Strontium sorption and precipitation behaviour during bioreduction in nitrate impacted sediments

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Abstract

The behaviour of strontium (Sr\(^{2+}\)) during microbial reduction in nitrate impacted sediments was investigated in sediment microcosm experiments relevant to nuclear sites. Although Sr\(^{2+}\) is not expected to be influenced directly by redox state, bioreduction of nitrate caused reduced Sr\(^{2+}\) solubility due to an increase in pH during bioreduction and denitrification. Sr\(^{2+}\) removal was greatest in systems with the highest initial nitrate loading and consequently more alkaline conditions at the end of denitrification. After denitrification, a limited re-release of Sr\(^{2+}\) back into solution occurred coincident with the onset of metal (Mn(IV) and Fe(III)) reduction which caused minor pH changes in all microcosms with the exception of the bicarbonate buffered system with initial nitrate of 100 mM and final pH>9. In this system ~95% of Sr\(^{2+}\) remained associated with the sediment throughout the progression of bioreduction. Analysis of this pH 9 system using X-ray absorption spectroscopy (XAS) and electron microscopy coupled to thermodynamic modelling showed that Sr\(^{2+}\) became partially incorporated within carbonate phases which were formed at higher pH. This is in contrast to all other systems where Sr\(^{2+}\) remained associated with the sediment. The behaviour of 90Sr, during changes in sediment biogeochemistry induced by bioreduction in nitrate impacted nuclear contaminated environments.

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In iron rich sediments, adsorption to both aluminosilicate clays and Fe(III)-oxyhydroxide minerals will have a significant control over Sr\(^{2+}\) mobility with pH important in controlling the mineral surface charge and therefore the extent of cation adsorption (Chiang et al., 2010). Clay minerals (eg illite, chlorite, kaolinite and montmorillonite) provide important adsorption surfaces for cations even at low pH due to their relatively low PZC (pH 4–6) and permanent structural charge, (Hussain et al., 1996; Dyer et al., 2000; Coppin et al., 2002; Zhuang and Yu, 2002; Alvarez-Silva et al., 2010) whilst Fe(III)-hydroxides tend to contribute significant adsorption sites at higher pH (PZC pH 7–8) (Small et al., 1999; Hofmann et al., 2005). With increasing pH and alkalinity, groundwater will become oversaturated with regard to carbonate phases and at high Sr\(^{2+}\) concentrations, this may allow the precipitation of Sr\(^{2+}\) as strontianite (Zachara et al., 1991; Tesoriero and Pankow, 1996; Greegor et al., 1997; Parkman et al., 2000; Coppin et al., 2002; Zhuang and Yu, 2002; Alvarez-Silva et al., 2010) whilst Fe(III)-hydroxides tend to contribute significant adsorption sites at rated into the newly formed mineral phase (Parmar et al., 2000; McKenzie and Armstrong-Pope, 2010). It should be noted that \(^{90}\text{Sr}\) has a relatively short half-life (28.8 years) and is therefore not a concern in groundwater environments.

