

A long-term experimental study of the reactivity of basement rock with highly alkaline cement waters: Reactions over the first 15 months

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ABSTRACT

A series of long-term laboratory experiments was started in 1995 to investigate longer-term dissolution/precipitation reactions that may occur in the alkaline disturbed zone surrounding a cementitious repository for radioactive waste. They consist of samples of UK basement rock reacting with either Na-K-Ca-OH water ('young' cement porewater) or Ca-OH water ('evolved' cement porewater) at 70°C. This paper summarizes results of reactions occurring over the first 15 months. Experiments of both fluid types showed many similar features, though primary mineral dissolution and secondary mineral precipitation were more extensive in the experiments involving Na-K-Ca (younger) cement porefluids compared to more evolved (Ca-rich) cement porefluids. Dissolution of dolomite, and to a lesser extent silicates (probably K-feldspar, but also possibly mica) occurred relatively rapidly at 70°C. Dolomite dissolution may have been a key factor in reducing pH values, and may be a key mineral in controlling the extent of alkaline disturbed zones. Dissolution was followed by precipitation of brucite close to dolomite grains, at least two generations of C-S-H phases (which may have contained variable amounts of K, Al and Mg); overgrowths of calcite; small crystals of hydroxyapophyllite; and elongate crystals of celestite. Though hydroxyapophyllite was observed (a phase commonly associated with zeolites), there was no evidence for the formation of zeolites in the experiments. Fluid chemical changes track the mineralogical changes, with C-S-H phases being a major control on fluid chemistry. In the 'young' porewater experiments there were decreases in pH, and K, Ca and Mg concentrations, together with transitory increases in SiO₂ concentrations. In the 'evolved' porewater experiments there were decreases in pH, Mg, Ca and Sr concentrations, together with small increases in K and SiO₂ concentrations. A number of experiments are still running, and will be sampled in coming years.

KEYWORDS: alkaline fluids, cement porewaters, alkali disturbed zone, laboratory experiments, repository evolution, radioactive waste, cement.

Introduction

THE current UK concept for the disposal of low- and intermediate-level wastes (L/ILW) involves a mined geological disposal facility (GDF) several hundreds of metres below the surface (Crossland, 1998; Nirex 1989; NDA,

2010). The waste will be encapsulated in a cement grout matrix within steel or concrete containers, and placed in disposal vaults backfilled with a cement-based material. One of the main purposes of the cement is to maintain high pH conditions (>pH 12.5) that will reduce radionuclide solubility and microbial activity. Cement-equilibrated water will migrate into the surrounding host rock forming a plume of elevated pH, alkaline water, forming a region known as the 'alkaline disturbed zone' (ADZ) (Atkinson, 1995; Savage, 1995, 2011). The pH of the cement porewater and

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of the alkaline plume will decrease over time (Atkinson, 1985; Baston *et al.*, 2012) as differing cement components buffer the pH (Fig. 1). The alkaline cement leachate will react with the rock causing dissolution of some minerals and precipitation of others, and this will potentially modify both the geochemical and hydrogeological properties of the host rock within the ADZ prior to any eventual radionuclide release (e.g. Baker *et al.*, 2002; Rochelle *et al.*, 1992; Savage, 1995; Savage *et al.*, 1992; Savage and Rochelle, 1993). Understanding these geochemical interactions is important for predicting changes to rock porosity, fracture widths (and hence permeability and changes in groundwater flow), and minerals that may have an effect on diffusivities and sorption characteristics.

One area of uncertainty in our understanding of alkaline fluid-rock reactions is how to link data from well-constrained (but short-term) laboratory experiments with data from less well-constrained (but very much longer-term) industrial or natural analogue sites (e.g. Thomassin and Rassinoux, 1992; Savage, 2011). Predictive computer modelling is one way to achieve this, but the outputs from such modelling are highly dependent upon the databases used, and the codes may not adequately represent the complexities of real systems. Uncertainty can also be reduced by running laboratory experiments for timescales well in excess of just a few months to produce a more accurate and quantitative understanding of reactions within the ADZ.

In July 1995 as part of the Nirex Safety Assessment Research Programme (NSARP), a series of long-term experiments was started to study the changes in chemistry, mineralogy and sorption properties resulting from the reaction of cement-equilibrated waters

with Ordovician lower-greenschist facies Borrowdale Volcanic Group (BVG) volcano-sedimentary rocks, which were being considered as a potential host rock for a UK GDF. At this time, there was some uncertainty as to the nature of the secondary phases that may form over time. For example, calcium silicate hydrate (C-S-H) phases are common products of alkaline fluid-silicate mineral reactions, but their Ca:Si ratio, degree of hydration and degree of crystallinity can change over time (Atkinson *et al.*, 1995). Additionally, there was uncertainty as to whether zeolite minerals would form. These minerals have been found to occur within a range of natural alkaline environments (e.g. Sand and Mumpton, 1978; Hay, 1986; Savage, 1998, 2011; and references therein), as a product of reaction of aggregate within Portland cement (Marfil and Maiza, 1993), and as products within alkaline fluid-rock laboratory experiments at a range of temperatures (Knauss *et al.*, 1985; Knauss, 1987; Chermak, 1992, 1999; Adler *et al.*, 1999). They have also been predicted to form within alkaline fluid-mineral experiments (Bateman *et al.*, 1999) and the ADZ (Savage *et al.*, 1992; Savage and Rochelle, 1993; Soler, 2003; Gaucher *et al.*, 2004; Soler and Mäder, 2007; Watson *et al.*, 2009; Savage, 2011) – especially as the pH of the alkaline plume decreases. However, they had not been definitively identified as products of the reaction between cement porewaters and samples of UK basement rock. If zeolites did form in the ADZ, then their large surface areas and ion-exchange properties may be beneficial in the retardation of radionuclide migration (Pansini, 1996).

The experiments were originally designed to run for about two years, and were sampled and analysed after 4, 9 and 15 months reaction. This paper

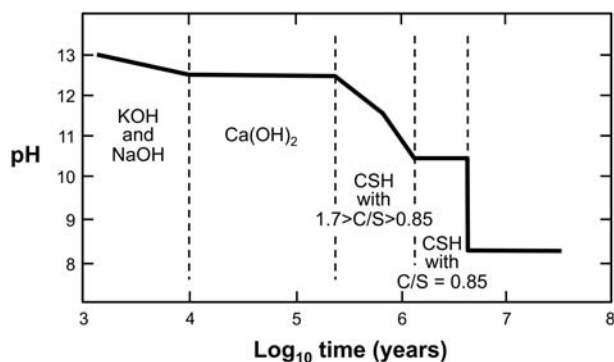


FIG. 1. Schematic illustration of the evolution of cement pore fluid pH as a function of time (redrawn after Atkinson (1985) for a repository diameter of 20 m containing 185 kg m^{-3} of cement, a temperature of 25°C , and with a groundwater flux density of $1 \times 10^{-10} \text{ m s}^{-1}$).

summarizes the setup and results from these experiments, plus an additional set of experiments that ran for 12 months. Several ‘back-up’ experiments were also started in 1995, and ran alongside those initially-sampled experiments. Following the rejection of the Nirex Rock Characterization Facility (RCF) planning application in March 1997 and the consequent cessation of the NSARP in the late 1990s, the remaining experiments were left unopened, but they were kept running. Some of these experiments have now been examined after 15 years of reaction and are the focus of a companion paper to the one presented here (Moyce *et al.*, 2014). The remaining experiments are still running, and will be sampled and analysed over the coming years. In time, the entire suite of experiments will provide a valuable and unique set of data to aid our understanding of decadal-scale alkaline fluid-rock reactions, and be a valuable resource with which to parameterize geochemical models.

Materials and methodology

A total of 32 experiments were assembled in 1995, comprising 16 that contained alkaline fluid and granulated BVG rock, and 16 ‘experimental blanks’ containing just alkaline fluid. Two different alkaline fluids were used (described in the following section), and consequently, the experiments could be grouped into ‘sets’ of four. A detailed description of the experiments is given in Rochelle *et al.* (1997) and a revised and updated summary of these is presented below. Subsequently, a further four sets of experiments were started (in 2013) to help address large gaps in time between some samples, and data from one of these (of 12 months duration) is also presented below.

Starting solutions

Two different synthetic cement porewaters were used, their compositions reflecting: the initial cement leachate dominated by dissolved NaOH and KOH and saturated with $\text{Ca}(\text{OH})_2$ – termed ‘young’ near-field porewater (YNFP); and an ‘evolved’ near-field groundwater (ENFG) representing a synthetic deep groundwater equilibrated with cement that was saturated with $\text{Ca}(\text{OH})_2$ and high in Na/Ca-Cl and Na_2SO_4 (Tables 1 and 2). The experimental fluid compositions were based on porewater compositions predicted by Bond and Tweed (1995) and Bond *et al.* (1995). They were prepared by the addition of pure chemical reagents to

de-aired, de-ionized water under a nitrogen atmosphere to minimize uptake of atmospheric CO_2 .

Starting rock sample

The disaggregated rock sample used in these experiments (sample NSF14A/30/P4) came from Sellafield borehole BH14A, west Cumbria (UK grid reference NY 0248 0569) at 858–859 m depth (below rotary table). It was selected to be mineralogically representative of material that would be encountered by present-day deep groundwaters in the fracture-controlled flow system in west Cumbria (Milodowski *et al.*, 1995; 2002). The fracture mineralization in the Sellafield boreholes has been described in detail by Milodowski *et al.* (1998, 2002). The sample contained a relatively high proportion of fracture mineralization and associated altered BVG wallrock from two potential flowing features identified in borehole BH14A (Fortey and Shepherd, 1997): (1) Feature No. 94/10 was a large fracture with vuggy cavities up to 20 cm long occurring in red, hematite-impregnated BVG wallrock. It was mineralized by coarse crystals of cream-to-pink-zoned euhedral ferromanganous dolomite and ankerite, and ferromanganous calcite up to 10 mm, with minor illite clay and fine earthy and specular hematite. (2) Feature No. 94/12 comprised a vein illitic clay and hematite.

The BVG host rock for this fracture mineralization consisted of a medium-grained, massive, volcanoclastic sandstone/tuff composed predominantly of quartz and K-feldspar, with minor albite, muscovite, chlorite and hematite.

Approximately 4 kg of the above sample was crushed gently to a 125–250 μm grain size, with as much fine material as possible removed by delicate sieving. Quantitative X-ray diffraction analysis of the major components of the sample showed it contained (in wt.%) ~41% quartz, 29% dolomite, 13% mica, 12% orthoclase, 3% calcite and 2% hematite. The dolomite, calcite, hematite and most of the mica came from the fracture mineralization. The high proportion of dolomite contrasts to that used in some other studies, but is typical for the mineralogy of groundwater flow paths associated with many parts of the BVG in west Cumbria (Milodowski *et al.*, 1995, 2002).

