Rock alteration in alkaline cement waters over 15 years and its relevance to the geological disposal of nuclear waste

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Abstract

The interaction of groundwater with cement in a geological disposal facility (GDF) for intermediate level radioactive waste will produce a high pH leachate plume. Such a plume may alter the physical and chemical properties of the GDF host rock. However, the geochemical and mineralogical processes which may occur in such systems over timescales relevant for geological disposal remain unclear. This study has extended the timescale for laboratory experiments and shown that, after 15 years two distinct phases of reaction may occur during alteration of a dolomite-rich rock at high pH. In these experiments the dissolution of primary silicate minerals and the formation of secondary calcium silicate hydrate (C–S–H) phases containing varying amounts of aluminium and potassium (C–(A)–(K)–S–H) during the early stages of reaction (up to 15 months) have been superseded as the systems have evolved. After 15 years significant dedolomitisation (MgCa(CO3)2 + 2OH- → Mg(OH)2 + CaCO3 + CO3(aq)2) has led to the formation of magnesium silicates, such as saponite and talc, containing variable amounts of aluminium and potassium (Mg–(Al)–(K)–silicates), and calcite at the expense of the early-formed C–(A)–(K)–S–H phases. This occurred in high pH solutions representative of two different periods of cement leachate evolution with little difference in the alteration processes in either a KOH and NaOH or a Ca(OH)2 dominated solution but a greater extent of alteration in the higher pH KOH/NaOH leachate. The high pH alteration of the rock over 15 years also increased the rock’s sorption capacity for U(VI). The results of this study provide a detailed insight into the longer term reactions occurring during the interaction of cement leachate and dolomite-rich rock in the geosphere. These processes have the potential to impact on radionuclide transport from a geodisposal facility and are therefore important in underpinning any safety case for geological disposal.

1. Introduction

A widely recognised concept for the disposal of radioactive waste, which will remain hazardous for hundreds of thousands of years, is emplacement in a geological disposal facility (GDF). Many proposed GDF concepts for Intermediate Level Waste (ILW), such as those in the UK, France, Canada and Switzerland (NDA, 2010a; Andra, 2012; Nuclear Waste Management Organisation, 2010; Nagra, 2014), involve cement e.g. as a waste-form, backfill and construction material. Post-closure, groundwater will saturate the facilities and cement dissolution will produce a high pH leachate which will evolve in composition and pH over the lifetime of the GDF (Atkinson, 1985; Berner, 1992). Initially, dissolution of KOH and NaOH within the cement will form a leachate of pH ~13. The leachate pH will then decrease to ~12.5 where it will be buffered by equilibration with portlandite (Ca(OH)2). The leachate will remain at this pH until all the Ca(OH)2 has dissolved, after which, pH will be controlled by equilibrium with calcium silicate hydrate (C–S–H) gel and will decrease to ~10.5. The leachate will form a chemically disturbed zone (CDZ) in the geosphere surrounding the GDF, also known as an alkaline disturbed zone (ADZ; NDA, 2010b). Previous studies have shown that in the CDZ high pH leachates could cause the dissolution of aluminosilicate minerals and formation of secondary mineral phases (e.g. C–S–H phases, Gaucher and Blanc, 2006 and references therein). This could...
change the physical (e.g. porosity and permeability) and chemical (e.g. reactive surface area and sorption capacity) properties of the host rock thereby affecting radionuclide transport. For example, mineral dissolution could increase rock permeability and promote radionuclide transport, or the formation of secondary solid phases could block flow paths and have the opposite effect. The formation of secondary phases will also change the nature of the surfaces with which radionuclides could interact e.g. any increased sorption capacity of secondary phases could retard contaminant transport. The potential for cement leachates to alter host rock properties shows that understanding the chemical and mineralogical changes which occur will be key to developing a long term safety case for any cementitious GDF. However, there is a lack of long-term (>10 years) experimental studies which investigate these processes.

Overall, it has been suggested that CDZ alteration can be divided into two regions (Savage, 2011). Firstly, a zone closest to the cement/rock interface (zone 1) where extensive mineral alteration occurs, including dissolution of primary silicate minerals and precipitation of secondary solid phases. Secondly, a zone further from the interface (zone 2) where fluid chemistry is perturbed and ion exchange reactions are important, but rock alteration is significantly diminished. Many short term laboratory and underground rock laboratory experimental studies have investigated rock and mineral alteration in high pH cement leachates (zone 1; e.g. Gaucher and Blanc, 2006; Bateman et al., 1999; and references therein; Braney et al., 1993; Cuevas, 2004; Mäder et al., 2006).

Generally, these studies have found that reaction in high pH, Ca-bearing cement-type leachates results in the dissolution of silicate minerals (Eq. (1)) followed predominantly by the precipitation of secondary C–S–H phases (Eq. (2)) of varying Ca:Si ratio (e.g. 0.5–1.5 (Gaucher and Blanc, 2006)), morphology and crystallinity e.g. C–S–H gel (Savage and Rochelle, 1993; Hodgkinson and Hughes, 1999).

\[
\text{KAlSi}_3O_8 + 3\text{OH}^- + 2\text{H}_2\text{O} \rightarrow \text{Al(OH)}_4^- + 3\text{HSiO}_4^{aq} + K^{aq} (1)
\]

\[
\text{HSiO}_3^{aq} + \text{Ca}^{aq}_n + \text{H}_2\text{O} \rightarrow \text{C–S–Hgel} (2)
\]

where aluminium (e.g. from primary mineral dissolution) and potassium (e.g. dissolved in cement leachate) are present, secondary aluminium and potassium bearing C–S–H (C–(A)–(K)–S–H) phases have also been identified (e.g. Braney et al., 1993; Savage et al., 1992). Studies of clay alteration (e.g. bentonite) at high pH found the formation of a number of Na/K/Ca bearing silicate phases including zeolites (e.g. phillipsite (K,Na,Ca)\textsubscript{1.2}(Si,Al)\textsubscript{1.0}O\textsubscript{16}•9H\textsubscript{2}O, analcime NaAlSi\textsubscript{2}O\textsubscript{6}•H\textsubscript{2}O) and apophyllite (KCa\textsubscript{4}Si\textsubscript{6}O\textsubscript{20}(OH)•8H\textsubscript{2}O) (Gaucher and Blanc, 2006; Ramirez, 2005). Carbonate may also be released into solution during high pH rock alteration. For example, cement pore water can promote the breakdown of dolomite (CaMg(CO\textsubscript{3})\textsubscript{2}) according to the reaction shown in Eq. (3) (Poole and Sotiropoulos, 1980; Bérubé et al., 1990; Braithwaite and Heath, 2013 and references therein) releasing carbonate to solution leading to the formation of calcium carbonate minerals e.g. calcite. However, these processes have not been studied in the context of the CDZ.

\[
\text{MgCa(CO}_3\textsubscript{2}) + 2\text{OH}^- \rightarrow \text{Mg(OH)}_2 + \text{CaCO}_3 + \text{CO}_3^{aq} (3)
\]