Microbial metabolism has the ability to affect the geochemistry and mineralogy of subsurface sediments, as a result “bioreduction” systems have been considered for the remediation of groundwater containing the redox active radionuclides Tc and U (Lloyd and Renshaw, 2005). Tc and U have been shown to be immobilised by reduction from the more soluble Tc(VII) and U(VI) to poorly soluble Tc (IV) and U(IV) during Fe(III) reducing conditions (Lloyd, 2003; Law et al., 2010; Bellenger and Staunton, 2008; Chorover et al., 2008). As radioactive \(^{90}\text{Sr}\) is often found as a co-contaminant in Tc/U contaminated land (Riley and Zachara, 1992; Hartman et al., 2007; McKenzie and Armstrong-Pope, 2010), understanding the behaviour of Sr\(^{2+}\) during bioreduction is essential in predicting and managing the mobility of this problematic contaminant in both natural and engineered bioreduction scenarios. During bioreduction the solution pH will be affected by the reaction products which include OH\(^-\) and HCO\(_3^-\), and metal reduction will affect sediment mineralogy (Law et al., 2010; Thorpe et al., 2012). Reductive dissolution of bioavailable Fe(III)/Mn oxides and formation of new Fe(II) mineral phases may result in Sr\(^{2+}\) that was sorbed to Fe(III) oxide surfaces being released due to mineral dissolution (Langley et al., 2009a, b). However, it has been shown that during Fe(III) oxide crystallisation adsorbed contaminant metals (e.g. Pb\(^{2+}\)) can become incorporated into the newly formed phase, therefore, the effect of Fe(III) oxide recrystallisation has the potential to increase or decrease Sr\(^{2+}\) environmental mobility. At the same time, the increase in pH caused by bioreduction processes may lead to enhanced removal of Sr\(^{2+}\) through increased sorption to mineral surfaces and carbonate precipitation/substitution (Roden et al., 2002; Mitchell and Ferris, 2005; Chorover et al., 2008). Microbial metabolism can result in the production of CO\(_2^-\)/HCO\(_3^-\) which promotes alkaline pH conditions and supersaturation with regard to carbonate mineral phases (SrCO\(_3\) or CaCO\(_3\)) in which Sr\(^{2+}\) can be precipitated (Coleman et al., 1993; Fujita et al., 2004; Mitchell and Ferris, 2005). These processes can also lead to siderite (Fe(II)CO\(_3\)) formation during microbial reduction of Fe(III), which may result in minor Sr\(^{2+}\) becoming incorporated into the newly formed mineral phase (Parmar et al., 2000; Roden et al., 2002).

Here, we consider the effects of microbial metabolism on the biogeochemistry and speciation of Sr\(^{2+}\) in conditions relevant to radioactively contaminated sites using stable Sr\(^{2+}\) as an analogue for \(^{90}\text{Sr}\). Specifically, sub-surface nitrate concentrations are often elevated and have been reported in excess of 100 mM some nuclear facilities (Riley and Zachara, 1992; Finneran et al., 2002; Fredrickson et al., 2004; Istok et al., 2004; Senko et al., 2005; McKenzie and Armstrong-Pope, 2010). In this study, we have examined the behaviour of Sr\(^{2+}\) during the development of bioreducing conditions in sediments representative of the Sellafield nuclear facility that have been amended with between 0.3 and 100 mM nitrate. We tested the hypothesis that an increase in OH\(^-\) and CO\(_3^{2-}\)/HCO\(_3^-\) during nitrate reduction may lead to increased adsorption of Sr\(^{2+}\) to mineral surfaces and, once over-saturation was reached, the precipitation and or incorporation of Sr\(^{2+}\) into carbonate phases at the high Sr/Ca ratio used in this study (1:1:5). Overall, our aim is to assess whether bioreduction approaches may be relevant to a range of problematic radionuclides including redox active U and Tc as well as \(^{90}\text{Sr}\) and thus provide a holistic remediation strategy where co-contamination of these radionuclides occurs.

### 2. Methods

#### 2.1. Experimental section

##### 2.1.1. Sample collection

Sediments representative of the Quaternary unconsolidated alluvial flood-plain deposits that underlie the UK Sellafield reprocessing site were collected from the Calder Valley, Cumbria, during December 2008 (Law et al., 2010). The sampling area was located ~2 km from the Sellafield site and sediments were extracted from the shallow sub-surface (Lat 54°26′30 N, Long 03°28′09 W). Sediments were transferred directly into sterile containers, sealed, and stored at 4°C prior to use.

