td>
</tr>
<tr>
<td>Ba-137 m</td>
<td>4.14E+15</td>
<td>1.06E-13</td>
<td>4.39E+02</td>
</tr>
<tr>
<td>Ce-144</td>
<td>3.33E+13</td>
<td>1.78E-14</td>
<td>5.92E-01</td>
</tr>
<tr>
<td>Pr-144</td>
<td>3.33E+13</td>
<td>1.88E-13</td>
<td>6.06E+00</td>
</tr>
<tr>
<td>Eu-154</td>
<td>1.72E+14</td>
<td>2.42E-13</td>
<td>4.16E+01</td>
</tr>
<tr>
<td>Am-241</td>
<td>7.65E+13</td>
<td>9.04E-13</td>
<td>6.91E+01</td>
</tr>
<tr>
<td>Cm-242</td>
<td>9.93E+09</td>
<td>9.95E-13</td>
<td>9.88E-03</td>
</tr>
<tr>
<td>Cm-244</td>
<td>1.21E+14</td>
<td>9.46E-13</td>
<td>1.14E+02</td>
</tr>
<tr>
<td>Total</td>
<td></td>
<td></td>
<td>1.53E+03</td>
</tr>
</tbody>
</table>

*Above values for 45 GWd/t U burn up and 8-year cooling post reactor.

Table 2: Dissolver liquor concentrations*.

<table>
<thead>
<tr>
<th>Radionuclide</th>
<th>Approximate concentration</th>
</tr>
</thead>
<tbody>
<tr>
<td>U</td>
<td>~300 g/l</td>
</tr>
<tr>
<td>Pu</td>
<td>3.6 g/l</td>
</tr>
<tr>
<td>Np</td>
<td>170 mg/l</td>
</tr>
<tr>
<td>Am</td>
<td>225 mg/l</td>
</tr>
<tr>
<td>Cm</td>
<td>8 mg/l</td>
</tr>
<tr>
<td>Alkali metals (Cs, Rb)</td>
<td>1.3 g/l</td>
</tr>
<tr>
<td>Alkaline earth metals (Sr and Ba)</td>
<td>1.05 g/l</td>
</tr>
<tr>
<td>Y and lanthanides</td>
<td>4.4 g/l</td>
</tr>
<tr>
<td>Zr</td>
<td>1.5 g/l</td>
</tr>
<tr>
<td>Se and Te</td>
<td>220 mg/l</td>
</tr>
<tr>
<td>Mo</td>
<td>1.4 g/l</td>
</tr>
<tr>
<td>Tc</td>
<td>350 mg/l</td>
</tr>
<tr>
<td>Ru, Rh, Pd</td>
<td>1.8 g/l</td>
</tr>
<tr>
<td>Ag, Cd, Sn, Sb</td>
<td>60 mg/l</td>
</tr>
</tbody>
</table>

*Concentrations are based on a typical irradiated PWR 3.7 % U-235 fuel with a
burn up of 45GWd/t HM, cooled for 3 years.

Table 3: Adsorbents used and their functionality and polymeric properties.

<table>
<thead>
<tr>
<th>Resin Name</th>
<th>Functional Group</th>
<th>Structure</th>
</tr>
</thead>
<tbody>
<tr>
<td>100H</td>
<td>Sulphonic Acid</td>
<td>Gel Polystyrene with divinylbenzene at 8%</td>
</tr>
<tr>
<td>C100X10MBH</td>
<td>Sulphonic Acid</td>
<td>Gel Polystyrene with divinylbenzene at 10%</td>
</tr>
<tr>
<td>S910</td>
<td>Amidoxime</td>
<td>Polycrylic croslinked with divinylbenzene</td>
</tr>
<tr>
<td>D5530</td>
<td>Unknown</td>
<td>Unknown</td>
</tr>
</tbody>
</table>

any chromatographic separation, batch studies of the effect of nitric
acid strength (up to 4 M) on the uptake of ions by all adsorbents were
initially investigated. Experiments were carried out in 250 ml Durran
bottles with 100 ml of the required concentration of nitric acid; to this,
a requisite amount of the cation stock solution was added to bring the
cation concentration to 500 ppb. 1 g of the adsorbent was then added to
the bottle, sealed and placed in the water bath and agitated at 25°C for
24 hours (24 hrs. taken to approximate equilibrium). Percentage uptake
values were calculated and plotted against acid strength (Figures 2-4).