Generally, experimental studies have limited timescales, with few longer than 1–2 years (e.g. a 540 day experiment is the longest study reviewed by Gaucher and Blanc, 2006) and no longer-term experimental studies examine the stability of the secondary phases formed. However, GDFs will evolve over tens to hundreds of thousands of years. To investigate the effect of high pH alteration at timescales more comparable to GDF scenarios, natural and anthropogenic analogue sites have been studied. At natural analogue sites such as Maqarin, Jordan (Milodowski et al., 1998; Alexander, 1992; Alexander et al., 2012; Linklater, 1998; Savage, 2011) and Troodos, Cyprus (Alexander et al., 2011), alkaline groundwaters have interacted with rock at timescales extending beyond 1 million years. Whereas anthropogenic analogue sites (e.g. the Tournemire Tunnel; Tinsley et al., 2006; Techer et al., 2012) bridge the gap between laboratory experiments and natural analogues. A review of many such sites representing timescales of alteration from ~30 years to >1 million years is provided by Savage (2011). Overall, these studies indicate that over time a variety of secondary phases can form, predominantly alkali-silica gels (e.g. C–S–H gel), which can crystallise with time to zeolites, C–S–H minerals e.g. okienne (CaSiO\textsubscript{2}•2H\textsubscript{2}O), and feldspars. The key factors controlling which phases form are primarily solution composition (e.g. pH) and reaction time. Modelling has also been used to predict the chemical and physical evolution within the CDZ (e.g. Savage et al., 1992; Savage and Rochelle, 1993; Braney et al., 1993; Bateman et al., 1999; Piffigsten et al., 2006; Soler and Mäder, 2007; Fernandez et al., 2010; Alexander et al., 1992). These studies generally support the experimental findings that silicate mineral dissolution is followed by secondary solid phase (e.g. C–S–H) formation, with subsequent transformation of C–S–H to feldspar and zeolite over time, as found at analogue sites. However, modelling predictions are limited by the ability to constrain which solids will form due to slow reaction rates, and a lack of reliable thermodynamic data for some phases (e.g. C–S–H gel).

The secondary solid phases produced during high pH rock alteration may affect radionuclide migration through the CDZ by changing the sorption properties of the material. A key radionuclide of concern is U(VI) which is highly mobile and hazardous over the long timescales relevant to geological disposal (NDA, 2010c). As C–S–H has been found to be the predominant secondary phase produced by high pH rock alteration and is also the most abundant phase in hardened cement paste (Taylor, 1990), which is used as an ILW wasteform, the interaction of U(VI) with these phases has been studied in some detail (Harfouche et al., 2006; Tits et al., 2011; Gaona et al., 2012; Atkins and Glasser, 1992). However, few experimental studies have looked at the interaction of U(VI) with secondary phases in-situ following high pH mineral/rock alteration and these have generally been limited to investigation after only several months of high pH reaction (e.g. Berry et al., 1999). Continued evolution of altered material over decades may further change surface properties and so affect U(VI) interactions. Therefore experimental investigation of these interactions with rock after extended periods of alteration could help fill this knowledge gap.

1.1. Review of alteration experiment up to 15 months

In this study rock alteration by high pH cement waters has been characterised in unique experiments lasting over 15 years, providing new insight into longer-term rock alteration. These experiments were originally part of the Nirex Safety Assessment Research Programme (NSARP) run by the British Geological Survey (BGS). The experiments were started in 1995 and the rock type and solution compositions used reflect the focus of NSARP at that time (Rochelle et al., 1997; details of how to access this report are provided in Supporting Information). However, as the rock contains many common rock-forming minerals the long-term alteration processes which have occurred will be representative of many rock types. The experiments investigate reaction of disaggregated dolomite-rich fracture fill rock (Borrowdale Volcanic Group (BVG), UK) with fluids representative of young (pH 13, Young Near Field Pore-water (YNFP) and intermediate (pH 12, Evolved Near Field Groundwater (ENFG)) cement leachates. The products of these experiments (fluid and solid phases) were initially investigated up to 15 months of reaction, and a full description of the results
up to that point is presented elsewhere (Rochelle et al., 1997). However, a brief summary of the results is given here. Characterisation of the rock surfaces indicated intensive dissolution of primary silicate minerals (e.g. feldspar), and the formation of poorly crystalline alkali silicate gels (i.e. C–(A)–(K)–S–H phases), and less abundant crystalline apophyllite–KOH. Although the solid secondary phases produced in both leachates were similar, alteration was more extensive in the pH 13 YNFP system. Analysis of the pore fluids showed that the pH of the YNFP and ENFG reduced to 11.8 and 9.7 respectively. During this time the concentration of dissolved Si and Al increased significantly, due to the dissolution of the silicate minerals, and the Ca concentration decreased due to the precipitation of Ca-rich secondary phases. Overall, the reaction followed the 2-stage dissolution and precipitation process observed in other studies. However, the data indicated that the system was not at equilibrium (i.e. the concentration of ions in solution was not stable) and that further evolution of the system may occur over a longer timescale. For example, the dolomite in the rock may undergo dedolomitisation (Eq. (3)) as observed by (Bérubé et al., 1990), leading to the release of significant amounts of carbonate and magnesium.

The aim of this study was to characterise the evolution of the BVG alteration experiments after 15 years of reaction and determine the influence of rock alteration on its reaction with U(VI). This was achieved through (1) characterisation of the changes in fluid composition and solid phase stability, crystallinity and composition between 15 months and 15 years of reaction; (2) assessment of any difference in reactions due to leachate composition; (3) comparison of U(VI) interactions with unaltered rock and rock altered at high pH for 15 years. This work will aid understanding of the longer term effects of the cement leachate on rock alteration and radionuclide transport in the CDZ.

2. Materials and methodology

A full description of the experimental method is provided by Rochelle et al. (1997). A summary is given here. The rock used in the experiments was altered wallrock and dolomite-mineralised fracture fill from a hydrogeologically conductive fracture zone in the BVG, Ordovician basement volcanic rocks, UK (Milodowski et al., 1998). It was collected from the United Kingdom Nirex borehole B114A in Cumbria, UK at 859 m depth (UK grid reference NY 0248 0569) (Rochelle et al., 1997, see Supporting Information for access details). A large (1–2 kg) sample of the rock was disaggregated and sieved. A sub-sample of the 125–250 μm size fraction was then reacted with two synthetic cement leachates. The leachates were designed (see Table 1) to represent a young near field porewater (YNFP; pH 13.0 at 25 °C) and an evolved near field groundwater (ENFG; pH 12.2 at 25 °C). The YNFP was dominated by dissolved KOH and NaOH, and saturated with respect to Ca(OH)₂, and the ENFG represented a synthetic deep groundwater (i.e. high salinity, Na/Cl and NaSO₄ rich), saturated with respect to Ca(OH)₂. The ENFG included the influence of deep groundwater, but the YNFP does not because a significant amount of time would be required for groundwater to ingress into the repository. Each solution was prepared under a nitrogen atmosphere to minimise interactions with CO₂. For further details see Rochelle et al., 1997 in Supporting Information).