##### 2.1.2. Bioreduction microcosms

Microcosms (10±0.1 g Sellafield sediment, 100±1 ml groundwater) were prepared using a synthetic groundwater representative of the Sellafield region (Wilkins et al., 2007) that was manipulated to produce a range of treatments (Table 1). Aerated systems were first established at variable pH (4.5, 5.5 and 7) to assess Sr\(^{2+}\) sorption in oxic systems. Following this a range of sealed microcosms were prepared. Unbuffered systems with an initial pH of ~5.5, and representative of the mildly acidic in situ pH at the sample site, were prepared with 0.3, 10, and 25 mM nitrate amendments. Bicarbonate buffered systems with an initial pH of ~7 were prepared with 0.3, 10, 25, and 100 mM nitrate amendments. Sodium acetate was added as an electron donor in excess of available electron acceptors (10 mM for 0.3–10 mM nitrate treatments, 20 mM for 25 mM nitrate treatments and 70 mM for 100 mM nitrate treatments) and a deoxygenated NaN\(_3\) solution was used for NO\(_3^-\) amendment. Finally, Sr\(^{2+}\) (as stable SrCl\(_2\)) was added to each microcosm to achieve 1.15 mM which promotes alkaline pH conditions and HCO\(_3^-\) during nitrate reduction may lead to increased Sr\(^{2+}\) sorption to mineral surfaces and carbonate precipitation substitutions (Parmar et al., 2000; Roden et al., 2002).

<table>
<thead>
<tr>
<th>System</th>
<th>Initial pH</th>
<th>Nitrate (mM)</th>
<th>Acetate (mM)</th>
<th>Ionic strength</th>
</tr>
</thead>
<tbody>
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<td>Bicarbonate unamended</td>
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<td>0.3</td>
<td>10</td>
<td>0.024</td>
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<td>5.5</td>
<td>10</td>
<td>10</td>
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<tr>
<td>Bicarbonate unamended</td>
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<td>25</td>
<td>20</td>
<td>0.059</td>
</tr>
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<td>7</td>
<td>100</td>
<td>70</td>
<td>0.190</td>
</tr>
</tbody>
</table>

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*Table 1: Initial geochemical composition of microcosm systems.*
short half-life and thus high specific activity and even for the most impacted sites where groundwater concentrations of >1000 Bq l−1 have been reported the molar concentration of 90Sr will be very low (<10−11 mol l−1) compared to our experimental concentrations (Riley and Zachara, 1992; McKinley et al., 2007; Mckenzie and Armstrong-Pope, 2010). Triplet microcosms were then sealed with butyl rubber stoppers and incubated anaerobically at 20 °C in the dark for 110–250 days. At appropriate time points, sediment slurry was extracted under an O2-free Ar atmosphere using aseptic technique and centrifuged (15,000 g; 10 min) to provide wet sediment pellets and porewater samples for analysis of bioreduction products and strontium.

2.1.3. Geochemical analyses and imaging

During microcosm sampling, total dissolved Fe, Mn(II), and NO2 concentrations were measured with standard UV–vis spectroscopy methods on a Jenway 6715 UV–vis spectrophotometer (Goto et al., 1977; Viollier et al., 2000; Harris and Mortimer, 2002). Aqueous NO3−, SO42−, HCO3−/CO3− and acetate were measured by ion chromatography ( Dionex 4000i liquid chromatograph). Aqueous Sr2+ and Ca2+ were measured by ICP-AES (Perkin-Elmer Optima 5300). Total bioavailable Fe(III) and the proportion of extractable Fe(II) in the sediment was estimated by digestion of ~0.1 g of sediment in 5 ml of 0.5 N HCl for 60 min followed by the ferrozine assay (Stooy, 1970; Lovley and Phillips, 1986). The pH and Eh were measured with a pH/Eh metre (Denver Instruments, UB10) and probes calibrated to pH 4.7 and 10. Standards were routinely used to check the reliability of all methods and calibration regressions typically had R2 ≥ 0.99. The elemental composition and bulk mineralogy of the sediment were determined by X-ray fluorescence (Thermo ARL 9400XR and X-ray diffraction (Philips PW 1050 XRD). Selected end point samples were imaged using Environmental Scanning Electron Microscope (ESEM) in combination with Backscattering Electron Detection (BSE) and Energy Dispersive X-ray Analysis (EDAX) (Philips XL30 ESEM-FC).