Selectivity co-efficient at the same acid concentration were calculated by Equation 1:

\[ S = \frac{\text{Percent uptake of ion } A}{\text{Percent uptake of ion } B} \]  

(1)

The amount of each ion adsorbed per g of resin was calculated by Equation 2:

\[ \mathcal{Q} = \frac{(C_i - C_f) \times V}{m} \]  

(2)

Where \(C_i\) is the initial ion concentration (mg/l), \(V\) the volume of
nitric acid solution (l), \(m\) the weight of resin (g) and \(C_f\) the concentration
at the end of the experiment taken as equilibrium (mg/l).

Distribution coefficients were calculated from Equation 3:

\[ K_{oe} = \frac{(C_i - C_f) \times V}{C_i \times m} \]  

(3)

Rate of cation uptake

Small samples were periodically withdrawn from the 3,000 ppm
Zr/2 M nitric acid solution (100 ml) equilibrating at the appropriate
temperature and subsequently analysed. It was considered that the Zr
ions had reached equilibrium with exchange materials after 24 hours
(C_e). Rates of Zr ion uptake as a function of time were calculated either
as a percentage of \(C_i\) or Zr mass uptake (mg/g).
Results and Discussion

Capacity and selectivity

Selectivity and distribution coefficients were determined for six ions on three adsorbents with four ions being tested with the S910. Only one sulphonic acid resin graph is displayed as the similarity between the sulfonic C100H and C100x10 MBH resins, is differentiated only by the amount of divinyl benzene cross-linking agent within the resin structure (Figure 2). It is appreciated however that the degree of cross-linking does affect both capacity and selectivity of sulphonic acid ion exchange resins but in this study the affect and implications were marginal [14].

Uptake was observed for most of the cations over quite a large nitric acid molar range for the sulfonic resins, with the exception of cesium, the adsorption of which falls off quickly as the acidity is raised above 1.0 M.

At this acid concentration, the dissociation of sulphonic acid will be slightly impaired, and as the nitric acid concentration increases, the dissociation of SO3H will be further reduced in order to maintain the equilibrium, viz:

R- SO3H ↔ R-SO3⁻ + H⁺ \[ pK_a \approx -2.8 \] [15]

Ce and Zr ions are capable of overcoming this lack of dissociation by displacing the hydrogen ion attached to the functional group. As the measured maximum uptake for these ions is of the order 50 µg/g this would entail only 10% dissociation of the sulphonic acid group for a 1:1 SO3⁻-Zr complex. The uptake of ions is higher for the more charged species, so zirconium (IV) > cerium (IV) > strontium (II) > copper and cobalt (II) > caesium (I); this sequence is consistent with previously measured selectivities for sulphonic acid resins [16]. Cerium as reported earlier and zirconium ions are capable of forming nitrate- complexes in nitric acid which may assist in the uptake of these ions by the sulphonic acid functional group, in particular for monovalent species such as Ce(NO3)3⁺.

The acid concentrations for 50% extraction for Cs, Co, Cu and Sr ions from Zr ions are 0.15, 0.4, 0.65, 1.40, 1.50 and 2.10 M respectively (Figure 2). These extraction profiles indicate that an effective separation of Cs, Co, Cu and Sr ions from Zr ions could be achieved in 2.0 M nitric acid, but the separation would be less efficient for Ce ions from Zr ions.

This separation factor trend for sulphonic acid resins is underpinned by the data reported in Tables 4 and 5. In 2 M acid, the separation factor for Cs ions from Zr ions is about 29 and nearly 20 and 16.4 for Co and Sr ions respectively for C100H. The performance of 100x MBH is slightly inferior with values of 22, ~ 10 and 6.5 respectively, which may be due to the higher degree of cross-linking.

The difference in the sulphonic acid distribution coefficients is interesting; although they gave similar shaped graphs, the more highly cross-linked resin gave nearly double the distribution coefficient. This may be due to the interesting chemistry displayed by zirconium speciation at low acid concentrations (ZrO2⁺, Zr⁴⁺ and [Zr₆(OH)₁₆(H₂O)₁₆]⁶⁺). The variation in charges, hydrated radii and ionic diameters displayed at these low nitric acid concentrations will have a dramatic effect on the adsorption on to the resin. The pore size of the resin, density of active sites on its surface and capacity will all be limiting factors affecting uptake.