Experimental equipment

The equipment consisted of 150 ml (for the experiments) and 100 ml (for the blanks),

TABLE 1. Summary of fluid chemical information for YNFP experiments. Data are presented for the composition of the starting fluid, together with final compositions of YNFP+BVG experiments and YNFP 'blanks' at 4, 9, 12 and 15 months.

Starting fluid = synthetic 'young near-field porewater' (YNFP)									
Temperature 70 °C									
Time (days)	0	134	134	276	276	366	366	442	442
Approximate time (months)	0	4	4	9	9	12	12	15	15
Run number	–	486	487	488	489	1521	1522	490	491
Vessel number	–	B497	B212	B498	B213	B362	B207	B500	B214
Experimental charge	–	Crushed BVG	Blank	Crushed BVG	Blank	Crushed BVG	Blank	Crushed BVG	Blank
Weight of BVG	–	35.0000 g	–	34.9999 g	–	35.0002 g	–	35.0001 g	–
Initial weight of fluid	–	140.01 g	102.00 g	140.02 g	102.01 g	139.998 g	101.994 g	139.98 g	101.99 g
Measured Eh at 70°C (mV)	n/a	–115.5	–56.6	–71.5	–2.0	–58.2	–77.7	n/a	n/a
Corrected Eh at 70°C (mV)	n/a	38.5	97.4	82.5	152	146	126	n/a	n/a
pH measured at 25°C	12.96	12.03	13.12	11.79	12.93	11.93	13.03	11.75	12.94
pH recalculated to 70°C	11.67	10.79	11.83	10.60	11.64	10.69	11.74	10.56	11.65
Ca	51.8	1.84	24.8	0.840	12.6	<7.00	<7.00	1.27	7.33
Mg	0.089	<1.00	<1.00	<1.00	<1.00	<0.50	<0.50	<0.20	<0.20
Na	1890	1870	1770	1760	1810	2014	1938	2010	1890
K	3230	2750	3450	2410	3160	2323	3005	2230	2670
OH	2520	1064	2604	884	2680	823	2542	795	2674
CO ₃	166	2225	144	2590	129	1307	n/a	2769	226
HCO ₃	n/a	n/a	n/d	n/a	n/a	n/a	n/a	n/a	n/a
Cl	63.7	298	85.8	423	131	280	0.588	412	123
SO ₄	<1.00	20.8	0.800	31.4	2.09	31.6	0.527	52.7	<0.50
NO ₃	<1.00	<1.00	<0.50	0.800	0.600	0.344	0.052	<1.00	<0.50
Br	<0.50	<0.50	<0.25	<1.50	<1.50	0.562	0.060	0.416	<0.15
Total S	n/a	16.4	1.70	8.72	<2.00	<17.0	<17.0	9.96	<0.40
F	19.0	30.0	24.8	25.8	25.8	0.432	n/a	23.6	27.3
Si	9.20	1320	2.37	1090	2.97	1033	6.00	988	4.72
SiO ₂	19.7	2824	5.07	2332	6.35	2211	14.0	2114	10.1
Ba	<0.010	<0.010	<0.012	<0.010	<0.010	<0.004	<0.004	<0.002	0.004
Sr	0.018	0.012	0.032	<0.010	<0.010	<0.024	0.038	0.014	0.016
Mn	<0.010	<0.010	<0.010	<0.010	<0.010	0.004	<0.002	<0.002	<0.002
Total Fe	<0.10	<0.10	<0.10	<0.10	<0.10	0.013	0.019	<0.02	<0.02
Al				<0.20	<0.20			<0.04	<0.04
B				<0.50	<0.50			0.123	<0.10
Li				<0.05	<0.05			<0.01	<0.01
U				<0.20	<0.20			<0.1	<0.1
Th				<0.20	<0.20			0.400	0.500

Concentrations in mg/L except for U and Th which are in µg/L.

All solutions were filtered through a 0.2 µm filter prior to analysis.

n/a signifies that no data are available.

Eh corrected for filling solution 90–00–11 = measured Eh + 224 mV – temperature (°C).

< signifies concentrations below the detection limit.

EXPERIMENTAL STUDY OF CEMENT POREWATER INTERACTION

TABLE 2. Summary of fluid chemical information for ENFG experiments. Data are presented for the composition of the starting fluid, together with final compositions of ENFG+BVG experiments and ENFG ‘blanks’ at 4, 9, 12 and 15 months.

Starting fluid = synthetic ‘evolved near-field groundwater’ (ENFG)									
Temperature 70°C									
Time (days)	0	134	134	276	276	366	366	442	442
Approximate time (months)	0	4	4	9	9	12	12	15	15
Run number	–	470	471	472	473	1530	1529	474	475
Vessel number	–	B362	B207	B363	B208	B498	B433	B364	B209
Experimental charge	–	Crushed BVG	Blank	Crushed BVG	Blank	Crushed BVG	Blank	Crushed BVG	Blank
Weight of BVG	–	35.0002 g	–	35.0000 g	–	34.919 g	–	35.0001 g	–
Initial weight of fluid	–	140.02 g	101.98	140.01 g	102.02 g	140.040 g	102.001 g	140.00 g	102.02 g
Measured Eh at 70°C (mV)	n/a	44.7	–92.0	–3.0	–38.0	–89.1	–28.4	–95.0	–188.5
Corrected Eh at 70°C (mV)	n/a	199	62.0	151	116	115	176	59.0	–34.5
pH measured at 25°C	12.22	9.82	12.14	9.75	11.6	9.63	12.18	9.66	12.01
pH recalculated to 70°C	10.84	8.87	10.76	8.82	10.21	8.72	10.81	8.75	10.62
Ca	1930	1400	1970	1340	1950	1244	1799	1530	2090
Mg	0.117	<1.00	<1.00	<0.10	<0.10	<0.50	<0.50	<0.20	<0.20
Na	9160	9550	9110	8630	8850	9954	9624	9110	9300
K	185	197	156	204	176	198	169	236	214
OH	459	11.0	493	n/a	451	<20.0	444	n/a	473
CO ₃	<20.0	9.00	n/a	n/a	39	n/a	39.0	n/a	40
HCO ₃	n/a	n/a	n/a	47	n/a	n/a	n/a	50	n/a
Cl	15,100	16,270	15,100	16,700	16,400	17,030	16,305	15,796	15,659
SO ₄	1090	1060	1090	1060	1160	1038	1123	1084	1182
NO ₃	<20.00	<20.0	<20.0	<10.0	<10.0	<12.0	<12.0	<5.00	<5.00
Br	23.2	25.6	25.2	24.8	22.7	14.6	15.6	27.5	27.3
Total S	n/a	364	378	326	369	373	410	330	382
F	0.03	0.45	0.07	0.480	0.060	<2.0	<2.0	0.550	0.080
Si	2.07	32.4	<0.20	28.5	0.090	28.4	0.300	28.1	0.050
SiO ₂	4.43	69.3	<0.43	61.0	0.193	60.8	0.700	60.0	0.107
Ba	0.017	<0.010	0.022	0.005	0.013	0.014	0.297	0.006	0.017
Sr	166	80.8	174	65.0	180	54.8	143	66.3	184
Mn	<0.010	<0.010	<0.010	<0.001	<0.001	<0.002	<0.002	<0.002	<0.002
Total Fe	0.120	<0.10	<0.10	<0.01	<0.01	<0.013	<0.013	<0.02	<0.02
Al	4.17	<0.50	<0.50	0.259	0.316	0.008	0.107	0.396	0.430
B	0.335	<0.50	<0.50	0.097	<0.05	<0.144	<0.144	0.110	<0.10
Li	0.153	3.93	<0.10	3.01	0.150	2.66	<0.024	2.84	0.154
U	n/a	<0.20	<0.20	<0.1	<0.1	<0.00007	<0.00007	<0.1	<0.1
Th	n/a	<0.20	<0.20	<0.4	<0.4	<0.00009	<0.00009	<0.2	<0.2

Concentrations in mg/L except for U and Th which are in µg/L.

All solutions were filtered through a 0.2 µm filter prior to analysis.

n/a signifies that no data are available.

Eh corrected for filling solution 90–00–11 = measured Eh + 224 mV - temperature (°C).

< signifies concentrations below the detection limit.

Teflon[®]-lined, stainless-steel digestion vessels. The experiments were assembled and sampled in a nitrogen (CO₂-free) atmosphere inside a glove box. The wetted Teflon[®] parts of the equipment were first cleaned by boiling in dilute hydrochloric acid, and washed in de-ionized water followed by boiling in de-ionized water. An accurately known quantity of crushed BVG (~35 g) was weighed carefully into each vessel, followed by ~140 g of either synthetic YNFP or ENFG. For the blanks, ~100 g of either porewater was added to the vessels. The lids of the vessels were fastened down securely prior to removal from the glove box. The combined volume of the alkaline solution and crushed BVG was just less than that of the vessel, so that a small N₂ gas headspace/bubble resided at the top of the experiment. The experiments were placed in a thermostatically controlled oven at 70°C (accurate to ± 0.5°C), and were at approximately atmospheric pressure. The vessels were gently shaken/inverted by hand at regular intervals to ensure uniform reaction (approximately three times per week for the duration of the tests reported here). The presence of the N₂ gas headspace/bubble helped ensure complete mixing of the fluid and solid.

The initial fluid:rock mass ratios in the experiments were 4:1. Though the volume of the water-conducting features in the original sample of BVG core were small relative to the amount of rock, groundwater movement would result in exposure of this rock to a far larger volume of water over time. The relatively high fluid:rock ratio in the experiments was therefore chosen to represent advection of a significant amount of cementitious water through the water-conducting features. Under *in situ* conditions, the actual amount of fluid reacting with the rock will be significantly dependent on fracture properties and fluid flow rates, and we acknowledge the uncertainties in what these values may be. In terms of an experimental study, the higher fluid:rock ratio also made for more straightforward analysis of reaction products – if a significantly lower ratio had been used, then there would have been less fluid available for reaction, resulting in more subtle mineral dissolution features and much reduced precipitation of secondary phases.

Fluid sampling and analysis

Sampling of fluids and solids was carried out within a nitrogen-filled glove box and performed as soon as possible after the reaction vessels left the oven. Once in the glove box, each reactor was maintained at as

close to 70°C as possible in order to limit artefactual dissolution/precipitation reactions.

On sampling the reacted fluids, each of them was split into several subsamples, with the analysis of pH, Eh and bicarbonate/carbonate being conducted as soon as possible. For the 4-, 9- and 15-month duration experiments, pH was measured using an Orion[®] 520A pH meter calibrated using Whatman[®] NBS traceable buffers at pH 7, 10 and 13, and Eh was measured using an Orion 96–78 platinum electrode (with an internal silver/silver chloride reference electrode) connected to an Orion SA250 meter. For the 12-month duration experiments, pH was measured using a Thermo Scientific Orion VersaStar meter with an Orion 9103BNWP semi-micro combination pH electrode calibrated using Whatman[®] NBS traceable buffers at pH 7, 10 and 12.46, and Eh was measured using a Hanna Instruments HI3230 platinum electrode (with an internal silver/silver chloride reference electrode) connected to a Thermo Scientific Orion VersaStar meter. For the data presented here, room-temperature pH measurements and mineral saturation states were recalculated to those present in the experiments at 70°C using *PHREEQC* (version 3, Parkhurst and Appelo, 2013) with the Thermochemie database (version 9a, and implemented as SIT.dat [specific ion interaction theory]). This results in slightly different pH values to those originally reported by Rochelle *et al.* (1997) using the EQ3/6 geochemical modelling software, which appears to be due to the differing species included in the databases.