2.1.4. X-ray absorption spectroscopy

Selected samples from the bicarbonate buffered pH 7 systems with 10, 25 and 100 mM nitrate amendments were chosen to examine Sr2+ speciation in: (1) oxic sterile control pH 7 sediment; (2) Fe(III)/SO42− reducing end point pH 7.2 sediments; (3) Fe(III)/SO42− reducing end point pH 8.1 sediments; and (4) Fe(III) reducing end point pH 9.3 sediments. Typical concentrations of Sr2+ in these samples were in the range 600–1000 ppm. Standards: (1) SrCl2 (aq), 3000 ppm (Fisher Scientific), (2) SrCO3 (s) (Fisher Scientific) and (3) natural Sr2+ substituted aragonite from crushed aragonite mineral sample (Sr2+ concentration ~1000 ppm), were prepared and diluted with boron nitride where necessary. Samples were transferred to XAS cells under anoxic conditions, cooled to ~ 80 K with a liquid nitrogen cryostat (see Nikitenko et al., 2008), and Sr K-edge XAS spectra were collected on beamline BM26A at the European Synchrotron Radiation Facility (ESRF). For sediment samples, Sr K-edge XAS spectra (16115.26 keV) were collected in fluorescence mode using a 9 element solid state Ge detector. Multiple scans were averaged in Athena version 0.8.061 (Ravel and Newville, 2005) and normalised XANES data plotted. Background subtraction for EXAFS analysis was performed using PySpline v1.1 (Tenderholt et al., 2007). EXAFS data were fitted using DLEXCURV v1.0 (Tomic et al., 2005) using full curve wave theory (Gurman et al., 1984) by defining a theoretical model which was informed by the relevant literature (e.g. O’Day et al., 2000; Finch et al., 2003) and comparing the model to the experimental data. Shells of backscatterers were added around the Sr2+ and by refining an energy correction Ef (the Fermi Energy; which for final fits typically varied between −3.8 and −2.6), the absorber–scatterer distance, and the Debye–Waller factor for each shell. Model iterations were performed until a least squares residual was minimised. Shells were only included in the model fit if the overall least square residual (the R-factor; Binsted et al., 1992) was improved by >5%.

3. Results and discussion

3.1. Sediment characteristics

Sediment composition was measured by X-ray fluorescence and was found to comprise Si (31.57%), Al (7.63%), Fe (3.64%), K (2.79%), Na (0.99%), C (0.96%), Mn (0.87%), Ti (0.45%), Ca (0.23%) and P, S and Cl (<0.1%). We note that XRF analyses show that phosphate is present in our systems at very low concentrations (<0.008%) and is not likely to be a significant control on Sr2+ behaviour in this system.

Trace metal analysis showed natural Sr2+ to be present in sediments at 62.8 ± 0.2 ppm and natural aqueous Sr2+ was <1 ppm. Strontium was added to groundwater media in significant excess to the natural background at 100 ppm (1.15 mM Sr2+) and Ca2+ was present at 67 ppm (1.67 mM) thus a Sr/Ca ratio of ~1:1.5 was present in the synthetic groundwater media. The concentration of 0.5 N HCl extractable Fe(III) in the sediment was 5.6 ± 0.5 mmol kg−1 prior to incubation and the sediment pH was ~5.5.