Large ions such as the [Zr₆(OH)₁₆(H₂O)₁₆]⁶⁺ will be less able to penetrate the internal structures and active sites within a resin than that of smaller ion such as ZrO₂⁺ or Zr⁴⁺. It will also take more active sites on the resin to adsorb a 8⁺ charged hydrated ion than that of a 4⁺ or 2⁺ ion. This will effectively reduce the capacity of the resin for Zr. The relatively high cross-linking will reduce pore size within the resin, possibly making it too small for the hydrated ions, thus slowing the uptake. The ion does however behave as expected at the higher acid concentrations of most interest to this project, on both C100 resins.

The separation performance of the amidoxime resin for Zr ions from other cations is far superior to the sulphonic acid ion exchangers, for example at 50% extraction the acid concentrations are 0.1 and 0.2 M for Zr and Ce(IV) ions respectively (Figure 3). At these acid concentrations uptake of other cations is below 10%.

The distribution ratios recorded for Zr ions from 1 to 3M nitric acid solutions exceed 900 demonstrating the binding capacity of the amidoxime group (Tables 6 and 7).

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**Figure 1:** UCLan’s Continuous chromatography separation process.

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The reduction in uptake for the sulfonic resins can be described by the equilibrium between hydrogen ions on the resin, those in solution and the cation as previously explained. The S910 being a chelating resin is slightly different, it has the ability to complex with specific ions but reject others. The selectivity can be further enhanced by increasing acidity; this is demonstrated in the experiments where in deionised water the uptake of Ce (IV) ion is greater than that of Zr ion but is reduced close to zero once HNO$_3$ is above 1 mol.l$^{-1}$ [1].

In general hydroxamic acids are weak donors with the pKa values varying from 7.05 (nitrobenzohydroxamic acid) to 11.33 (N-phenyl-n-butylhydroxamic acid) [17], amidoxime resins to remove cations from acid solutions has been reported on several previous occasions [18-20]. Vernon studied various transition metals such as Ni, Co, Cu, Pb V, Fe and U, Hg and Au in near neutral to acid solutions (pH values 6.0 to 1.0) and reported that V and Au had the highest capacities of ~0.8 mmoles/g and ~2.5 mmoles/g respectively at zero pH value [21]. The resin also had a high uptake for copper but at pH value of 6 (3.2 mmoles/g). This high capacity for copper ions is also a characteristic of S910 (40 g/l equivalent to ~0.5 mmoles/g) [14].

There are extensive reviews of the stability constants and metal complexation properties of hydroxamic acids but with emphasis on benzo hydroxamic acid (BHA) complexes [22,23]. The metals of interest to UCLan’s chromatographic separation process that have been studied with BHA and AHA are Fe(iii), U(vi), Ru(iii), Zr(iv), Pu(vi) and rare earths [24].

The stability constants reported in Table 7 were measured in slightly...
acid to modest alkaline pH conditions i.e., ~2 to 10 unlike the nitric acid conditions employed in this study. Below pH values 1.0, the proportion of metal hydroxamate in solution will be small. This for most cations in our study is confirmed by the extraction-acid profiles in Figure 3, the exception is Zr ions. These profiles and the data in Table 7 would suggest that only mono-dentate or at worst bi-dentate ligands are formed.

The most interesting commercially available material supplied by Purolite Ltd., was D5530 that had strong uptake of cations from 1 to 3 M nitric acid (Figure 4). In 2 M, acid Zr ion was adsorbed totally with Cs, Sr, and Ce cations in the 60% region. At a higher acidity of 4M the separation was enhanced with Zr ion still at 100% uptake but the other cations with the exception of Cs reduced to around 20% uptake. Unfortunately an explanation of this trend cannot be provided, as the functionality of the material has not been disclosed.

The largest distribution coefficients for Zr ions were demonstrated by both the amidoxime chelate resin S910 and the inorganic D5530.