A subsample of ~8 ml was taken using a polythene syringe and filtered using a 0.2 µm and 0.4 µm Nuclepore[™] syringe filter, into a polythene tube for the subsequent analysis of major and minor anions by ion chromatography (only the 0.2 µm data are presented in this paper, but they are similar to the 0.4 µm data – see Rochelle *et al.*, 1997). An accurately known, and similarly filtered, subsample of ~8 ml was taken, and placed into a polystyrene tube for the subsequent analysis of major and minor cations (by ICP-OES and ICP-MS). Initially the ICP samples were acidified with 0.08 ml of concentrated 'ARISTAR[®]' nitric acid as a standard preservation procedure. However, after initial analysis of the acid-preserved samples, it was found that the results for high concentrations of Si (mainly the YNFP samples) were not consistent with each other, and that instrumental problems occurred for some samples. It was noted that if analysis was conducted within a relatively few hours of sampling and preservation, the analytical

equipment performed well, and measured Si concentrations were higher. However, if the 'preserved' samples were left in cold storage for several weeks (following standard procedures at the time), the analytical equipment did not perform well, and measured Si concentrations were only ~10% of the expected values. It is noteworthy that the equipment operator also described these solutions as appearing to have an 'oily' viscosity.

High concentrations of Si are stable in alkaline solutions, and it appears that acidification, a usual method of sample 'preservation', actually causes problems. We believed that acidification initiates Si polymerization, or gel formation, which increases the viscosity of the sample, and impairs analysis. This process appears to be kinetically controlled, rapid analysis does not allow the formation of larger Si 'structures', whereas at longer timescales these had formed to a sufficient extent to cause problems. To avoid this problem, Si analysis (by ICP-OES) was carried out on the highly alkaline 'unpreserved' samples, with standards also being prepared in alkaline medium. Although such an approach might not be suitable for many solutions, it appears very suitable for our Si-rich samples, mainly due to its high solubility in alkaline solutions. Samples from the ENFG experiments contained much less Si in solution, did not suffer problems during analysis, and hence were analysed by standard methods (sample preservation by acidification prior to analysis by ICP-OES).

Note that the data reported in Tables 1 and 2 were generated over several years. Whilst the analyses passed laboratory quality assurance criteria, there are slight differences in analytical detection limits between samples because of instrument changes and slightly different calibration procedures over time.

Solids sampling and analysis

After dismantling the experiments, the solid remaining at the bottom of each vessel was removed and subsampled: some were kept damp and sealed in an airtight container, whilst some were frozen rapidly in liquid nitrogen prior to freeze drying in an Edward's Modulyo freeze drier. The latter procedure was found to be more appropriate for subsequent mineralogical characterization.

Surface area analysis

The surface areas of subsamples of both unreacted and reacted disaggregated BVG material were

determined by the nitrogen BET adsorption/desorption method (Brunauer *et al.*, 1938) using a Micromeritics Flowsorb II 2300 analyser. The samples were prepared by degassing at 30°C in a vacuum oven for a minimum of 24 h to remove water vapour and adsorbed gases prior to analysis. Surface area data were obtained by cycling the sample from room temperature to liquid nitrogen temperature (at ~-196°C) whilst immersed in a 30:70 molar ratio stream of nitrogen and helium. Due to the highly hydrated nature of potential secondary reaction products it is possible that some dehydration and consequent change in structure will have occurred. However, this effect is essentially unavoidable.

Scanning electron microscopy

Unreacted and reacted BVG samples were examined by scanning electron microscopy (SEM) and environmental scanning electron microscopy (ESEM), using energy-dispersive X-ray microanalysis (EDXA) to aid phase identification on the basis of qualitative microchemical information. Samples of disaggregated material were mounted on aluminium SEM stubs using double-sided carbon adhesive tabs. These were then coated with a thin layer of carbon (~25 nm) in order to make the surfaces electrically conductive prior to examination by secondary electron imaging. Initial SEM-EDXA was carried out using a Cambridge Instruments Stereoscan S250 SEM instrument equipped with a Link Systems 860 EDXA system. Subsequently, more detailed observations were recorded using either: a LEO 435VP variable pressure SEM instrument with an Oxford Instruments ISIS 300; or a FEI QUANTA 600 ESEM instrument with a FEI QUANTA 600 tetrode gun, and equipped with an Oxford INCA 450 EDXA system. The SEM observations were made using an electron beam accelerating voltage of 15–20 kV.

Polished thin sections were also prepared from the reacted BVG material (following vacuum impregnation with epoxy resin) and examined by back-scattered scanning electron imaging (BSEM). However, in most cases, observations were limited by the very small amounts of reaction product intersected in the polished sections.

Analytical transmission electron microscopy

Analytical transmission electron microscopy (ATEM) of the reaction products was undertaken using a Jeol 2010 TEM fitted with an Oxford

Instruments ISIS 200 EDXA system (Nottingham Trent University). TEM with EDXA and selected-area electron diffraction (SAED) was conducted using an FEI Tecnai TF20 FEGTEM with an Oxford Instruments INCA 350 EDXA system with 80 mm X-Max SDD detector and a Gatan Orius SC600A CCD camera (University of Leeds). Samples were prepared for TEM by ultrasonication in ethanol to liberate reaction products from the altered BVG grain surfaces. The resultant suspension was then deposited onto copper TEM grids with holey carbon films.

X-ray diffraction analysis (XRD)

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, with a cobalt-target tube, and X'Celerator detector operated at 45 kV and 40 mA. They were scanned at 4.5–85°2θ at 2.76° 2θ/min. The diffraction data were analysed using PANalytical X'Pert Highscore Plus version 2.2e software coupled to the latest version of the International Centre for Diffraction Data (ICDD) database. After identification of the mineral species present quantification was determined by Rietveld modelling (Snyder and Bish, 1989). Although different mica species may be present in these materials, these could not easily be differentiated and for simplicity were described simply as 'mica'.

Results

Analyses of the solutions reacted for 4, 9, 12 and 15 months are given in Tables 1 and 2. The following section gives a summary of mineralogical observation for samples from the 4-, 9- and 15-month duration experiments. Tables 1 and 2 also show analyses of experimental 'blanks' (i.e. from solutions kept in Teflon®-lined reactors for several months, but with no BVG present), plus analyses of de-ionized water (DIW) that served as an indicator of any sample handling/analytical issues. A selection of the data from the experiments is plotted in Fig. 2.

Note that for most of the experimental 'blanks' there are small decreases in concentrations of Ca and Si (± Al and K). These decreases are minor compared to changes in the experiments containing crushed BVG, but they are consistent with potential slight initial oversaturation with respect to C-S-H

phases (plus related calcium aluminosilicate hydrate [C-A-S-H] phases) and hence their precipitation. This may have originated from 'uncertainties' in the thermodynamic data for the C-S-H phases used in the modelling packages that predicted the starting fluid compositions. The continued decrease in concentrations over time may reflect not just continued precipitation, but also a shift towards more-ordered phases having lower solubility than initially-formed disordered gels, and some modelling work is included below to investigate this.

Reactions of young cement porewater (YNFP) with BVG

Changes in fluid chemistry

Sodium and chloride concentrations remain relatively stable throughout the 15 month duration of the experiments (Na at ~2000 mg L⁻¹, and Cl⁻ from ~64 mg L⁻¹ to 400 mg L⁻¹). The increase in chloride concentration may reflect some leaching from the rock sample – the deep groundwater from where the sample came from being appreciably saline. The relatively conservative behaviour of these elements is to be expected. Importantly however, these data indicate that significant loss of water had not occurred over the 15 month lifetime of these experiments (the steel vessels around the Teflon® liners appear to have restricted diffusional loss of water vapour). Water loss is a potential problem in any long-term experimental study as it could lead to enhanced precipitation (effectively an artefact of the experiment), increased salinity, and increased complexity in interpreting the results. This feature will be monitored in on-going, longer-term experiments.

For pH there was a noticeable drop from 11.7 (recalculated to 70°C) to 10.6 (recalculated to 70°C). The majority of this decrease occurred during the first 4 months of reaction, with a progressively slower rate of decrease after that. This decrease is consistent with alkaline fluid-rock reaction. This could include the dissolution of dolomite (shown in previous studies by Rochelle *et al.* (1994a, 1998a, e) to be effective at reducing pH, see Discussion) or the formation of C-S-H phases.

The likelihood of C-S-H phase precipitation is further enhanced by decreases in calcium concentrations from ~52 mg L⁻¹ to 1.3 mg L⁻¹ over the 15 months of the experiments. However, the small amounts of calcium initially in solution means that only limited C-S-H would have formed, unless