3.2. Sorption to oxic sediment

In sterile control microcosms, increased Sr2+ sorption was observed in microcosms with a high pH and a low ionic strength. For example, for a constant ionic strength system (1 = 0.027 mol dm−3) run at pH 4.5, 5.5 and 7, the Sr2+ removal from solution was 35.6 ± 1.9%, 47.4 ± 5.3% and 63.2 ± 2.1% respectively (equating to Kd values of 5.5, 9.0 and 17.1 ml g−1). Differences in strontium behaviour in the sterile microcosms were attributed to pH dependent differences in sorption to mineral surfaces present in the sediment. Sorption to both clays and Fe(III)-oxyhydroxide surfaces is possible although clay minerals is likely to predominate in unbuffered microcosms in which the pH of 4.5–5.5 is above the PZC for many clay minerals (Coppin et al., 2002; Zhuang and Yu, 2002; Alvarez-Silva et al., 2010) whilst Fe-oxyhydroxides become more significant as pH approaches their PZC at pH ~7 (Dyer et al., 2000; Hofmann et al., 2005). Additionally, in control experiments at pH 7 and with increasing ionic strength (0.027, 0.037, 0.062 and 0.190 mol dm−3) resulting from sodium nitrate and sodium acetate additions, Sr2+ sorption was 68, 65, 55 and 28% respectively (equating to Kd values of 21.2, 18.5, 12.2 and 3.8 ml/g), presumably reflecting increased competition for Sr2+ sorption sites at higher ionic strengths due to cation exchange processes (Hull and Schafer, 2008).

3.3. Biogeochemistry in sediment microcosms

The unbuffered (initial pH 5.5; nitrate range 0.3–25 mM) and bicarbonate buffered (initial pH 7.0; nitrate range 0.3–100 mM) experiments all underwent progressive anoxia and electron acceptors were utilised in the order NO3− > NO2− > Mn/Fe(III) > SO42− (Fig. 1 and 2). Microbially mediated nitrate reduction caused a decrease in porewater nitrate and transient accumulation of nitrite in all systems. The onset of Fe(III) reduction was indicated by an increase in sediment extractable Fe(II) and this was then followed by a decrease in porewater SO42− indicating sulfate reduction. No geochemical changes were observed in sterile control microcosms. In unbuffered systems (initial pH 5.5), as expected, microbial activity was inhibited at low pH and terminal electron accepting processes proceeded more slowly than in the parallel bicarbonate buffered microcosms (initial pH 7.0) (Figs. 1 and 2) (e.g. Law et al., 2010; Thorpe et al., 2012). However, in the unbuffered systems, nitrate reduction led to the release of OH− and HCO3−, amending the pH such that a pH increase from 5.5 to 6.8, 7.5 and 8.3 occurred during the period of 0.3, 10 and 25 mM nitrate respectively (Fig. 1; Table 2; Thorpe et al., 2012). Metal reduction commenced once nitrate reduction had occurred and mid-point 0.5 N HCl extractable Fe(III) reduction was observed at approximately 25, 35 and 45 days for systems with 0.3, 10 and 25 mM nitrate (Fig. 1). Nitrate reduction and the associated pH increase in all microcosms coincided with removal of Sr2+.
and Ca\(^{2+}\) from solution (Fig. 1). Interestingly, as bioreduction progressed through Fe(III) and SO\(_4^{2-}\) reduction in these dynamic systems, a small amount of both Sr\(^{2+}\) and Ca\(^{2+}\) (<10% of that sorbed after nitrate reduction) was remobilised to solution. This re-release coincided with a slight decrease in pH (<0.5 pH units) presumably due to re-equilibration of the microcosm system following nitrate reduction. The re-release of sorbed Sr\(^{2+}\) and Ca\(^{2+}\) may be due solely to pH dependent sorption/desorption to mineral surfaces or in some systems (for example above pH 7) there may be release of Sr\(^{2+}\) and Ca\(^{2+}\) sorbed to Fe(III)-oxyhydroxides as reductive dissolution of the Fe(III) phases occurred (Small et al., 1999; Roden et al., 2002; Langley et al., 2009a, b).