### Table 4: Separation factors and distribution coefficients for C100H.

<table>
<thead>
<tr>
<th>Nitric Acid Strength (M.)</th>
<th>Selectivity Factor</th>
<th>Equilibrium Coefficient ($K_{eq}$)</th>
<th>Distribution Ratio ($K_{ion}$)</th>
<th>Retention Factor ($K_{ion}$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Co (II)</td>
<td>Cu (II)</td>
<td>Sr (II)</td>
<td>Zr (IV)</td>
<td>Cs (I)</td>
</tr>
<tr>
<td>1</td>
<td>1.46</td>
<td>0.54</td>
<td>1101</td>
<td>2.07</td>
</tr>
<tr>
<td>1</td>
<td>1.84</td>
<td>2.69</td>
<td>1102</td>
<td>180</td>
</tr>
<tr>
<td>1</td>
<td>102.04</td>
<td>55.54</td>
<td>21129</td>
<td>4.08</td>
</tr>
<tr>
<td>1</td>
<td>1148</td>
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<td>0.11</td>
<td>1102</td>
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<tr>
<td>1</td>
<td>24.99</td>
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<td>13.60</td>
<td>1124</td>
</tr>
<tr>
<td>1</td>
<td>28.37</td>
<td>41.49</td>
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<td>2</td>
<td>121.18</td>
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<td>1102</td>
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<tr>
<td>2</td>
<td>12.82</td>
<td>5.86</td>
<td>10.60</td>
<td>132</td>
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<td>2</td>
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<td>613</td>
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<td>3</td>
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<td>0.47</td>
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<td>3</td>
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<td>3</td>
<td>2.86</td>
<td>1.53</td>
<td>2.94</td>
<td>1147</td>
</tr>
</tbody>
</table>

### Table 5: Separation factors and distribution coefficients for C100X10MBH.
Nitric Acid | Selectivity Factor | Equilibrium Coefficient ($K_d$) | Distribution Ratio ($K_{ion}$) | Retention Factor ($k_{ion}$)  
--- | --- | --- | --- |  
Strength (M) | Co (II) | Sr (II) | Zr (IV) | Cs (I) | Ce (IV) | Ce (III)  
--- | --- | --- | --- | --- | --- | ---  
1 | 2.55 | 0.00 | 1.30 | 164 | 2.06 | 4.50  
1 | 0.39 | 0.00 | 0.51 | 25 | 0.81 | 0.64  
1 | 1401.30 | 3671.12 | 1818.06 | 89129 | 2880.01 | 2258.14  
1 | 0.77 | 1.96 | 0.00 | 0.49 | 1.58 | 1.25  
1 | 1.57 | 4.01 | 0.00 | 2.04 | 3.23  
1 | 0.48 | 1.24 | 0.00 | 0.63 | 131 | 2.53  
2 | 0.00 | 0.00 | 0.01 | 0.13 | 12.32 | 0.25  
2 | 111.65 | 111.65 | 111.65 | 111.65 | 111.65 | 111.65  
2 | 0.00 | 0.00 | 0.00 | 0.00 | 0.00 | 0.00  
2 | 0.00 | 0.00 | 0.00 | 0.00 | 0.00 | 0.00  
3 | 0.76 | 0.01 | 72.92 | 2.83 | 2.71 | 31.29  
3 | 1.31 | 0.01 | 95.82 | 313 | 3.56 | 4155  
3 | 196.73 | 149.72 | 14346.70 | 557.73 | 533.18 | 962.00  
3 | 0.01 | 0.01 | 0.00 | 0.014 | 0.04 | 1.89  
3 | 0.35 | 0.27 | 0.00 | 0.96 | 5.50 | 0.11  
3 | 0.37 | 0.28 | 0.00 | 26.91 | 1.05 | 0.96  

Table 6: Separation factors and distribution coefficients for S910.

| Cations | Hydroxamic Acid | Logβ1 | Logβ2 | Logβ3 | Logβ4 | Reference  
--- | --- | --- | --- | --- | --- |  
U(vi) | BHA | 8.72 | 16.77 |  
U(vi) | BHA | 9.89 | 18.0 |  
Pu(vi) | BHA | 12.73 | 26.32 |  
Zr(iv) | BHA | 12.43 | 24.08 |  
Fe(III) | AHA | 11.42 | 21.10 |  
Fe(III) | BHA | 12.18 |  
Ce(III) | AHA | 5.45 | 9.79 |  
La(III) | AHA | 5.16 | 9.33 |  

Table 7: Stability constants for certain cations with BHA and AHA.

( functionality not disclosed) (Table 8). The distribution coefficient is lower or remains similar as the nitric acid concentration increases for all ions tested, with the exception of Zr on the D5530 adsorbent. Some of the cation’s unpredictable uptake with varying acid concentration could be due to the formation of cation-nitrate complexes as already described for Ce and Zr ions; this could be equally true for the transition metals Cu and Co (Figure 4). This complex formation will also affect ion hydration influencing hydration size. A third contributory factor could be the potential for protonation of the D5530 functional group thus transforming the cation properties to anionic but this would be expected with sulphonic acid resins. This irregular behaviour would be slower than simple salt formation, as unexpected as S910 is a chelating resin and it would be expected that Sr (II) ions only as it is the only cation to display any significant uptake on each of the adsorbents evaluated in nitric acid conditions.