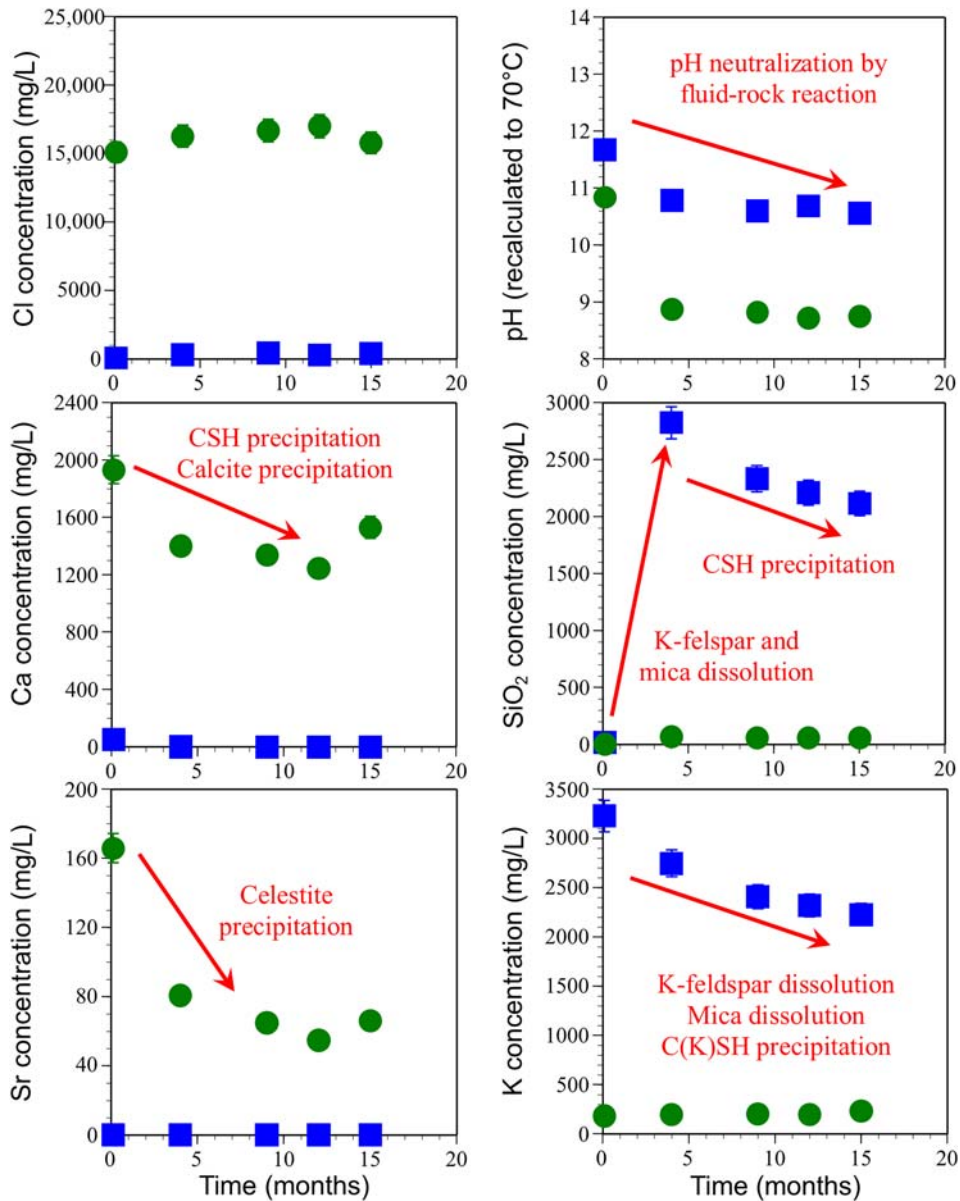


FIG. 2. Plots of selected fluid chemical data for the starting solution, and at 4, 9, 12 and 15 months. Squares represent the young porewater experiments, circles represent the evolved porewater experiments.

calcium was also contributed from dissolution of minerals. Silica concentrations show an increase from an initial value of 9 mg L^{-1} , to 2824 mg L^{-1} in the first 4 months. Subsequent to this however, the concentrations fall noticeably. Dissolution of (alumino)silicate minerals present within the BVG undoubtedly accounts for the large increase

in silica, with precipitation as secondary phases (such as C-S-H phases) probably accounting for the later, decreasing concentrations. It is probable that the decreasing concentration would stabilize at a steady state value if C-S-H were controlling silica concentrations, though the few data points do not allow for estimation of a value for this. The initial

rate of increase of silica concentration in solution appears high. This may reflect the reactivity of the BVG towards the YNFP, although it might, in part, reflect the physical state of the starting solid. Crushing the BVG prior to reaction will have produced many small (and hence highly reactive) grains. Although care was taken to remove most of these, it is still possible that some were adhering to the larger grains and were only eventually removed by dissolution into the YNFP. In addition, it is possible that larger grains of fault-gouge material disaggregated in solution, contributing to the proportion of fine-grained reactive material.

Potassium concentrations decreased from 3230 mg L⁻¹ to 2230 mg L⁻¹ over the 15 months of the experiments, signifying removal of potassium from solution by precipitation as a secondary phase. This interpretation is supported by the mineralogical analysis of the reacted solids (see below). Indeed, the decreasing rate of removal of potassium is consistent with a precipitation rate controlled by the concentration of potassium in solution and the formation of a K-rich C-S-H phase. However, this does not preclude the dissolution of K-rich minerals within the BVG sample, so long as their rate of supply of potassium to solution was less than its rate of removal by precipitation.

Aluminium concentrations were 12.2 mg L⁻¹ in the starting solution and decreased initially to values below the limit of detection (<0.5 mg L⁻¹). Although it is possible that dissolution of aluminosilicate minerals (such as feldspars – see Sections below) within the BVG was occurring (i.e. Al concentrations would be expected to increase over time), there appears to have been an efficient process removing aluminium from solution. Many previous studies of both natural and experimental systems (e.g. Cliff *et al.*, 1975; Richardson *et al.*, 1993; Linklater, 1998; Rochelle *et al.*, 1998*a,c*; Richardson, 2014; Myers *et al.* 2015) have noted in excess of 5 wt.% aluminium within C-S-H precipitates, so it is very possible that much of the released aluminium was being incorporated into secondary C-S-H identified by petrographic analysis (see below).

All the experiments had magnesium concentrations below the detection limits. This is consistent with previous studies that have shown that magnesium has very low solubility in alkaline solutions, and will precipitate rapidly as brucite (Bérubé *et al.*, 1990; Cheng, 1986; Rochelle *et al.*, 1994*b*). If, as seems probable, magnesium was released from the BVG (see comments about dolomite dissolution in the following sections),

then sufficient hydroxide ions were present to ensure its immediate precipitation. The solutions also contained relatively low concentrations of strontium that were close to the limits of detection.

Sulfate concentrations are generally low, but the steady increase from below detection (<1 mg L⁻¹) to ~53 mg L⁻¹ suggests dissolution of a S-containing phase. Unfortunately, no S-containing phase was identified during the initial characterization of the BVG sample. It must therefore be present in trace quantities, and so probably has a relatively fast dissolution rate to produce the observed increase in sulfate. This is consistent with either anhydrite or pyrite dissolution, both of which have relatively fast dissolution kinetics and commonly occur as fracture-filling phases within the BVG in the Sellafield area and elsewhere in west Cumbria (Milodowski *et al.*, 1998, 2002).

Whilst not a major component of the solutions, fluoride concentrations in the original starting solution and 4-, 9- and 15-month experiments are unexpectedly high, as are those in the experimental blanks. We consider it unlikely that fluoride leached from the Teflon[®] reaction vessel liners as they were thoroughly boiled in dilute hydrochloric acid and water prior to use. However, the 12-month-long experiment and blank show almost no fluoride present in solution (this experiment was initiated more recently, and with a different batch of starting solution). It seems probable therefore, that much of the fluoride in the 4-, 9- and 15-month experiments was an impurity in the starting solution. Although the original batches of chemicals used to make the starting solution were no longer available for examination, ‘trace’ fluoride is noted in current silica standard solutions (similar to that used to add silica to the original starting solution), and this appears to be the most probable source. The potential for dissolved fluoride to stabilize the formation of F-containing phases is discussed below.

Mineralogical observations

Significant mineralogical alteration of the BVG was observed in the YNFP experiments. Dolomite grains were particularly reactive and grains were strongly etched (Fig. 3*a*), with dissolution increasing progressively with time from 4 to 15 months. The etched dolomite surfaces were coated with a thin veneer of reaction product, which ATEM indicated to be predominantly fine (typically <1 µm), Mg-O-rich platy crystallites, which are identified as brucite (Fig. 3*b*). The presence of

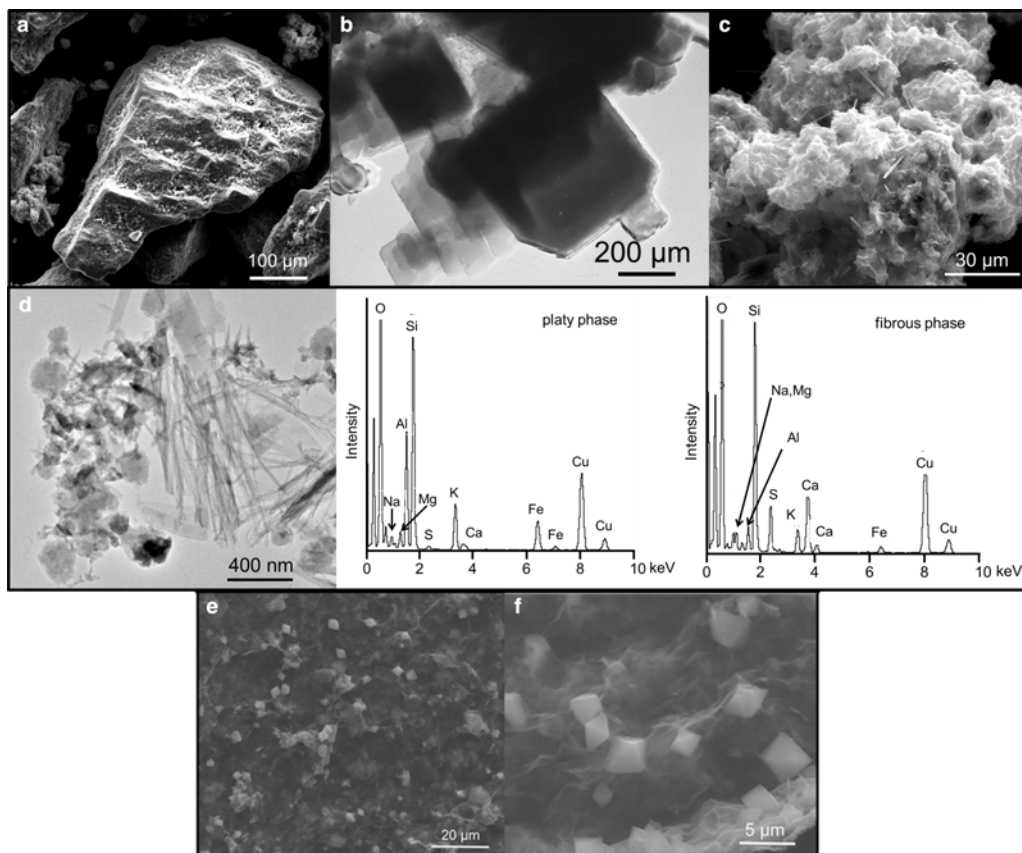


FIG. 3. (a) SEM (secondary electron) image of corroded dolomite grain (9 months YNFP experiment). (b) TEM image of well-developed brucite crystals adjacent to dolomite (9 months YNFP experiment). (c) SEM image showing extensive C-S-H reaction product coating altered BVG fragments (9 months YNFP experiment). (d) TEM image and EDXA spectra of fibrous C-S-H reaction product and primary platy illite (9 months YNFP experiment). (e) and (f) SEM images of euhedral tetragonal bipyramidal apophyllite crystals formed as reaction products (4 months YNFP experiment).

(relatively reactive) dolomite in these experiments could explain why the reduction in pH was so large, and the precipitation of brucite as a reaction product appears to have provided an initial sink for magnesium when the pH was at its highest. Minor silicate reaction is also evident with etching of quartz and K-feldspar. In most cases the extent of primary silicate mineral surface dissolution was difficult to assess because most grain surfaces were usually obscured by extensive fine-grained primary illite clay (or sericite). Results from SEM-EDXA showed that a secondary gel-like C-S-H reaction product was typically intimately mixed with, and difficult to resolve from, the primary illite or sericite fines on the grain surfaces that were originally present in the starting material.