In bicarbonate buffered systems with an initial pH of 7, the final pH following the reduction of 0.3, 10, 25 and 100 mM nitrate was 7.5, 8.0, 8.5 and 9.4 (Fig. 2: Table 2). Terminal electron accepting processes proceeded faster than in unbuffered microcosms with mid-point 0.5 N HCl extractable Fe(III) reduction occurring at ~20 days for 0.3 and 10 mM nitrate systems and around 40 and 160 days for systems with 25 and 100 mM nitrate (Fig. 2). As with the unbuffered systems, Sr\(^{2+}\) and Ca\(^{2+}\) were removed from solution during nitrate reduction with increasing pH and a small amount (<10% of that sorbed after nitrate reduction) of Sr\(^{2+}\) and Ca\(^{2+}\) was re-released into solution in all systems apart from the bicarbonate buffered,
100 mM nitrate system (Fig. 2). Here, the final pH was 9.3 and interestingly, Sr\(^{2+}\) remained associated with the sediment throughout Fe(III) and sulfate reduction. In this high nitrate loaded system, the utilisation of 70 mM acetate resulted in the accumulation of 207 ± 4.9 mM of dissolved inorganic carbon and amended the pH to alkaline conditions.

For comparison with other studies it is useful to examine distribution coefficients for Sr\(^{2+}\) (Kd = (solid in g kg\(^{-1}\)/aqueous in g ml\(^{-1}\))). Distribution coefficients are only relevant to the specific geochemical conditions of each system of study and give an indication of the extent of Sr\(^{2+}\) partitioning onto the solid phase in different systems. Here Kd values ranged from <10 ml g\(^{-1}\) in systems with a high ionic strength (0.190 mol dm\(^{-3}\)) or a low pH (5.5) and increased to >50 ml g\(^{-1}\) with increasing pH. The distribution coefficient in systems with a final pH >9 was calculated to be 133 ml g\(^{-1}\). These Kd values compare well with literature values where surface sediment Kd values are typically between 10 and 200 ml g\(^{-1}\) (Deldebbio, 1991; Liszewski et al., 1998; Dewiere et al., 2004).

Modelling of the solution chemistry in bioreduced systems (PHREEQC-2: LLNL database) suggested that for all unbuffered bioreduced system end-points, Fe(II)CO\(_3\) and SrCO\(_3\) were oversaturated in (PHREEQC-2: LLNL database) suggested that for all unbuffered bioreduced system end-points, Fe(II)CO\(_3\) and SrCO\(_3\) were oversaturated in.

Table 2

<table>
<thead>
<tr>
<th>System</th>
<th>Sterile oxic</th>
<th>90% nitrate reduction</th>
<th>90% Fe(III)/SO(_4) reduction</th>
<th>End point</th>
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</thead>
<tbody>
<tr>
<td></td>
<td>% Sr(^{2+}) on sediment</td>
<td>pH</td>
<td>% Sr(^{2+}) on sediment</td>
<td>pH</td>
</tr>
<tr>
<td>Bicarbonate unamended 0.3 mM nitrate</td>
<td>47.0</td>
<td>5.5</td>
<td>54 ± 2.2</td>
<td>6.6</td>
</tr>
<tr>
<td>Bicarbonate unamended 10 mM nitrate</td>
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<td>5.5</td>
<td>75 ± 1.8</td>
<td>8.0</td>
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<td>Bicarbonate unamended 25 mM nitrate</td>
<td>45.5</td>
<td>5.5</td>
<td>82 ± 2.1</td>
<td>8.2</td>
</tr>
<tr>
<td>Bicarbonate amended 0.3 mM nitrate</td>
<td>67.9</td>
<td>7.0</td>
<td>63 ± 1.4</td>
<td>7.0</td>
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<tr>
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<td>7.0</td>
<td>78 ± 0.7</td>
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<td>7.0</td>
<td>84 ± 0.4</td>
<td>8.5</td>
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<td>Bicarbonate unamended 100 mM nitrate</td>
<td>32.6</td>
<td>7.0</td>
<td>93 ± 1.4</td>
<td>9.3</td>
</tr>
</tbody>
</table>