The experiments were conducted as sacrificial batch experiments in stainless steel pressure vessels lined with Teflon® (Rochelle et al., 1997). Each 150 ml vessel was loaded under a nitrogen atmosphere with 35 g of rock and 140 g of fluid. Non-reacting ‘blank’ experiments for both leachates were also run containing approximately 100 g of fluid in 100 ml vessels. The cells were held in an oven at 70 °C ± 0.5 °C and shaken regularly by hand to achieve mixing.

The experiments reported here were sampled after 15 years and 4 months in a CO₂ controlled anaerobic chamber, with a hydrogen–nitrogen atmosphere. Solution samples were filtered to <0.2 μm with a nylon filter. A sub-sample of the solution was then immediately acidified (with 2% HNO₃) for cation analysis and another frozen (−20 °C) for anion analysis. Suspendeed fines were collected by preserving the 0.2 μm filter papers inside a CO₂ free desiccator. The pH and Eh of the filtrates were measured at the point of sampling at room temperature within the anaerobic chamber.

Cation concentrations in the solutions were analysed by Inductively Coupled Plasma Atomic Emission Spectroscopy (ICP-AES) using a Perkin–Elmer Optima 5300 dual view system. Ion Chromatography (IC) was carried out to quantify anion concentrations using a Dionex DX120 ion exclusion system with a Dionex ICE AS1 column for carbonate analysis and a Dionex AS9-HC column for all other anions. During the 15 years of reaction some experimental fluid evaporated and the extent of this has been estimated using the change in Cl⁻ (expected to be a conservative species) concentration over time. In the ENFG Cl⁻ concentration increased at the same rate in both the reacted and blank solutions (Table S1) and indicated approximately 3% fluid loss. In the YNFP system it was not possible to use Cl⁻ due to its very low concentration. However, a similar level of evaporation was observed in all the experiments, therefore it was assumed that 3% of the fluid was lost from every experiment and the data are corrected for this (Table S2). BET surface area analyses of the reacted rock and a sample of unaltered rock were performed using a Micromeritics Gemini V Surface Area Analyser. Quantitative analysis of the bulk mineralogy and the <2 μm size fraction of the altered and unaltered samples were determined by powder X-ray Diffraction (XRD) using a PANalytical X’Pert Pro series diffractometer, in conjunction with Rietveld refinement (for sample preparation and data analysis details see Supporting Information).

The rock samples were imaged using an FEI QUANTA 600 environmental scanning electron microscope (ESEM) and an FEI QUANTA 650 field emission gun ESEM (FEG–ESEM) both equipped with Oxford Instruments INCA 450 energy dispersive X-ray (EDX) microanalysis systems with a 50 mm X-Max silicon drift detector (SDD). The reacted solids and the particles retained on the 0.2 μm filter papers were imaged uncoated under low vacuum. Sections through the solids were created by embedding samples in resin and polishing under ethane-2-diol. These were then viewed uncoated under low vacuum and coated with 2.5 nm of carbon under high-vacuum. Transmission electron microscopy (TEM) with EDX and selected area electron diffraction (SAED) was conducted using an FEI Tecnai TF20 FEG–TEM with an Oxford Instruments INCA 350 EDX system with 80 mm X-Max SDD detector and a Gatan Orion SC600A CCD camera. Samples were prepared for TEM by ultrasonication in ethanol to create a suspension which was then deposited onto copper TEM grids with holey carbon films.

### Table 1

<table>
<thead>
<tr>
<th>Chemical component</th>
<th>YNFP (mg l⁻¹)</th>
<th>ENFG (mg l⁻¹)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Na</td>
<td>1640</td>
<td>7730</td>
</tr>
<tr>
<td>K</td>
<td>3630</td>
<td>174</td>
</tr>
<tr>
<td>Mg</td>
<td>67.1</td>
<td>1980</td>
</tr>
<tr>
<td>Si</td>
<td>–</td>
<td>174</td>
</tr>
<tr>
<td>Cl</td>
<td>–</td>
<td>11400</td>
</tr>
<tr>
<td>Br</td>
<td>–</td>
<td>25.2</td>
</tr>
<tr>
<td>CO₃²⁻</td>
<td>8.14</td>
<td>1.50</td>
</tr>
<tr>
<td>SO₄²⁻</td>
<td>–</td>
<td>1120</td>
</tr>
<tr>
<td>SiO₄²⁻</td>
<td>7.60</td>
<td>–</td>
</tr>
<tr>
<td>pH (at 70 °C)</td>
<td>11.8</td>
<td>10.9</td>
</tr>
</tbody>
</table>
U(VI) reaction with unaltered rock, and 15 year reacted samples from the YNFP and ENFG systems was also examined. 0.1 g of the unaltered and reacted solids were each reacted in 35 ml of a 0.1 M NaCl solution (pH 7.0; representative of a simplified saline groundwater) with UO$_2^{2-}$ spiked to a concentration of 3 mg l$^{-1}$ under N$_2$. The suspensions were then agitated for 24 h prior to sampling. The solutions were filtered to $<0.45\mu$m (nylon filter) and acidified (1% HNO$_3$), while the solids were frozen moist at $-80^\circ$C. The solutions were analysed by inductively coupled plasma mass spectroscopy (ICP-MS) for uranium concentration. From these data, distribution coefficient ($K_d$) values for U(VI) sorption (assuming no U(VI) phase precipitation) were calculated according to Eq. (4) (Fetter, 1999), where $C^*$ is the mass of solute sorbed per dry unit weight of solid (mg kg$^{-1}$) and $C$ is the concentration of solute in solution in equilibrium with the solid (mg l$^{-1}$).

$$C^* = K_dC$$

The speciation of U associated with the rock samples was analysed using X-ray absorption spectroscopy (XAS). For this, the moist solid samples (~50% moisture) were mounted in double contained cells. Uranium L$_3$-edge XAS spectra were collected at beamline B18 of the Diamond Light Source. The data were collected in fluorescence mode using a 9 element Ge solid state detector at room temperature. Standards were collected in transmission mode for scooter (U(VI)) and uraninite (U(IV)). The software package Athena (Ravel and Newville, 2005) was used to average multiple scans for each sample to improve the signal to noise ratio of the data and for background subtraction.

3. Results

3.1. Solution chemistry

After 15 years, the pH of the reacted YNFP and ENFG solutions were 9.9 and 8.8, respectively, while those of the corresponding 'blank' solutions remained oxic throughout the experiment. The Eh of the reacted YNFP and ENFG solutions were +112 mV and +258 mV respectively, and those of the corresponding 'blank' solutions were +28 mV and +90 mV respectively, indicating the experimental solutions remained oxic throughout the experiment.