The rate factor in ion-exchange adsorption is recognized for its importance for the economic and industrial employment of ion-exchange resins, no more so than the application under consideration. The kinetic data reported in Figures 5-7, show all four resins increased Zr ion uptake with time reaching near 100% after several hours for most resins but with the amidoxime resin having a more rapid uptake initially. At the 300 min. time interval the amidoxime (S910) is the more superior resin with the other three having similar rates. This is unexpected as S910 is a chelating resin and it would be expected that formation of complexes would be slower than simple salt formation, as would be expected with sulphonic acid resins. This irregular behaviour may be due to either the resin porosities being affected differently when in contact with 2 M nitric acid and/or the sulphonic acid resins forming more complex binding to the Zr$^{iv}$ ions than a simple salt formation, i.e., the incorporation of nitrate ion to achieve electron neutrality.

The largest increase in the uptake rate observed over the temperature range 25 to 45°C is for that of the inorganic D5530 whilst the S910 resin is the least affected by the change in temperature. This however may be due to the capacity of the resin for the ion and the particle diffusion mechanisms are prevalent in ion exchange process, although normally the slowest step (rate-limiting step) for a given system controls the speed of ion exchange.
The rate constant $k$ can be derived from:

$$\ln (1-\alpha) = -kt$$  \hspace{1cm} (5)

and the activation energy $E_a$ calculated using:

$$\ln k = \ln A - \frac{E_a}{RT}$$  \hspace{1cm} (6)

Where, $A$ is the pre exponential factor, $R$ is the gas constant, and $T$ is temperature.

The plots of $\ln (1-\alpha)$ versus time (not shown) at different temperatures were linear confirming that the ion exchange process is particle diffusion controlled. The value of the overall rate constant, $k$ was deduced from the slope of the plots. This was substantiated by comparing the coefficient of determination ($R^2$) measurements from the $\ln (1-\alpha)$ and $\ln (1-\alpha^2)$ plots for the Vermeulen particle and film diffusion models (Equations 7 and 8).

$$-\ln \left(1 - \alpha^2\right) = t \frac{D_p \pi^2}{r_p^3}$$  \hspace{1cm} (7)

$$-\ln (1-\alpha) = t \frac{3D_f C_o}{r_p \delta C_e}$$  \hspace{1cm} (8)

Where, $D_p$ is the diffusion coefficient in the ion exchanger ($\text{m}^2/\text{sec}$), $r_p$ is the radius of the particle ($\text{m}$), $D_f$ is the diffusion coefficient in the film ($\text{m}^2/\text{sec}$), $C_o$ and $C_e$ are concentrations in the solution and in the exchanger respectively ($\text{mol}/\text{m}^3$), and $\delta$ is the film thickness.

Although coefficient of determination values from the two plots at 25°C for the four exchange materials studied were close (more so for the sulphonic acid resins) suggesting an interplay of film and particle diffusion in the exchange process the best fits for all four were with the Vermeulen's particle diffusion model which exhibited higher values of coefficient determination (Table 8).

These values indicate that the rate controlling mechanism for the exchange reaction is particle diffusion. The activation energy values
Figure 5: Mass uptake of Zr (iv) on four adsorbents and resins at 25°C.

Figure 6: Mass uptake of Zr (iv) on four adsorbents and resins at 45°C.