* Differences in Sr\(^{2+}\) sorption to sterile controls occur due to varying pH and ionic strength due to the addition of NaHCO\(_3\), Na-acetate and NaNO\(_3\).
Further analysis showed that a model EXAFS fit was possible with additional shells of 2.6 carbon atoms at 3.03 Å, 2.8 strontium atoms at 4.18 Å and 2.5 strontium atoms at 4.87 Å; these values are approximately 50% of what is expected for pure SrCO\textsubscript{3}, which is consistent with a model where approximately half of Sr\textsuperscript{2+} is present in a SrCO\textsubscript{3} like environment (Table 4; Fig. 5). Indeed, this model, which is geochemically sensible, resulted in a better fit to the spectrum and a 27% reduction in the least square residual when compared to the data modelled as 100% adsorbed Sr\textsuperscript{2+} suggesting that both adsorption and incorporation occurred in this system (Table 4).

In natural and engineered environments concentrations of Sr\textsuperscript{2+} and \textsuperscript{90}Sr are generally much lower than in these experiments (eg 0.1 ppm natural Sellafield groundwater) (Wilson, 1996). Under Sellafield conditions, model simulations predicted that bioreduced systems would remain supersaturated with respect to CaCO\textsubscript{3}. It is therefore feasible that substitution of Sr\textsuperscript{2+} into CaCO\textsubscript{3} rather than precipitation as SrCO\textsubscript{3} will be important in controlling the mobility of both natural Sr\textsuperscript{2+} and artificial \textsuperscript{90}Sr in such systems. Indeed, it is well documented that Sr\textsuperscript{2+} can substitute for Ca\textsuperscript{2+} within the calcium carbonate lattice (Pingitore et al., 1992; Tesoriero and Pankow, 1996; Greegor et al., 1997; Warren et al., 2001; Finch et al., 2003). Recent studies, focused on bacterial urolysis, have found that Sr\textsuperscript{2+} incorporation into the CaCO\textsubscript{3} lattice was enhanced by the rapid precipitation rates resulting from HCO\textsubscript{3}\textsuperscript{−} production and the pH rise associated with microbial respiration (Fujita et al., 2004; Mitchell and Ferris, 2005). Both a pH rise and dissolved organic carbon production were observed during bioreduction by indigenous microorganisms in this study suggesting that nitrate reduction might also result in enhanced Sr\textsuperscript{2+} uptake into calcite compared to those observed under slower precipitation rates.
Overall, our experiments showed that there is increased Sr$^{2+}$ removal from solution during bioreduction in nitrate impacted sediments compared to sterile control systems. In systems with an initially low pH (5.5), removal of Sr$^{2+}$ from solution after bioreduction was particularly enhanced, presumably due to the increased sorption onto deprotonated mineral surfaces as the pH increased above 6. After nitrate reduction, system re-equilibration and an associated decrease (<0.5 pH units) in pH resulted in modest (<10%) re-release of Sr$^{2+}$ into solution highlighting the vulnerability of adsorbed Sr$^{2+}$ to re-release due to changing geochemical conditions. In extreme environments with very high (100 mM) nitrate concentrations, bioreduction led to a final pH of >9 and enhanced removal of Sr$^{2+}$ from solution occurred throughout the bioreduction cascade. This study has shown that in very high nitrate systems an increase in pH and dissolved inorganic carbon associated with microbial reduction and particularly denitrification can promote the precipitation and incorporation of Sr$^{2+}$ into carbonate phases although the engineering aspects of this process are as yet unexplored. Clearly, radio-strontium incorporation into carbonate phases is desirable in remediation scenarios as they are redox insensitive phases and are potentially more resistant to remobilization than sorbed Sr$^{2+}$. It is also clear that bioreduction scenarios have the potential to impact Sr$^{2+}$ mobility in the subsurface and that understanding the bioreduction behaviour of redox inactive radionuclides can be of significance in assessing the efficacy of bioreduction schemes at nuclear facilities. We further suggest that under constrained conditions, bioreduction may have the potential to co-treat redox active radionuclides and $^{90}$Sr increasing the range of applications for this clean-up technology across the global nuclear waste legacy.

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