The Na, K, Ca, Mg, Si, CO$_3^-$, Sr and SO$_4^{2-}$ concentrations of all the solutions at 15 years (corrected for evaporation) are shown in Fig. 1 along with the data for 0, 4, 9 and 15 months of reaction (Rochelle et al., 1997). Al concentration was <0.5 mg l$^{-1}$ after 15 years of reaction and in all but the starting solutions (Rochelle et al., 1997). The concentrations of K and Na did not change between 15 months and 15 years of reaction (see Fig. 1 a and b). Ca, Si, Sr and SO$_4^{2-}$ concentrations decreased in both leachates over the same period (see Fig. 1c-f). However, differences in the Mg and CO$_3^-$ concentrations existed between the two leachates. After 15 years Mg concentration was elevated in the ENFG, and CO$_3^-$ concentration was elevated in the YNFP (see Fig. 1g and h). It is highlighted that the solution concentration of Mg in the YNFP and ENFG is very low (<0.01 mg l$^{-1}$) up to 15 months, with the exception of the concentration in the YNFP at t = 0. It should also be noted that at 9 and 15 months of reaction, an elevated concentration (approx. 40 mg l$^{-1}$) of CO$_3^-$ was measured in the ENFG solution blank. However, as these levels were not observed throughout the experiment, it was attributed to the ingress of atmospheric CO$_2$(g) during analysis.

3.2. Mineral alteration

Quantitative XRD analysis of the unaltered rock showed it comprised of quartz, dolomite, mica, orthoclase feldspar, calcite, hematite and anatase (Table 2). Note that it was not possible to distinguish between different mica/clay phases (e.g. muscovite, biotite and illite) and they are grouped into ‘mica’ XRD characterisation of the 15 year altered fracture fill identified the same bulk solids, with the addition of halite in the ENFG system, which is an artefact of sample drying. The XRD data further indicate that relative to the unaltered material, the YNFP altered rock is enriched in calcite, from 2.6% to 6.4%, and depleted in dolomite from 29.5% to 23.7% (see Table 2). Data for the ENFG altered rock suggest a similar but less definitive trend in this system with an increase in calcite from 2.6% to 4.5%, and a decrease in dolomite from 29.5% to 28.5%. The proportions of the other minerals remain unchanged with the exception of an increase in ‘mica’ in the YNFP altered material. However, it should be noted that the error on these analyses are up to ±2.5%, therefore some of these changes may be within analytical error.

Further XRD characterisation of the fraction identified as ‘mica’ was undertaken by isolating the <2 µm size fraction from the samples. In the unaltered rock, the clay minerals were identified as predominantly illite, with a trace amount of chlorite (Table 3) which confirms the findings of Rochelle et al. (1997). In the reacted YNFP and ENFG samples, XRD indicated that the predominant clay mineral was still illite. However, in the reacted samples, the patterns were characterised by broad, low intensity peaks centred on ~12.2 Å in the XRD pattern from the YNFP altered material and between 12.5 and 15 Å from the ENFG altered material. Following ethylene glycol-solvation a peak centred on ~17 Å appeared in the XRD patterns for both samples (Figs. S1 and S2). This suggests the presence of swelling clay, not present in the unreacted material, which is more abundant in the ENFG altered material than the YNFP material (Table 3). NEWMOD-modelling of the XRD data (Reynolds and Reynolds, 1996) from the YNFP altered material suggested that this interstratified illite/smectite consisted of 80% smectite, 20% illite phase. The absence of a peak at ~14 Å also suggests that there is no significant chlorite present in the altered samples.

Characterisation of the minerals using SEM revealed that the rock grains reacted for 15 years in both the YNFP and ENFG retained their angular shape and 125–250 µm size distribution indicating that no large scale dissolution or precipitation has occurred (Fig. S3). Dolomite reacted in YNFP for 15 years showed extensive pitting, indicating dissolution of the mineral surface (Figs. 3a and S4). However, examination of the material in section indicated the dissolution was limited to grain surfaces. Some dissolution of dolomite had been observed after 15 months of reaction, but to a much lesser extent than observed after 15 years (Rochelle et al., 1997). Euhedral, rhombohedral crystals of calcite, 5–10 µm in length, (Fig. 3b) were observed on silicate grain surfaces. This is thought to be secondary as calcite is not observed with this euahedral morphology in the early stages of the experiment (Rochelle et al., 1997).

After reaction in ENFG, grains of dolomite also exhibit surface pitting indicative of dissolution although to a much lesser extent than in the YNFP altered material (Fig. 2c and d). As in the YNFP system CaCO$_3$ crystals with rhombohedral morphology were observed on silicate mineral surfaces and are also identified as secondary calcite. However, in the ENFG system CaCO$_3$ identified as calcite, was also observed as poorly developed crystal coatings on dolomite grains (Fig. 2d). These coatings were not present on the unaltered dolomite or the material reacted for up to 15 months (Rochelle et al., 1997).

The surfaces of the quartz and feldspar altered in YNFP were completely coated with secondary phases with ‘sheet-like’ and acicular morphologies (Fig. 3a). The sheets were typically several micrometers in size (see Fig. 3b) and the needles several micrometers in length and <0.5 µm in width. EDX showed these
Fig. 1. Concentrations of (a) K, (b) Na, (c) Ca, (d) Si, (e) Sr, (f) SO$_4$, (g) Mg and (h) CO$_3$ in solution in the starting solution and after 4, 9, 15 months and 15 years of reaction in YNFP and ENFG (solid triangle and square markers respectively) and in corresponding 'blank' experiments (corresponding unfilled markers).
particles contained Mg, Al, K and Si and, coupled with their morphology, indicated they were aluminosilicate clays, potentially the interstratified illite/smectite clay phases identified via XRD.

The aluminosilicate surface coatings on the YNFP altered rock were further examined using TEM to eliminate interferences from the underlying primary mineral grains. The most abundant phase exhibited a ‘sheet-like’ morphology with particles ranging from 100’s of nm to a few μm in size (Fig. 4a). Typically, this was identified via EDX as an Mg–silicate, though low levels of additional K and Fe were noted in a few examples of this phase. SAED patterns for these particles show the typical hexagonal pattern indicative of sheet silicate structured phases (Fig. 4a), and together with the EDX identify this phase as talc (Mg₃Si₄O₁₀(OH)₂). Two acicular morphology phases were also identified; (i) elongate rods approximately 100 nm in width and several μm in length, identified as a Mg–Al–silicate based on EDX analysis and exhibiting the typical hexagonal SAED pattern indicative of aluminosilicate clays (Fig. 4b), and; (ii) shorter rods approximately 100 nm in width and <1 μm in length, identified by EDX as a Mg–Al–K–silicate (Fig. 4c). There was also a phase which consistently occurred as groups of radiating acicular needles forming a ‘sheet-like’ morphology up to 2 μm in size (Fig. 4d). EDX indicated this phase was an Mg–Al–K–silicate with low and variable Fe content.

SEM analysis of the ENFG altered rock also revealed surface coatings of aluminosilicate clay particles on the primary silicate grains, similar to those observed in the YNFP altered system (Fig. 5a). These coatings comprised of particles with either ‘sheet-like’ or elongate

### Table 2
Quantitative XRD analyses of the bulk mineralogy of the rock altered in YNFP and ENFG for 15 years and a sample of unreacted rock (‘mica’ represents undifferentiated mica/clay species and may include muscovite, biotite, illite etc.).