Some quartz and feldspar grains locally showed much more extensive development of this C-S-H reaction product, which displayed a discrete wispy and sheet-like morphology, and formed an extensive honeycomb-like coating that loosely cemented the grains together (Fig. 3c). Significant amounts of K, Al and S were detected within this reaction product, but it was difficult to determine whether these elements were present within discrete C-S-H or if the SEM-EDXA data represented an unresolved mixture of secondary C-S-H and fine primary illite / sericite or other phases. In contrast, higher-resolution TEM / ATEM analyses were able to resolve more clearly the C-S-H phase from primary silicate mineral fines and showed that initially (after 4 months reaction) it comprised a

microfibrinous phase with a low Ca:Si ratio (Fig. 3*d*). ATEM analysis of individual C-S-H fibres still identified the presence of K, Al and S (Fig. 3*d*), which suggests that these elements are actually intrinsically incorporated within the C-S-H. It is possible that the K, Al and S might be partially contributed from amorphous material or soluble sulfate salts admixed with the fibrous C-S-H that is still unresolved at the ATEM scale. However, significant aluminium substitution for silicon can occur in C-S-H phases to produce calcium aluminosilicate hydrate (C-A-S-H) gels (e.g. Richardson *et al.*, 1993; Richardson, 2014; Myers *et al.*, 2015). It is also feasible that at least part of the potassium might be bound within the C-S-H phase, particularly when Ca:Si is low (e.g. Hong and Glasser, 2002; L'Hôpital, 2014). Furthermore, some previous research has indicated that potassium uptake by C-S-H is enhanced by aluminium substitution (Hong and Glasser, 2002), though this is refuted in other studies (e.g. L'Hôpital, 2014). This reaction product developed a more flaky morphology as reaction progressed (after 9 and 15 months). No electron diffraction patterns could be obtained from the C-S-H reaction products, which therefore appear to be amorphous or very poorly crystalline.

In addition to the K-Al-bearing C-S-H gel, tiny euhedral crystals of a K-Ca-silicate were also observed by SEM-EDXA as a reaction product in the 4 month experiment. These crystals have a distinctive tetragonal bipyramid morphology (Fig. 3*e*). Semi-quantitative EDXA indicated this phase has a K:Ca ratio of ~1:4. The morphology and composition are characteristic of hydroxyapophyllite (ideal formula $\text{KCa}_4\text{Si}_8\text{O}_{20}(\text{OH})_8\text{H}_2\text{O}$) (cf. Deer *et al.*, 1992). Hydroxyapophyllite was not observed in the 9- and 15-month experiments, and therefore appears to an early-formed transitory reaction product in these experiments. The initial formation of hydroxyapophyllite accompanied by K-Al bearing C-S-H in the first 4 months, subsequently followed by continued K-Al-bearing C-S-H precipitation explains the progressive decrease of potassium in the fluid whilst sodium (which is not incorporated in these phases) does not mirror potassium (Fig. 2). Due to their small size (<5 µm), it was not possible to analyse these tiny crystals for F content, and so we were unable to determine if they contained a fluorapophyllite component.

No evidence for zeolite formation was found in any of the experimental residues. Similarly, there was also no evidence for fluorite (CaF_2) in either the starting material or the reaction products. Whilst we

note that minor-to-trace amounts of fluorite can be found in some fracture assemblages within the BVG (Milodowski *et al.*, 1998, 2002), if it was not observed in either the starting material or the reaction products, to be present it must have occurred in minute concentrations (i.e. below the levels that we could identify with the techniques used).

Reactions of evolved cement porewater with BVG

Changes in fluid chemistry

Relative to the starting solution, there was a noticeable drop in pH from 10.8 to 8.7 (recalculated to 70°C). The majority of this decrease occurred during the first 4 months of reaction, with a progressively slower rate of decrease after that. After 15 months reaction the pH still appears to be decreasing. Such a decrease would be expected as a result of fluid-rock reaction and has been observed in previous studies under the NSARP using ENFG (Rochelle *et al.*, 1994*a*, 1998*a,e*). As with the YNFP experiments above, the magnitude of the pH decrease is probably due to relatively reactive minerals in the BVG, especially the dolomite present in the fracture mineral assemblage (see comments in Discussion section).

As with the YNFP experiments, sodium and chloride appear relatively conservative, and have approximately constant concentrations (~9000 mg L⁻¹ and ~16,000 mg L⁻¹ respectively). These relatively constant concentrations again suggest minimal loss of water during the experiments.

Calcium concentrations decreased from the initial value of 1930 mg L⁻¹ to 1340 mg L⁻¹ over the first nine months of reaction, but then showed a small increase to 1530 mg L⁻¹ after 15 months reaction. The initial decrease is not unexpected, as the relatively high pH of the solution would have favoured the formation of two potential secondary phases: calcite, and C-S-H phases. For the latter, the presence of silica from dissolution of minerals within the BVG would have caused their precipitation. This concurs with previous studies under the NSARP using ENFG reacting with BVG (Rochelle *et al.*, 1994*a*, 1998*a,c,e*), and is consistent with the YNFP experiments above. The ENFG experiments initially contained relatively little silica in solution, and the increase in silica from 2 mg L⁻¹ to 60 mg L⁻¹ suggests dissolution of (alumino)silicate minerals within the BVG (e.g. quartz, muscovite and orthoclase). Though there are no clear trends within the data, the generally low concentrations

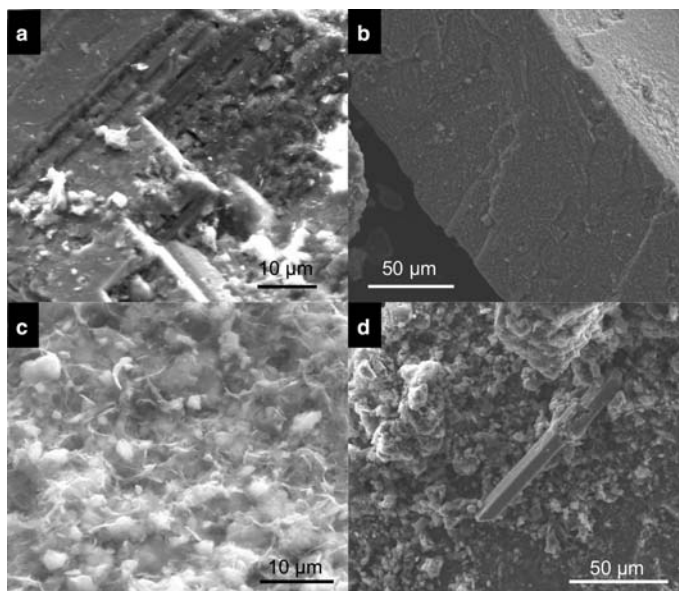


FIG. 4. (a) SEM (secondary electron) image of corroded dolomite grain displaying deep etching along cleavage (9 months ENFG experiment). (b) SEM image showing sugary coating of very fine secondary calcite overgrowths on reacted dolomite surface (9 months ENFG experiment). (c) C-S-H reaction product coating altered BVG fragments (9 months ENFG experiment). (d) Euhedral acicular crystal of celestite growing on the surface of a BVG fragment encrusted with fine particulate material cemented by C-S-H (9 months ENFG experiment).

in solution are consistent with C-S-H-type phase equilibrium (i.e. high calcium concentrations necessitate low silica concentrations).

Potassium concentrations are relatively low in the ENFG compared to the YNFP, but they do show a steady increase from 185 mg L^{-1} to 236 mg L^{-1} over the 15 months of the experiments. This increase probably reflects mineral dissolution in the BVG, the most probable candidates being K-feldspar and/or muscovite. Dissolution of these phases would have also contributed aluminium to solution, though concentrations remained low throughout the experiments. It is possible therefore, that an efficient process was removing aluminium from solution, potentially via incorporation into secondary phases, such as C-S-H or C-K-S-H gel-like phases noted earlier.

The decrease in strontium from 166 mg L^{-1} to 66 mg L^{-1} indicates precipitation of small amounts of a secondary phase. The trend in the data, with a falling rate of decrease, suggests an approach to a steady-state value in the region of 50 mg L^{-1} . This decrease is consistent with the observation of secondary celestite in the reacted mineral assemblage (see below). The driving mechanism for celestite precipitation may well have been initial

oversaturation with this phase, however, its formation after ettringite breakdown can not be ruled out (see below). Celestite precipitation would have been favoured by the relatively high concentrations of dissolved sulfate in the ENFG, which remained at $\sim 1060 \text{ mg L}^{-1}$ throughout the experiments.

As per the YNFP experiments, magnesium concentrations remained below detection limits, and these are consistent with previous studies showing that appreciable amounts of magnesium are not stable in alkaline solutions, and will rapidly precipitate as brucite (Bérubé *et al.*, 1990; Cheng, 1986; Rochelle *et al.*, 1994b).

Fluoride concentrations were much lower in the ENFG experiments ($< 1 \text{ mg L}^{-1}$) compared to the YNFP experiments, including the 'blank' runs. This appears to strengthen the case for the fluoride having originated from an impurity in the silica solution added to the starting solution of the YNFP experiments (silica was not added to the starting ENFG solution).

Mineralogical observations

Reaction of the BVG with the ENFG also resulted in corrosion and dissolution of dolomite (Fig. 4a). Although the dolomite reaction is dominated by dissolution in these experiments, the reacted

dolomite after 9 months showed evidence for the secondary formation of very fine grained (up to 2 μm) overgrowths of secondary calcite on the etched grains. These overgrowths appeared to be syntaxially (crystallographically) orientated to the underlying dolomite substrate, and gave the dolomite a 'sugary' appearance under the SEM (Fig. 4b).

Quartz, K-feldspar, albite and muscovite were also all observed to have reacted with ENFG after 9 and 15 months, but little alteration was evident in the 4 month experiment. The extent of secondary precipitation was more limited than in the YNFP experiments, probably because of the lower pH (i.e. lower reactivity) of the starting fluid. The principal reaction product is an amorphous Ca(\pm K,Al) silicate hydrate gel, which formed a very thin film over the primary silicate minerals (Fig. 4c). This phase was more abundant in the 4 month experiment compared to the 9- and 15-month experiments, which may indicate dissolution of this phase as the fluid pH decreased during the course of the experiments, as reaction progressed. Although apophyllite was identified tentatively as a trace phase by Rochelle *et al.* (1997) in the 4-month duration experiments, subsequent re-analysis of the very limited amounts of remaining sample showed no evidence for it. No zeolites or fluorite were observed.

Well-formed, acicular crystals of celestite (SrSO_4) were also found in the residues from the 4-, 9- and 15-month experiments (Fig. 4d) (this phase was originally miss-reported by Rochelle *et al.* (1997) as strontianite). The crystals range from 1 to 200 μm in size and typically display very smooth crystal faces. The precipitation of this phase explains the progressive decrease in strontium in the fluid from 4 to 15 months (Fig. 2).