Figure 7: Mass uptake of Zr (iv) on four adsorbents and resins at 65°C.
Nitric Acid Strength (M) | Selectivity Factor | Equilibrium Coefficient (K)<sub>e</sub> | Distribution Ratio (K<sub>ion</sub>) | Retention Factor (K<sub>Re</sub>)
--- | --- | --- | --- | ---
1 | Co (II) | 1.07 | 0.62 | 0.1 | 3.44 | 1041 | 90.64 | 1.81 | 18.56
1 | Cu (II) | 0.93 | - | 0.58 | 0.09 | 3.22 | 0.38 | 84.41 | 1.69 | 17.34
1 | Sr (II) | 1.62 | 1.73 | - | 0.15 | 5.57 | 0.66 | 148.3 | 2.97 | 30.03
1 | Zr (IV) | 10.48 | 11.21 | 6.47 | - | 36.1 | 4.26 | 955.2 | 19.1 | 194.4
1 | Cs (I) | 0.29 | 0.31 | 0.18 | 0.03 | - | 0.12 | 155.6 | 3.11 | 31.34
1 | Ce (IV) | 2.48 | 2.63 | 1.52 | 0.23 | 8.47 | - | 78.52 | 0.79 | 7.87
3 | Co (II) | 1.02 | 0.94 | 0.00 | 0.68 | 0.93 | 104.9 | 2.1 | 21.26
3 | Cu (II) | 0.98 | - | 0.92 | 0.00 | 0.67 | 0.91 | 103.8 | 2.08 | 20.92
2 | Sr (II) | 1.07 | 1.09 | - | 0.00 | 0.73 | 0.99 | 117.7 | 2.35 | 23.63
2 | Zr (IV) | 854.2 | 868.3 | 798.9 | - | 580 | 792.77 | 93737 | 1875 | 18920
2 | Cs (I) | 1.47 | 1.5 | 1.38 | 0.00 | - | 1.37 | 201.2 | 4.02 | 40.37
2 | Ce (IV) | 1.08 | 1.1 | 1.01 | 0.00 | 0.73 | - | 94.56 | 1.89 | 19.05

Table 8: Separation factors and distribution coefficients for D5530.

| Temperature (°C) | Zr (IV) Ion Exchange Capacity (mg/g) |
| | 25 | 45 | 60 |
| Resin | +/- | +/- | +/- |
| C100H | 14.6 | 0 | 17 | 0.4 | 19.8 | 0.9 |
| C100 × 10MBH | 14.0 | 1.3 | 18.4 | 1.7 | 19.0 | 0 |
| SN10 | 4.7 | 1.7 | 4.6 | 0.6 | 2.4 | 0.7 |
| D5530 | 2.4 | 1.2 | 6.1 | 0.5 | 4.2 | 0.2 |

Table 9: Mass of Zr ion uptake on different adsorbents at differing temperatures.

<table>
<thead>
<tr>
<th>Equation</th>
<th>Ion exchange material coefficients of determination</th>
</tr>
</thead>
<tbody>
<tr>
<td>C100H</td>
<td>C100×10MBH</td>
</tr>
<tr>
<td>-ln(1-α)&lt;sup&gt;2&lt;/sup&gt;</td>
<td>0.9914</td>
</tr>
<tr>
<td>-ln(1-α)</td>
<td>0.9426</td>
</tr>
</tbody>
</table>

Table 10: Comparison of coefficients of determination (R<sup>2</sup>) for the Vermeulen particle and diffusion models for Zr ions at 25°C.

predicted using equation 6 are 20.25, 20.87, 18.25 and 18.02 kJ/mole for C100H, C100X10MBH, S910 and D5530 respectively. These values suggest that the mechanism is one of ion exchange of the Zr ions with hydrogen ions on the sulphonic acid exchange materials, but for D5530 physisorption could also be playing a part and complexation for the S910 (Tables 9 and 10).

Conclusions

Of the four ion exchange materials reported in this paper the sulphonic acid resins offer little opportunity for chromatographic separation of Cs and/or Sr from other fission products when in nitric acid solutions of 1 to 3 M concentration. The data reported suggest it is modest i.e., less than 24 hrs to attain equilibrium for the sulphonic acid resins, much quicker for the amidoxime resin.

The rate of extraction of Zr ions from 2 M nitric acid solution is modest i.e., less than 24 hrs to attain equilibrium for the sulphonic acid resins, much quicker for the amidoxime resin.

There was an indication that the exchange materials under investigation in nitric acid solutions at ambient temperature could be undergoing oxidative/hydrolytic damage which was exacerbated at higher temperatures. The stability of these materials is the subject of another paper.

The kinetic data confirms that the rate controlling uptake step for Zr ions is particle diffusion and the mechanism is one of ion exchange for all four materials, the predicted Ea values were similar (~ 20 kJ/ mole). The inorganic exchanger (D5530) had the lowest Ea value (18.02 kJ/mole) which infers that ion exchange could be supplemented by physisorption; with complexation dominating the mechanism for the amidoxime resin (S910).

The separation factors recorded for the four materials were sufficiently encouraging for column studies and chromatographic separations to be undertaken. This is the subject of further publications.

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