<table>
<thead>
<tr>
<th>Mineral</th>
<th>Unreacted material (weight%)</th>
<th>YNFP altered sample (weight%)</th>
<th>ENFG altered sample (weight%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Quartz</td>
<td>41.3</td>
<td>39.1</td>
<td>39.3</td>
</tr>
<tr>
<td>‘Mica’ (muscovite/biotite/illite etc.)</td>
<td>12.8</td>
<td>15.9</td>
<td>13.0</td>
</tr>
<tr>
<td>Dolomite</td>
<td>29.5</td>
<td>23.7</td>
<td>28.5</td>
</tr>
<tr>
<td>Calcite</td>
<td>2.6</td>
<td>6.4</td>
<td>4.5</td>
</tr>
<tr>
<td>Orthoclase</td>
<td>11.9</td>
<td>12.9</td>
<td>11.9</td>
</tr>
<tr>
<td>Hematite</td>
<td>1.7</td>
<td>1.7</td>
<td>1.6</td>
</tr>
<tr>
<td>Anatase</td>
<td>&lt;0.5%</td>
<td>&lt;0.5%</td>
<td>&lt;0.5%</td>
</tr>
<tr>
<td>Halite</td>
<td>nd</td>
<td>nd</td>
<td>1.0</td>
</tr>
</tbody>
</table>

### Table 3
Summary of the clay minerals identified through XRD analysis in the <2 μm size fraction of unaltered and YNFP and ENFG altered rock after 15 years of reaction.

<table>
<thead>
<tr>
<th>% Of &lt;2 μm fraction mineral assemblage</th>
<th>Unreacted material</th>
<th>YNFP altered sample</th>
<th>ENFG altered sample</th>
</tr>
</thead>
<tbody>
<tr>
<td>Illite</td>
<td>99</td>
<td>76</td>
<td>96</td>
</tr>
<tr>
<td>Chlorite</td>
<td>1</td>
<td>&lt;1</td>
<td>1</td>
</tr>
<tr>
<td>Interstratified illite/smectite</td>
<td>nd</td>
<td>24</td>
<td>3</td>
</tr>
</tbody>
</table>

Fig. 2. SEM images of dolomite altered for 15 years: After reaction with YNFP (a) dolomite grain surface pitting, (b) calcite rhombs (highlighted in black ovals) on silicate grain surface. After reaction with ENFG (c) dolomite grain exhibiting minor pitting and a CaCO₃ coating of limited extent, (d) detail of pitting on dolomite grain surface (highlighted in the white circle) with CaCO₃ on surface indicated by the white arrow.
was found to contain Sr and S (Fig. 6d) and based on chemistry and phase occurred as euhedral, tabular, lath shaped crystals, 50–150 μm in size, composed of pure Mg–silicate which were identified as talc. The elongate particles had two distinct morphologies, elongate rods (approximately 100 nm in width and several μm in length, e.g. Fig. 5c), and short rods (approximately 100 nm in width and <1 μm in length, e.g. Fig. 5d) which were both identified as Mg–Al–K–silicates using EDX. The Mg, Al and K content of all the particles varied significantly, though all three elements were always present with minor Fe also present in some examples (Fig. 5c and d). It is believed that these Mg–(Al)–(K)–silicate phases, identified in both the YNFP and ENFG altered rock, correspond to the interstratified illite/smectite clay phase indicated in the XRD analyses, most likely a mixture of Mg-rich smectite (e.g. Mg–rich saponite–K) and Mg-rich interlayered illite/smectite. As expected, the magnitude of these alteration features were less extensive in the less aggressive ENFG system and additional minor secondary phases of nontronite and celestite were observed in this more chemically complex system. Interestingly, SEM and TEM analyses did not identify C–S–H phases or appophyllite–KOH in the materials reacted for 15 years in the YNFP or ENFG experiments although these were pervasive in the altered BVG up to 15 months of reaction (Rochelle et al., 1997).

### 3.3. U(VI) sorption

To assess whether alteration affected the sorption properties of the rock, the reaction of U(VI) with unaltered, and 15 year altered rock samples was studied. The surface areas of the unaltered and YNFP and ENFG 15 year altered rock were determined to be 5.31, 6.04 and 4.93 m² g⁻¹ (±3%), respectively, and indicated that any differences in sorption capacity between the samples was unrelated to surface area.

The concentrations of U(VI) in solution following the adsorption experiments (see Table 4) indicate that less than 6% of the U(VI) was taken up by the unaltered BVG and 30% and 40% was taken up by ENFG and YNFP altered material, respectively.

The calculated Kd values for both 15 years reacted rock samples are an order of magnitude greater than the unaltered material (Table 4). Assuming no U(VI) phases have precipitated (no evidence for discrete U(VI) minerals was observed), this indicated that the secondary phases or altered primary mineral surfaces have a higher sorption capacity for U(VI) than the unaltered rock. Interestingly, the Kd of the YNFP altered system was modestly elevated compared to that of the ENFG altered system, suggesting greater U(VI) retention on this sample. This indicates that the secondary phases/ altered surfaces created during alteration are key to U(VI) adsorption and that the extent of alteration directly correlates with U(VI) uptake.

Comparison of the size, shape and positions of the peaks in the uranium X-ray absorption near edge spectroscopy (XANES) spectra from the samples with those from the standards indicated that the U(VI) sorbed to the rock samples has the same local bonding environment as U(VI) in schoepite (Fig. 7). This indicates that the uranium is present as U(VI) coordinated by six oxygen atoms, two axial oxygen atoms at ~1.8 Å and four equatorial oxygen atoms at ~2.4 Å, in a standard uranyl geometry (Grenthe et al., 1997).
These XANES spectra and U speciation are consistent with other studies of U(VI) adsorption at neutral pH to a range of mineral phases e.g. clay (Schlegel and Descostes, 2009) and goethite (Sherman et al., 2008) which support the conclusion that uranium was adsorbed to the surface of the rock fragments. Due to the low concentration of U(VI) associated with the samples, it was not possible to collect extended X-ray absorption fine structure (EXAFS) data of sufficient quality in the time available required for further interpretation of the U(VI) coordination environment.

4. Discussion

This investigation identified significant changes in both the mineralogy of the BVG rock and cement leachates after 15 years of reaction and demonstrated that this had an impact on radionuclide retention. However, in order to understand the evolution of these systems, it is necessary to discuss the results of this study in with those from the first 15 months of the reaction as reported by Rochelle et al. (1997). In summary, during the first 15 months of reaction poorly-ordered C-(A)-(K)-S-H phases and minor
secondary appophyllite–KOH formed in both the YNFP and ENFG. However, between 15 months and 15 years of reaction the solution composition of both the YNFP and ENFG changed significantly and a different secondary mineral assemblage stabilised. This demonstrated that continued chemical and mineralogical evolution occurred and indicated a change in the reaction processes in these systems. Importantly, between 15 months and 15 years, two interconnected reactions occurred: firstly, dedolomitisation; and secondly, the transformation of C–(A)–(K)–S–H and illite to Mg-bearing aluminosilicate clays. These processes are discussed in detail below.