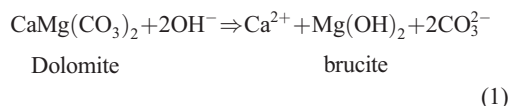
Discussion

This experimental study has aged samples of Borrowdale Volcanic Group material from a current, transmissive, groundwater flow zone, with young and evolved cement porewater at 70°C for 4, 9 and 15 months. Analyses revealed changes associated with dissolution of primary minerals and development of secondary phases.

Reaction of BVG with young cement porefluid (YNFP)

Initial fluid-rock reaction of these more alkaline fluids (i.e. dissolution of the starting BVG) is clearly demonstrated through a progressive reduction in pH

(from 11.7 to 10.6 when recalculated to 70°C). The majority of this decrease occurred during the first 4 months of reaction, and was probably aided by the preferential reaction of the finer grains of the BVG. The trend in pH suggests it is approaching a steady-state value of ~ 10.2 , though with prolonged reaction this would no-doubt eventually decrease further. This pH value is consistent with (possibly temporary) buffering by a secondary phase, probably the (K- and Al-rich) C-S-H phase. Such a decrease in pH has been observed in previous studies under the NSARP (Savage *et al.*, 1992; Rochelle *et al.*, 1994a, 1998a,b,e). Indeed, one of these studies (Rochelle *et al.*, 1998b) noted that dolomite was particularly influential in this respect due to its relatively fast reaction kinetics and tendency to react to form brucite (Mg-hydroxide), effectively removing hydroxyl ions from solution (the sample of BVG used in that study contained 25% dolomite, cf. 29% used in this study). Other studies have also noted the formation of brucite (and potentially calcite) as a result of NaOH-rich fluids reacting with dolomite (Cheng, 1986; Choquette, 1991; Hadley, 1961; Mohnot and Bae, 1985):



Reaction of dolomite was observed, with pitting on dolomite grains associated with platy crystals of brucite (Fig. 3a,b). Other grains (silicate minerals) showed only minor etching, though were covered with secondary C-(K-A)-S-H gel, and grains of gouge material appeared to have disaggregated to produce a large amount of fine particles. The latter are thought to have provided a large surface area for a reaction to occur. Carbonate mineral dissolution is faster than silicate mineral dissolution by several orders of magnitude (Brady and Walther, 1989; Chou *et al.*, 1989; Rochelle *et al.*, 1994b, 1998d), and the higher degree of pitting on dolomite is consistent with this.

Continued reduction in pH through fluid-rock reaction will result eventually in the destabilization of brucite, though this did not appear to have occurred during the 15 month duration of the experiments describe here. It is anticipated that this will eventually result in the re-release of magnesium to solution. It is noteworthy that Rochelle *et al.* (1998b) suggest the possible formation of a smectitic clay as a product of the reaction of YNFP with dolomite-rich BVG. Adler *et al.* (1999) also identify the dissolution of dolomite/ankerite as a source of magnesium to solution, and that this

helped drive clay mineral formation in their experiments. Longer-duration experiments would reveal if clays were a stable reaction product as the pH decreased (cf. Moyce *et al.*, 2014).

The dissolution of dolomite (\pm silicate minerals) will also have contributed calcium to solution. This will have added significant amounts of calcium to the limited amount in the starting fluid, and combined with silica released through the dissolution of silicate minerals will have helped form the extensive C-S-H gel precipitate.

The observation that measured calcium concentrations decreased over time suggests the rate of C-S-H precipitation was more than sufficient to keep pace with the input of calcium from the dissolving BVG. The solutions also contained large concentrations of dissolved silica which appears to have resulted from relatively fast dissolution of finer grains of BGS relatively early in the experiments. Previous work (Savage *et al.*, 1992) has already noted the inverse solubility relationship between calcium and silica for conditions of C-S-H phase saturation. The gel was not pure C-S-H, but contained abundant potassium and aluminium, and this is also consistent with C-S-H precipitates in previous experimental studies (Rochelle *et al.*, 1998*a,c*; Savage *et al.*, 1992). The potassium and aluminium were probably mainly sourced from K-feldspar and/or muscovite dissolution, which would have also contributed silica to solution. The C-K-A-S-H (calcium potassium aluminosilicate hydrate) phase probably controlled the solubility of calcium, potassium, aluminium and silica throughout most of the duration of the experiments.

The initial gel appeared to have a more 'fibrous' morphology, whilst the later gel appeared to have a more 'flaky' morphology and contained more aluminium. It is unclear if the change in morphology is linked to the change in composition, or the relative age of the C-S-H (i.e. it may slowly transform to more ordered structures over time). The gel may have been showing some signs of undergoing re-dissolution in the 15 month experiment. This may be a sign of the pH reduction through continued fluid-rock reaction causing the initially formed C-S-H to become unstable, possibly itself breaking down to form other phases (such as C-S-H of a different composition or clays) as the pH of the system evolved. Longer-duration experiments would reveal if this were the case (see Moyce *et al.*, 2014). The surface area of the reaction products increased relative to the starting material, reaching a maximum increase of 39% after 9 months reaction, though this later fell to a 30% increase after 15 months reaction.

This trend in surface area is consistent with the precipitation, and subsequent partial dissolution, of the C-K-A-S-H gel.

The dissolution of dolomite will also have increased the partial pressure of CO_2 ($p\text{CO}_2$) in the reactor, and there is evidence for higher concentrations of bicarbonate/carbonate in the longer duration experiments. Interestingly, Adler *et al.* (1999) conducted experiments at 30°C reacting Opalinus Clay with a Na-K-Ca-OH alkaline fluid at pH 13.2 (equivalent to a young porefluid) for between 4 and 18 months. They report that zeolite formation was suppressed by addition of CO_2 /bicarbonate/carbonate to solution, and similar observations were made by Thompson (1971). Adler *et al.* (1999) also report that zeolite formation was favoured in rock pore spaces, where fluid fluxes and fluid: rock ratios were very low. We sought evidence for zeolite formation, though none was found. This may possibly be explained by our experiments having a relative abundance of water, and having significant dolomite dissolution that increased inorganic carbon concentrations. The aluminium and silica needed for zeolite formation appear to have mainly entered the C-S-H (C-A-S-H, C-K-A-S-H) phases mentioned previously. The potential exists therefore, for zeolites to still form within the BVG, though they might require a carbonate-poor rock-dominated environment, and hence not be expected in water-conducting, carbonate-lined fractures.

Though zeolites were not found, tiny crystals of hydroxyapophyllite (ideal formula $\text{KCa}_4\text{Si}_8\text{O}_{20}(\text{OH})_8\text{H}_2\text{O}$) were found amongst the C-S-H gel, having a euhedral tetragonal bipyramidal morphology (Fig. 3e). Hydroxyapophyllite is commonly associated geologically with zeolites, such as in the alkaline alteration of basalts (e.g. Deer *et al.*, 1992). Only small quantities of this phase were found, and it is thought that it may have only been formed early in the experiments when calcium and potassium concentrations were relatively high and when silica concentrations were increasing. Detailed analysis of these small crystals proved difficult, and we were unable to determine if they contained a fluor-apophyllite component. However, had this been significant in this phase, then we would have expected fluoride concentrations in solution to increase once it re-dissolved – this was not observed. Indeed, the experiments at 4, 9 and 15 months show, if anything, a decreasing trend in fluoride concentrations (12 month data showed negligible fluoride).

The continuous, linear increase of sulfate throughout the experiments appears to indicate that certain

TABLE 3. Selected mineral saturation state data for YNFP experiments. Data are presented for the starting fluid, together with final compositions of YNFP+BVG experiments and YNFP 'blanks' at 4, 9, 12 and 15 months.

Time (days)	0	134	134	276	276	366	366	442	442
Run number	-	486	487	488	488	1521	1522	490	491
Experimental charge	-	Crushed BVG	Blank	Crushed BVG	Blank	Crushed BVG	Blank	Crushed BVG	Blank
Acmite						4.59	-2.18	-3.26	-0.82
Afwillite	2.34	-1.46	0.32	-3.61	-0.49				
Albite-high	-6.98					-0.24	-10.61		
Albite-low	-5.87					0.86	-9.50		
Analcime	-3.18					0.40	-6.50		
Aragonite	0.91	0.51	0.44	0.20	0.24		0.38	0.38	0.22
C ₂ SH(alpha)	1.51	-2.09	0.45	-3.56	-0.23		-3.33	-3.33	-0.51
CaCO ₃ H ₂ O(s)	0.09	-0.31	-0.38	-0.62	-0.58		-0.44	-0.44	-0.60
Calcite	1.09	0.68	0.61	0.38	0.42		0.55	0.55	0.39
Clinoptilolite-K	-13.07					-0.62	-17.59		
Cristobalite	-3.56	-0.36	-4.40	-0.24	-4.01	-0.35	-3.85	-0.25	-3.82
CSH _{0.8}	-0.77	-0.28	-1.69	-0.80	-1.73		-0.71	-0.71	-1.73
Ferryhydrite						-0.11	-1.00	-1.71	-0.56
Fluorite	-0.03	-1.29	-0.25	-1.82	-0.35	2.11	1.22	2.09	-2.25
Goethite	0.29	3.09	-2.45	1.87	-2.34	6.03	4.26		
Hematite						2.73	-20.99		
Heulandite-Na	-13.09					0.01	-14.54	-4.25	-1.44
Hillebrandite	0.58	-3.02	-0.48	-4.49	-1.15	0.53	-14.04		
HNonttronite-K						-0.39	-13.97		
HNonttronite-Na									
Illite-Fe ^{II}									
Jaffeite	4.41	-9.59	2.05	-14.14	-0.35			-13.41	-1.39
Jennite	6.65	-4.74	0.60	-11.19	-1.83			-10.12	-2.82
Lepidocrocite						1.81	0.92		
Maghemite(disord)						2.29	0.52		
Maghemite(ord)						2.08	0.31		
Magnetite						0.60	-2.56		
Microcline	-3.78					2.87	-7.46		
Nonttronite-K						-0.16	-14.72		
Nonttronite-Na						-0.61	-15.17	1.13	-2.83
Okenite	-1.43	1.57	-3.22	1.02	-2.97	2.65	-7.69		
Phillipsite-K	-4.00					1.74	-8.63		
Phillipsite-Na	-5.00								

Portlandite	-0.40	-3.80	-0.51	-4.60	-1.05	0.12	-3.38	-4.48	-1.28
Quartz	-3.09	0.11	-3.93	0.24	-3.54	2.54	-7.80	0.23	-3.35
Sandine	-4.11								
SiO _{2(am)}	-3.95	-0.74	-4.78	-0.62	-4.39	-0.74	-4.23	-0.63	-4.20
SrSiO ₃₍₆₎	-1.83	-0.76	-2.18				-1.66	-1.09	-2.17
Strontianite	-1.95	-1.09	-1.83					-0.99	-1.84
Tobermorite-11A	2.01	4.22	-3.58	0.97	-3.89			1.54	-3.94
Tobermorite-14A	3.14	5.35	-2.45	2.11	-2.76			2.68	-2.81
Trona	-12.33	-9.43	-12.67	-9.20	-12.51	-9.59		-9.00	-12.01
Truscottite	-6.26	8.35	-17.10	4.25	-16.13			5.02	-15.52
Vaterite	0.61	0.20	0.13	-0.10	-0.07			0.07	-0.09
Xonotlite	0.31	-0.89	-5.40	-4.94	-6.24			-4.24	-6.53

Negative values signify degree of undersaturation, positive values signify degree of saturation.

trace components of the BVG were undergoing constant dissolution throughout the experiments. A S-rich phase was not identified by XRD, so it would appear to be present in only very minor quantities. The most probable candidates for the source of the sulfate would be anhydrite dissolution or pyrite oxidation, both of which are known to occur within some of the fracture mineral assemblages in the BVG.