4.1. Dedolomitisation

Electron microscopy and quantitative XRD show the dissolution of dolomite and formation of secondary calcite, providing evidence that dedolomitisation occurred (Eq. (3)). In addition, changes in solution compositions were indicative of dedolomitisation. In the YNFP carbonate concentration increased from 166 mg l\(^{-1}\) in the original solution to 5070 mg l\(^{-1}\) after 15 years of reaction (Fig. 1h). The gradual increase in CO\(_3\) concentration throughout the experiment (Fig. 1h) is attributed to dedolomitisation, and may correlate with the minor amount of dolomite dissolution.
The same increase in carbonate concentration was not observed in the ENFG system (Fig. 1f). However, as the concentration of Ca\(^{2+}\) in the ENFG is high, any CO\(_3^{-}\) released through dedolomitisation would rapidly form CaCO\(_3\). Dedolomitisation would also release significant amounts of Mg\(^{2+}\). Evidence for this is observed in the ENFG system ([Mg\(^{2+}\)] > 200 ppm at 15 years), but the concentration of Mg\(^{2+}\) in the YNFP remains low, <1 mg l\(^{-1}\) throughout the experiment. This is attributed to either the low solubility of Mg\(^{2+}\) in highly alkaline solutions (Baes and Mesmer, 1976) and/or the rapid uptake of Mg\(^{2+}\) into secondary clay phase formation (see Section 4.2).

The dedolomitisation reaction consumes hydroxyl ions (Eq. (3)) and is therefore favoured at high pH (Min and Mingshu, 1993). However, as only minor dedolomitisation was observed at high observed at the beginning of the experiment (Rochelle et al., 1997). The same increase in carbonate concentration was not observed in the ENFG system (Fig. 1f). However, as the concentration of Ca\(^{2+}\) in the ENFG is high, any CO\(_3^{-}\) released through dedolomitisation would rapidly form CaCO\(_3\). Dedolomitisation would also release significant amounts of Mg\(^{2+}\). Evidence for this is observed in the ENFG system ([Mg\(^{2+}\)] > 200 ppm at 15 years), but the concentration of Mg\(^{2+}\) in the YNFP remains low, <1 mg l\(^{-1}\) throughout the experiment. This is attributed to either the low solubility of Mg\(^{2+}\) in highly alkaline solutions (Baes and Mesmer, 1976) and/or the rapid uptake of Mg\(^{2+}\) into secondary clay phase formation (see Section 4.2).

Table 4

<table>
<thead>
<tr>
<th></th>
<th>Unaltered</th>
<th>Altered in YNFP</th>
<th>Altered in ENFG</th>
</tr>
</thead>
<tbody>
<tr>
<td>U(VI) in solution (mg l(^{-1}))</td>
<td>2.8 ± 0.06(^a)</td>
<td>1.8 ± 0.04(^a)</td>
<td>2.1 ± 0.06(^a)</td>
</tr>
<tr>
<td>% U(VI) adsorbed</td>
<td>6.7</td>
<td>40</td>
<td>30</td>
</tr>
<tr>
<td>Kd (kg l(^{-1}))</td>
<td>714</td>
<td>6670</td>
<td>4290</td>
</tr>
</tbody>
</table>

\(^a\) Error is 1 standard deviation from the mean (triplicate analyses).
pH up to 15 months we can say that the rate of this reaction was restricted in the systems studied. This highlights the importance of longer term (>10 years) experimental studies to fully resolve the chemical and mineralogical reactions occurring in the CDZ. Thermodynamic predictions suggest that dedolomitisation would not occur below pH 11 (Min and Mingshu, 1993). As pH had decreased below 10 in the ENFG by 15 months of reaction (Rochelle et al., 1997) dedolomitisation would likely be restricted during the latter stage of the experiment. However, a high Ca\(^{2+}\) concentration, as found in both leachates, has been shown to promote dedolomitisation at lower pH, for example at cement–dolomite aggregate interfaces in concrete (Min and Mingshu, 1993). The enhancement of dedolomitisation occurs as Ca\(^{2+}\) reacts with aqueous CO\(_3^{2-}\) released in dedolomitisation to form CaCO\(_3\) effectively removing CO\(_3^{2-}\) from solution and so driving the dedolomitisation reaction. Accordingly, the occurrence of dedolomitisation in the latter stage of this experiment is attributed to the presence of aqueous Ca\(^{2+}\) in the cement leachates. This suggests that this reaction will always be favoured in the CDZ.

### 4.2. Mg–silicate formation

Examination of the reacted fracture fill showed that after 15 years, in both leachate systems, granates were coated with secondary Mg–(Al)–(K)–silicate minerals, most likely a mixture of talc, smectite (Mg–rich saponite–K) and interstratified illite/smectite. Similar Mg-rich solid phases have been observed previously as minor components in high pH clay alteration experiments/analogues. For example, saponite was recognised as an alteration product of bentonite by Cuevas (2004), secondary Mg-enriched mineral phases have been observed at the Tournemire analogue site (Tinseau et al., 2006; Techer et al., 2012) and polygorskite was identified as an Mg-rich alteration product of bentonite by high pH groundwater in the Troodos natural analogue site (Alexander et al., 2012). The formation of these phases at these high pH conditions is not unexpected as they are commonly associated with alkaline, saline lake environments (Yeniyol, 2007, 2012; Hojati et al., 2010; Akbulut and Kadir, 2003) where they form from post-sedimentary alteration of soils and rocks in high pH fluids in the presence of dolomitic material (Derkowski et al., 2013; Xie et al., 2013; Schwarzenbach et al., 2013; Birsoy, 2002). Their formation has also been identified in cement–aggregate systems where dedolomitisation in the presence of Ca(OH)\(_2\) and amorphous silica resulted in the formation of a mixed calcium–magnesium silica gel (Gali et al., 2001).

The presence of secondary Mg-bearing phases provides further evidence of dedolomitisation as dolomite is the major source of Mg\(^{2+}\) in the rock. However, dedolomitisation generally results in the production of brucite (Mg(OH)\(_2\)) (see Eq. (3)), which was initially found as a minor reaction product up to 15 months of reaction, but was not identified after 15 years. We suggest that this is due to the preferential reaction of the released Mg\(^{2+}\) to form the secondary Mg–(Al)–(K)–silicates. This has been observed in high pH cement systems where the presence of aqueous Mg and Si results in the formation of hydrated magnesium silicates such as talc–serpentine group minerals (Glasser, 2001; Eglington, 2006). We propose that two independent reaction pathways result in the assemblage of secondary solid phases observed. Firstly, the reaction of Mg\(^{2+}\) released by dedolomitisation with aqueous silica to form the pure Mg–silicate, talc, by direct precipitation, as shown in Eq. (5).

$$3\text{Mg}^{2+} + 4\text{H}_2\text{SiO}_4 + 2\text{OH}^- \rightarrow \text{Mg}_4\text{Si}_2\text{O}_{10}\text{(OH)}_2 + 6\text{H}_2\text{O} \quad (5)$$

This is evidenced by the decrease in aqueous Si concentration in solution between 15 months and 15 years of reaction in both the YNFP and ENFG fluids (see Fig. 1d), and the occurrence of talc in suspension which is indicative of formation through direct precipitation from solution. Secondly, the reaction of Mg\(^{2+}\) with primary illite and early formed C–(A)–(K)–S–H leading to transformation to Mg–(Al)–(K)–silicates. The transformation of illite to smectite (i.e. Mg–rich saponite–K) is likely to occur via an interstratified illite/smectite clay. Such interstratified clays are known to form as intermediate phases during the transformation of clay minerals (Cuadros et al., 2010). In this study the high pH and Mg\(^{2+}\) concentrations may have led to the transformation of illite either via cation exchange reactions and/or dissolution and precipitation. In addition, the absence of any C–(A)–(K)–S–H after 15 months of reaction shows that these phases have also either dissolved or transformed during the latter stages of the experiment. The transformation of C–S–H as a result of dedolomitisation has been suggested in numerical simulations of cement–aggregate systems; the release of CO\(_3^{2-}\) from dolomite results in calcite precipitation at the expense of the Ca content of C–S–H, and Mg\(^{2+}\) substitutes into the C–S–H structure eventually forming Mg–silicates (Gali et al., 2001). We proposed that this process is responsible for the destabilisation of the C–(A)–(K)–S–H formed in the early months of this experiment, and further promoted the dedolomitisation reaction by removing CO\(_3^{2-}\) from solution.

### 4.3. YNFP and ENFG comparison

Qualitatively, the degree of primary mineral dissolution and secondary phase formation appears greater in the YNFP. This would be expected due to the YNFP’s more aggressive, higher pH. However, despite the difference in solution composition, no significant difference was found in the assemblage of alteration phases produced in the two cement waters or in the surface area of the two samples. This indicates that broadly similar reaction processes have occurred in both systems, with the extent of alteration directly related to solution pH. In a real repository environment i.e. an open system, there would be a constant replenishment of cement pore water into the geosphere, therefore the observed decreases in pH and changes in solution composition would not occur to the same extent within the CDZ. This would increase the overall rate of the alkaline alteration reactions, but would be unlikely to significantly change which reactions occurring or the products formed. Two minor additional secondary phases, celestite and nontronite, were however identified in the ENFG system. Celestite precipitated from solution during the first 15 months of reaction (Rochelle et al., 1997) but surface pitting after 15 years indicated re-dissolution of this phase. As Sr\(^{2+}\) is known to substitute into calcite (Tesoriero and Pankow, 1996; Lorens, 1981; Pingitore and Eastman, 1986), and as calcite formed between 15 months and 15 years of reaction, it is proposed that Sr\(^{2+}\) substitution into calcite has driven celestite dissolution. Nontronite was not identified up to 15 months of reaction and Fe concentration in solution was generally below detection limits throughout the experiment as would be expected at highly alkaline pH. However, a significant proportion of the primary dolomite in the fracture fill is known to be ferroan (Rochelle et al., 1997). Therefore the occurrence of nontronite after 15 years of reaction is attributed to the reaction of Fe released from ferroan dolomite with dissolved silica via a similar mechanism to that which formed the secondary Mg–bearing aluminosilicate phases.

### 4.4. Geochemical modelling

Geochemical speciation and reaction-path modelling (PHREEQC, Parkhurst and Appelo, 2010) has been used to investigate the potential for dedolomitisation to produce the observed formation of Mg–(Al)–(K)–silicate phases in the experiments observed between 15 months and 15 years of reaction. Firstly, saturation
indices (SI) of mineral phases were calculated for the YNFP and ENFG aqueous compositions at 70 °C at 15 years using the Lawrence Livermore National Laboratory (LLNL) database, see Table S2 for aqueous composition considered and Table S3 for calculated saturation indices of oversaturated silicate and carbonate phases. In both YNFP and ENFG disordered dolomite is undersaturated, consistent with the observed dolomite dissolution. More crystalline forms of dolomite are oversaturated in both solutions. Calcite is oversaturated in YNFP consistent with its precipitation, but it is undersaturated in ENFG. It should be noted that measured carbonate concentrations considered in these calculations and hence calculated SI could be higher (more positive) than in the unopened experiment as CO₂ might have dissolved in the solutions on sampling. The model results confirm undersaturation of dolomite and oversaturation or equilibrium with calcite in YNFP consistent with dedolomitisation.

Considering silicates, for YNFP saponite phases with exchangeable K, Na, Ca and Mg, talc, phlogopite and chrysotile are the only silicate phases in the database that are oversaturated. Reaction-path calculations showed that phlogopite followed by chrysotile were the most stable phases. However, these phases are only likely to able to crystallise at high temperatures, e.g. above 100 °C. The oversaturation of saponite and talc is consistent with their observed formation in YNFP. Reaction-path calculation where talc and the saponite phases are allowed to precipitate predicted the precipitation of 2.1e-07 mol/kg water of saponite–K would form.

In the case of ENFG, experimental Al solution concentration after 15 years of reaction was below the analytical detection limit (Table S2). Therefore in this speciation calculation the limiting concentration of 0.01 mol/l Al was assumed in order to consider Al containing phases. A larger number of phases are oversaturated than for YNFP, which may be a consequence of the assumed Al concentration, although several high temperature pyroxene and olivine pure Mg silicate phases are thermodynamically stable at the lower pH of ENFG. Saponite and talc phases are again oversaturated, but the SI are higher than for YNFP. Interestingly brucite (Mg(OH)₂) is very close to saturation in ENFG (SI -0.04). Reaction path calculations that allow saponite and talc to precipitate resulted in the formation 1.1e-06 mol/kg water of Saponite–Mg and 1.4e-05 mol/kg water of talc. These speciation and reaction-path calculations on the YNFP and ENFG solution compositions confirm the dedolomitisation reaction and the associated precipitation of the Mg silicate phases saponite and talc, which drive the reaction (Eq. (5)).

To further examine the controls on talc or saponite precipitation in the experiments the dolomitisation reaction was simulated as a Reaction with PHREEQC. In this reaction dolomite was reacted with C–S–H in both YNFP and ENFG fluid compositions reported at 15 months of reaction (Rochelle et al., 1997). C–S–H is represented in the reaction model as C–S–H–gel (Reardon, 1990, 1992) for which the thermodynamic data was added to the LLNL database (the thermodynamic data for all phases are given in Supplementary Information). Saponite and talc phases were allowed to precipitate during the reaction. For each solution two scenarios were examined (i) where the reaction occurred in solutions with no additional source of Al and (ii) where an additional source of Al containing phases. A larger number of phases are oversaturated than for YNFP, which may be a consequence of the assumed Al concentration, although several high temperature pyroxene and olivine pure Mg silicate phases are thermodynamically stable at the lower pH of ENFG. Saponite and talc phases are again oversaturated, but the SI are higher than for YNFP. Interestingly brucite (Mg(OH)₂) is very close to saturation in ENFG (SI -0.04). Reaction path calculations that allow saponite and talc to precipitate resulted in the formation 1.1e-06 mol/kg water of Saponite–Mg and 1.4e-05 mol/kg water of talc. These speciation and reaction-path calculations on the YNFP and ENFG solution compositions confirm the dedolomitisation reaction and the associated precipitation of the Mg silicate phases saponite and talc, which drive the reaction (Eq. (5)).