Mineral saturation-state modelling using *PHREEQC* and the Thermochemie database showed that fluorite was slightly undersaturated in all the experiments (Table 3). It seems unlikely therefore that the accidental addition of fluoride to the YNFG starting solution caused precipitation of this phase. The saturation states of a number of C-S-H phases were also calculated (Table 3), but the very low concentrations of aluminium in solution greatly hindered meaningful calculation of Al-phase saturation indexes (including C-A-S-H phases). Overall, the C-S-H phases that were saturated in the starting solution were relatively Ca-rich, with a high Ca:Si ratio. However, over time the phases that were saturated changed to those with lower Ca:Si ratios. This pattern is consistent with increasing dissolution of minerals in the BVG supplying progressively more silica as the reaction progresses.

Reaction of BVG with evolved cement porefluid (ENFG)

There were many similarities between these experiments and the ones using the YNFP. Again, initial fluid-rock reaction whilst fluids were more alkaline (i.e. dissolution of the starting BVG) was clearly demonstrated through a rapid reduction in pH (from 10.8 to a nearly steady value of 8.7 when recalculated to 70°C). Virtually all of this decrease occurred during the first four months of reaction, and was probably aided by the preferential reaction of the finer grains of the BVG. This pH value is again consistent with buffering by a secondary phase, probably the (K- and Al-rich) C-S-H phase. It is noteworthy that the relative pH decrease between starting solution and experiments is much larger for the ENFG compared to the YNFP. This probably reflects the amount of hydroxyl ions in each solution (in essence, the solution 'buffering capacity'). The ENFG is based upon only limited amounts of Ca(OH)₂, which could be consumed by reaction relatively quickly, and hence pH would decrease relatively rapidly. Conversely, the YNFP contains more hydroxyl ions due to the addition of significant NaOH and KOH as well as Ca(OH)₂,

TABLE 4. Selected mineral saturation state data for ENFG experiments. Data are presented for the starting fluid, together with final compositions of ENFG+BVG experiments and ENFG 'blanks' at 4, 9, 12 and 15 months.

Starting fluid = synthetic 'young near-field porewater' (YNFP)												
Temperature	70 °C		134		276		366		442		474	
Time (days)	0	134	134	471	472	473	1530	1530	474	474	Blank	Blank
Run number	—	470	471	Blank	Crushed BVG	Blank	Crushed BVG	Crushed BVG	Blank	Crushed BVG	Blank	Blank
Experimental charge	—	Crushed BVG	Blank	Blank	Crushed BVG	Blank	Crushed BVG	Crushed BVG	Blank	Crushed BVG	Blank	Blank
Afwillite	3.21	-3.08	-0.1309	-0.1309	-3.45	-1.89	-4.15	-4.15	1.18	-3.68	-0.79	-0.79
Albite-high	-6.16				1.23	-9.42	-0.11	-0.11	-10.15	1.50	-11.30	-11.30
Albite-low	-5.05				2.34	-8.32	1.00	1.00	-9.05	2.61	-10.20	-10.20
Analcime	-2.60				1.96	-5.15	0.59	0.59	-5.78	2.20	-6.36	-6.36
Anhydrite	-3.02				-0.259	-0.1128	-0.2654	-0.2654	-0.1589	-0.2155	-0.104	-0.104
Anorthite					0.22	-6.64	-2.80	-2.80	-7.89	0.68	-7.74	-7.74
Aragonite					1.2018	1.3254			1.23	1.2378	1.3218	1.3218
Barite					-1.7842	-1.3732	-1.3024	-1.3024	0.0172	-1.7165	-1.2582	-1.2582
Bassanite					-0.7956	-0.6494	-0.8023	-0.8023	-0.6957	-0.752	-0.6406	-0.6406
Beidellite-Ca	-10.29				1.43	-14.24	-1.80	-1.80	-16.93	2.09	-17.24	-17.24
Beidellite-K	-9.08				2.67	-13.04	-0.55	-0.55	-15.72	3.34	-16.01	-16.01
Beidellite-Na	-8.90				2.82	-12.87	-0.37	-0.37	-15.52	3.48	-15.86	-15.86
Boehmite	-1.14				-0.33	-1.64	-1.75	-1.75	-2.71	-0.08	-1.92	-1.92
CaCO ₃ .H ₂ O ₍₆₎					0.3778	0.5013			0.4055	0.4138	0.4976	0.4976
Calcite					1.3756	1.4992			1.4037	1.4115	1.4955	1.4955
Celestite					-0.2213	0.2201	-0.2571	-0.2571	0.1578	-0.2201	0.2325	0.2325
C ₂ SH(alpha)	2.02	-3.14			-3.38	-1.15	-3.86	-3.86	0.94	-3.54	-0.18	-0.18
Chabazite	-3.91				5.07	-8.96	2.13	2.13	-10.40	5.58	-11.41	-11.41
Clinoptilolite-Ca	-11.04				1.66	-15.73	0.15	0.15	-16.79	2.00	-18.90	-18.90
Clinoptilolite-K	-13.95				-1.15	-18.69	-2.65	-2.65	-19.71	-0.77	-21.77	-21.77
Clinoptilolite-Na	-12.91				-0.18	-17.64	-1.60	-1.60	-18.60	-0.44	-20.79	-20.79
Cristobalite	-3.33				-0.46	-4.04	-0.42	-0.42	-4.14	-0.44	-4.72	-4.72
CSH _{0.8}	-0.43				-0.86	-2.12	-1.03	-1.03	-1.34	-0.91	-2.14	-2.14
CSH _{1.2}	0.08				-2.01	-2.10	-2.28	-2.28	-0.89	-2.09	-1.79	-1.79
CSH _{1.6}	0.23	-3.31			-3.51	-2.44	-3.89	-3.89	-0.79	-3.64	-1.80	-1.80
Diaspore	-0.50				0.32	-0.99	-1.11	-1.11	-2.06	0.57	-1.27	-1.27
Eitringite					-8.1436	-1.59	-11.7966	-11.7966	-0.6131	-7.7768	0.3389	0.3389
Foshagite	2.68	-4.73			-5.25	-4.36	-6.16	-6.16	-0.29	-5.55	-3.11	-3.11
Gibbsite	-1.52				-0.71	-2.01	-2.13	-2.13	-3.09	-0.45	-2.29	-2.29
Gismondine	2.04				8.51	-5.22	2.46	2.46	-7.72	9.43	-7.41	-7.41
Gypsum					-0.4898	-0.3438	-0.4973	-0.4973	-0.3907	-0.4462	-0.3351	-0.3351
Gyrolite	1.25	1.91			1.60	-3.33	1.20	1.20	-1.44	1.48	-3.72	-3.72
HBeidellite-Ca	-8.45				3.26	-12.41	0.03	0.03	-15.09	3.92	-15.40	-15.40
HBeidellite-K	-8.91				2.84	-12.88	-0.39	-0.39	-15.55	3.51	-15.85	-15.85

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HBeidellite-Na	-7.78	3.94	-11.75	0.74	-14.40	4.60	-14.74
Heulandite-Ca	-11.66	5.37	-18.89	2.32	-20.69	5.97	-23.26
Heulandite-Na	-11.26	5.82	-18.57	2.96	-20.17	6.40	-22.89
Hillebrandite	1.09	-4.31	-2.08	-4.79	0.01	-4.47	-1.11
Illite-Al	-5.50	4.19	-9.69	0.17	-12.56	4.98	-12.20
Jaffeite	5.70	-13.37	-3.09	-14.84	3.26	-13.88	0.49
Jaffeite	9.25	-9.60	-6.05	-12.82	3.18	-11.42	-2.72
Kaolinite	-7.11	0.26	-9.51	-2.51	-11.86	0.81	-11.43
Laumontite	-3.97	5.01	-9.02	2.07	-10.46	5.52	-11.47
Margarite	-2.29	2.57	-6.90	-3.29	-10.30	3.54	-8.55
Merlinoite-Na	-4.96	-0.60	-7.53	-2.05	-8.18	-0.35	-8.70
Microcline	-4.90	2.56	-8.18	1.14	-8.96	2.86	-9.99
Mordenite-Ca	-11.70	1.29	-16.36	-0.12	-17.39	1.61	-19.60
Mordenite-Oregon	-12.84	0.53	-17.48	-0.70	-18.44	0.83	-20.78
Muscovite	-5.12	3.97	-9.37	-0.30	-12.30	4.79	-11.75
Natrolite	-4.32	1.84	-8.73	-0.97	-9.89	2.31	-10.45
Okenite	-0.84	0.78	-3.48	0.60	-2.59	0.73	-4.02
Paragonite	-6.35	2.66	-10.60	-1.53	-13.48	3.44	-13.04
Phillipsite-Ca	-5.11	2.26	-8.34	0.83	-9.16	2.53	-10.24
Phillipsite-K	-5.14	2.32	-8.42	0.90	-9.20	2.63	-10.23
Phillipsite-Na	-4.19	3.20	-7.46	1.85	-8.19	3.47	-9.34
Portlandite	-0.27	-4.40	-1.49	-4.66	-0.40	-4.49	-0.67
Pyrophyllite	-13.12	0.00	-16.94	-2.69	-19.48	0.59	-20.21
Quartz	-2.86	0.02	-3.57	0.05	-3.67	0.04	-4.25
Sanidine	-5.24	2.22	-8.51	0.80	-9.29	2.53	-10.32
Scolecite	-2.19	3.92	-6.53	0.94	-7.88	4.40	-8.30
SiO ₂ (am)	-3.71	-0.84	-4.42	-0.80	-4.52	-0.82	-5.10
SrF ₂ (cr)	-7.18	-5.15	-6.57	0.09	0.54	-5.04	-6.32
SrSiO ₃ (s)	0.66	-0.92	-1.29	-1.16	-0.32	-1.03	-1.15
Stellerite	-30.59	4.63	-44.94	-1.04	-48.42	5.76	-53.90
Stilbite	-23.92	7.19	-38.70	0.03	-42.52	8.51	-46.83
Strontianite		0.1436	0.5624		0.4507	0.1372	0.5623
Tobemmorite-11A	4.08	0.64	-6.31	-0.42	-1.45	0.31	-6.27
Tobemmorite-14A	5.19	1.75	-5.20	0.69	-0.34	1.43	-5.16
Truscottite	-2.51	3.03	-19.61	1.69	-13.14	2.65	-21.99
Vaterite		0.8946	1.0182		0.9228	0.9306	1.0145
Wairakite	-5.95	3.04	-10.99	0.09	-12.44	3.54	-13.44
Zeolite-CaP	-2.94	3.54	-10.19	-2.51	-12.70	4.46	-12.38

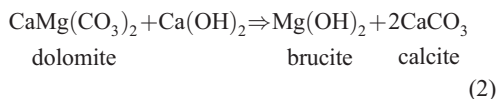
Experiments to react crushed BVG with synthetic cement pore fluids.