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In these models the extent of reaction is defined by the assumption that the reaction is driven by the rate and extent of dolomite dissolution. The difference in the weight% of dolomite in the unaltered and altered rock samples, identified by quantitative XRD (Table 2), indicates that over 15 years of reaction ~0.019 and ~0.001 mol of dolomite have dissolved in the YNFP and ENFG systems respectively. A constant rate of dedolomitisation was assumed; therefore the amount of dolomite dissolved between 15 months and 15 years can be calculated as 1.7 × 10⁻² moles in the YNFP and 9.2 × 10⁻⁴ moles in the ENFG. The reaction was modelled at 70 °C and in order to examine the sequence of mineral precipitation products, it was divided into 18 equal reaction-steps. At each step the system was equilibrated with all components. As C–S–H was observed up to 15 months of reaction but not after 15 years, for the purpose of the model it is necessary to assume C–S–H dissolves during this time. Since there is no quantitative data for the amount of C–S–H formed in the experiment, the quantity of C–S–H dissolving in each leachate was estimated and refined using trial and error to get good agreement between the modelling results and experimental data for pH and Ca²⁺ concentration at 15 years of reaction. As a result it is assumed 2.1 × 10⁻⁴ moles and 1.1 × 10⁻⁵ moles C–S–H dissolve in the YNFP and ENFG, respectively. The solution chemistry and mineral SI evolution over time predicted by the model are summarised in Tables S4 and S5. Broadly, the modelled solution chemistry at 15 years of reaction is similar to the observed solution chemistry (same order of magnitude). However, modelled SI concentrations are higher and Mg concentrations are lower.

Throughout the modelled reaction the SI of dolomite and C–S–H are negative in both the YNFP and ENFG, in the presence and absence of a dissolving Al-bearing solid, indicating that these phases were undersaturated (the predicted SI of all phases over time in both leachates are provided in Supplementary Information). The amount of the solid reactants and precipitated secondary solid phases predicted to form in YNFP is shown in Fig. 8. Results for ENFG are provided in Supplementary Information.

In both leachates in the absence of a dissolving Al-bearing phase, (Fig. 8a) a very small amount of saponite forms initially, but talc is the main alteration product formed. In the case where muscovite is included in the model (Fig. 8b), saponite forms in preference to talc. The formation of talc where Al is limited is consistent with the observed occurrence of talc present as suspended particles in the experiment fluids. The formation of saponite associated with the presence of muscovite is consistent with the association of saponite and mixed layered illite–smectite with the BVG rock which provides a source of Al to stabilise saponite in preference to talc.

Approximately the same quantity of saponite is predicted to form in both YNFP and ENFG in the absence of muscovite and results also confirm the experimental observations that calcite and talc can form in both leachate systems (Figs. 8a and 56). In YNFP talc is the main secondary phase predicted to form in association with a minor amount of secondary saponite (Fig. 8a) while in ENFG saponite initially precipitates followed by talc (Fig. 56). The relative amounts of saponite–K and talc that form is related to the amount of K available for saponite–K formation. Interestingly, in ENFG saponite–Ca is predicted to form rather than saponite–K attributed to the significantly higher concentration of Ca in this fluid (Table S4). The model also reproduced the drop in pH observed in both leachate systems (Table S5). Overall, thermodynamic considerations show that dedolomitisation can lead to the transformation of silicate phases (i.e. C–S–H), leading to the formation of talc and saponite–K, which is in agreement with experimental observations.

4.5. Uranium sorption

The investigation of U(VI) sorption to the materials altered in this study indicated that, despite no significant detectable change in surface area, uranyl sorption to the BVG reacted at high pH for 15 years was greater than to unreacted BVG. This suggests alteration of the rock and the formation of secondary phases, including Mg–silicates, may increase the rock’s sorption capacity for U(VI).
Results also show that sorption is modestly greater to the material reacted in YNFP than to that reacted in ENFG. This supports the hypothesis that the presence of secondary phases increases the sorption capacity of the rock as secondary phases are more abundant in the YNFP system. This indicates that long term, high pH mineral reactions could be beneficial for the geosphere as a barrier to radionuclide migration and highlights the need for further study of the secondary minerals formed in this study in relation to radionuclide transport.

5. Summary and conclusions

This study investigated the reaction of a fracture fill material with high pH cement leachates within 15 year laboratory experiments. As a result of the extended timescale of these experiments, two different phases of the alkaline rock alteration reaction were identified. In phase 1, a Ca–silicate dominated secondary mineral assemblage (i.e. containing C–S–H and apophyllite–KOH) stabilised as reported by Rochelle et al. (1997) (Fig. 9). Subsequently these initial phases destabilised and Mg–(Al)–(K)–silicates and calcite formed under the reaction conditions of this study (Phase 2 – Fig. 9). The evolution of the secondary mineral assemblage is attributed to a change in the dominant reactions between the early and late stages of the experiment. Initially, the high pH of the leachates caused the dissolution of primary silicate minerals and formation of C–(A)–(K)–S–H in the Ca²⁺ rich solutions (Gaucher and Blanc, 2006; Savage, 2011). Between 15 months and 15 years of reaction (Phase 2) significant dedolomitisation occurred and the early formed C–(A)–(K)–S–H destabilised (Fig. 9) which led to the formation of secondary CaCO₃ and Mg–(Al)–(K)–silicates.

Previous studies of high pH-rock alteration have identified C–(A)–(K)–S–H, feldspars and zeolites as the dominant secondary phases likely to form in a GDF environment over time (e.g. Gaucher and Blanc, 2006; Savage, 2011). The formation of Mg–(Al)–(K)–silicates has only been suggested in a few studies of CDZ-type alteration as a minor product (Cuevas, 2004; Techer et al., 2012) and has not been widely considered. This study has shown that, in the presence of the common rock forming mineral dolomite, these phases may form and become the dominant secondary phases in a GDF scenario. An additional implication of this finding is that other sources of Mg²⁺ may exist at a GDF site (e.g. magnox fuel cladding within the waste) which could potentially contribute to the formation of secondary Mg–(Al)–(K)–silicates.

This study also found that the secondary mineral phases resulting from high pH rock alteration may increase the rock’s sorption capacity for U(VI).

At a broader level, the findings of this study illustrate that, in the CDZ around a GDF, there is potential for different mineral alteration reactions to occur over time. This may result in several different, transient mineral assemblages forming. The similar characteristics of the secondary phases produced in the two leachate systems studied here suggests that the alteration processes and resulting secondary mineral assemblages may not be significantly affected by the evolutionary stage of the cement leachate. However, the greater extent of alteration observed in the YNFP system, and the resulting increase in uranium sorption to the YNFP altered material observed, indicate that the composition of the leachate could impact radionuclide transport. Indeed, even at the relatively short timescale investigated here, the potential for significant mineralogical evolution has been demonstrated. It is also important to note that the Mg–clay mineral assemblage identified in this study cannot be shown to be a stable equilibrium assemblage, but may also be a metastable assemblage such as the C–S–H phases identified initially. The question is therefore raised as to whether
further reaction may be anticipated. This work has important implications for any GF safety case and highlights the need for longer-term experimental programmes to be considered during the implementation of geological disposal, and the need to develop the experimental-modelling interface so that predictions of CDZ evolution are robust.

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Appendix A. Supplementary material

Supplementary data associated with this article can be found, in the online version, at http://dx.doi.org/10.1016/j.apgeochem.2014.08.003.

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