Starting fluid = synthetic 'young near-field porewater' (YNFP).

Negative values signify degree of undersaturation, positive values signify degree of saturation.

and so (for a given rate of consumption of hydroxyl ions) would show a slower pH decrease.

The reaction of dolomite with fluids rich in $\text{Ca}(\text{OH})_2$ bears many similarities to the reaction of dolomitic aggregate with high-alkali Portland cement. The latter commonly produces a solids volume increase, expansion, and cracking of concrete (e.g. Galí *et al.*, 2001). Other studies (e.g. Bérubé *et al.*, 1990; Poole and Sotiropoulos, 1980) note that dolomite is reactive in the presence of lime and $\text{Ca}(\text{OH})_2$ -rich solutions to produce brucite and calcite.



Dissolution of the starting BVG was confined mainly to dolomite grains (in the form of pitting), however after 15 months of reaction K-feldspar also showed some signs of corrosion. It is thought that much of the pH reduction was due to the dissolution of dolomite, with most overall reactions occurring early in the experiments when the pH was at its highest. Dissolution of K-feldspar (or possibly muscovite) is thought to have added much of the potassium and silica to solution. The continued linear increase in potassium from 185 mg L^{-1} to 236 mg L^{-1} over 15 months appears to indicate that this continued throughout the experiments. It is noteworthy that such a trend is the opposite of the YNFP case, where (the initially much higher) potassium concentrations fall throughout the experiments. This can be explained by the continued, more extensive incorporation of potassium into C-K-A-S-H gels in the YNFP experiments (higher pH), but only limited amounts of K-rich C-S-H phases in the ENFG experiments (at least over the timescales sampled).

Compared to the YNFP experiments, there were relatively minor amounts of secondary precipitates present in these experiments. Secondary C-S-H ($\pm\text{K}$) products were more abundant in the 4 month experiment compared to the 15 month experiment, and it appears that some precipitates which initially developed dissolved subsequently as the pH of the fluid decreased. This dissolution appeared to add little extra calcium or silica to solution, possibly because of the generally minor abundance of early precipitates, or because they were subsequently re-precipitated as other phases. There was evidence for minor calcite formation on dolomite surfaces after 9 months of reaction. The surface areas of the reaction products initially decreased by 30%, but

after 15 months had risen to almost their starting value. This may be explained by infilling or blocking of porosity within the grains by the relatively small quantities of secondary products. Subsequent removal of these by re-dissolution would have exposed much of the original porosity. Re-dissolution might help explain the increase in calcium concentration at 15 months (though conclusions are tentative as only a single datum is involved). Whilst dissolution of C-S-H could have increased calcium concentrations, it might also be expected to increase silica in solution, but there is no evidence for this. An explanation for this could be an evolution of C-S-H phase composition (C-S-H phases can have a wide range of Ca:Si ratios), so there is a possibility that a more Si-rich C-S-H phase precipitated later in the experiments at the expense of a more Ca-rich one. The potential for this was explored through geochemical modelling of C-S-H saturation states using *PHREEQC* and the GEMS database, as per the YNFP cases (Table 4). A similar pattern emerged, with saturation of relatively Ca-rich C-S-H phases predicted in the starting solution. However, these changed over time, and became dominated by more Si-rich C-S-H phases through progressive reaction with the minerals in the BVG.

There is only tentative evidence for hydroxyapophyllite formation after 4 months of reaction, but not at longer timescales. If it did form, this again suggests rapid formation early in the experiment, followed by its re-dissolution as the pH of the solution evolved towards lower values.

Celestite was observed as per the YNFP experiments and was the only secondary sulfate phase identified. However, previous studies (Rochelle *et al.*, 1994*b*, 1998*a,c*) using more evolved cement porewaters noted initial precipitation of the sulfate mineral ettringite ($\text{Ca}_6\text{Al}_2(\text{SO}_4)_3(\text{OH})_{12}\cdot 26\text{H}_2\text{O}$), which can be followed by its dissolution as it becomes destabilized as a result of pH decrease (Rochelle *et al.*, 1994*b*). Unfortunately, there was no mineralogical evidence for this in the present sets of experiments (such as impressions of ettringite needles in mats of C-S-H as reported by Rochelle *et al.*, 1994*b*). A reason for this may be that previous studies reacting powdered BVG with ENFG (Rochelle *et al.*, 1994*b*, 1998*a,c,e*) had short durations, not exceeding 4 months long. It is entirely conceivable that ettringite precipitated initially in the current set of long-term experiments, but subsequently dissolved (though this remains conjecture). If ettringite did form and then dissolve, initially precipitated sulfate would have been returned to solution, possibly with some proportion of it

precipitating as celestite. Consequently, the observed sulfate concentrations are not inconsistent with previous experiments under the NSARP. Running repeat experiments over much shorter timescales would be one way to check this possibility.

Discussion and implications for the near repository environment

In the performance assessment for a cement-based repository for radioactive waste, it will be necessary to consider the alkaline, chemically-disturbed zone. In order to understand the temporal and spatial evolution of this zone, it will also be necessary to consider how fluid-conductive features such as fractures evolve when they are filled with a high-pH fluid that is out of equilibrium with the host rock. Previous experimental and modelling studies (Rochelle *et al.*, 1994a; 1998a,b,c,e; Savage *et al.*, 1992, Savage and Rochelle, 1993) undertaken as part of the NSARP have highlighted the importance of C-S-H phases and ettringite in reducing porosity, in coating fracture surfaces and increasing mineral surface areas available for sorption.

In the set of experiments presented here, the BVG sample used was taken from a potential flowing feature, and should thus provide information on reactions most likely to occur when hyperalkaline fluids migrate along currently open fractures. Use of a sample of fracture fill is more realistic than using samples of bulk rock, as the latter may not actually contact alkaline fluids to any significant extent. The sample used in this investigation contained much fine-grained gouge material and dolomite, both of which appeared to contribute to the relatively fast reaction of the rock, and relatively rapid reduction in pH of the alkaline solutions. This reaction appeared to be most noticeable when the pH was highest. One feature of such fast reaction was the rapid formation of secondary phases, which included: significant amounts of a secondary C-(K,A)-S-H gel, some hydroxyapophyllite, and some brucite in the YNFP experiments; and more limited amounts of C-(K)-S-H gel, some celestite, minor calcite and potentially trace hydroxyapophyllite in the ENFG experiments.

The presence of significant dolomite in the starting solid is particularly noteworthy. Whilst this phase might be relatively more uniformly distributed within sedimentary host rocks (e.g. Adler *et al.*, 1999), it could be restricted only to fracture fills within fractured basement rocks such as the BVG (Milodowski *et al.*, 1998, 2002). In the latter therefore, though it may make up a tiny fraction of

the total rock mass it could be a highly significant proportion of fracture assemblages. As alkaline fluids migrating through fractured rock will be focused along these fractures, the reactivity of dolomite will have an impact far greater than would be suggested by its overall abundance in the rock mass. The observations from this study have wider application than to just the BVG in the UK. A dolomite-bearing host-rock mineralogy is also relevant to a number of potential repository and underground research laboratory (URL) sites being considered elsewhere, including: the Opalinus Clay (Jurassic) in Switzerland (Wenk *et al.*, 2008); the Callovo-Oxfordian (Jurassic) shales in the Meuse/Haute-Marne site near Bure in France (Wenk *et al.*, 2008); the Boom Clay (Tertiary) at the Mol site in Belgium (Honty and De Craen, 2009); Tertiary mudrocks at the Horonobe URL site, Hokkaido, Japan (Kunimaru *et al.*, 2010); and fracture mineralization in the Archaen basement rocks of the Olkiluoto site in Finland (Kuva *et al.*, 2012).

Dolomite may also play two other important roles as well as its ability to reduce solution pH. Firstly, its dissolution will increase $p\text{CO}_2$ locally, and this may suppress the formation of secondary zeolites (Adler *et al.*, 1999; Thompson, 1971). This may therefore limit the benefit that would be gained through ion-exchange and retardation of migrating radionuclides into zeolites. Secondly, its dissolution followed by calcite precipitation may help seal the wall of fractures, reducing diffusion and uptake of radionuclides into the rock-matrix. This has been shown via diffusion-reaction calculations undertaken by Steefel and Lichtner (1994, 1998). Although their calculations were undertaken for fractures in a sedimentary rock, similar processes may occur for less permeable basement rocks. They predicted that carbonate production by dolomite dissolution, coupled with interdiffusion of calcium from the infiltrating high-pH solution, has the potential to reduce significantly matrix porosity of rock immediately adjacent to the fracture through calcite precipitation.

Insufficient information exists from these experiments to quantify how the solids volume changed during the experiments. However, given the apparent abundance of the (K/Al-rich) C-S-H gel in some of the experiments, then it (qualitatively) appears to have a potentially important role in modifying fracture porosity. This phase also appears to have appreciable surface area, which would be of benefit when considering the retardation of radionuclides by sorption. Conversely however, a coating of precipitate may also isolate the rock matrix to some extent, and so it may have an impact on the accessibility of

the rock matrix to radionuclide diffusion. Further investigations are needed to determine the relative contributions of each of these processes, and hence ascertain whether radionuclide migration would be enhanced or retarded.

Although phases such as hydroxyapophyllite, and possibly the very K-rich C-S-H gel, have not previously been observed within experiments conducted as part of the NSARP, they are not entirely unexpected. Potassium-rich C-S-H can occur within cements (Hong and Glasser, 2002; L'Hôpital, 2014), and hydroxyapophyllite is commonly associated with zeolites as an alteration product of basalts by moderately alkaline fluids (Deer *et al.*, 1992). In terms of contributing to improvements in predictive geochemical modelling of alkaline fluid-fracture assemblages, it is important that thermodynamic data for these phases are included within model databases. No zeolites were observed in the experiments. There are several possible reasons for their absence, and it is not yet possible to identify which of these is the controlling factor. One or more of the following could have been key: rapid removal of aluminium and silica from solution by the formation of a range of C-S-H (C-A-S-H, C-K-A-S-H) phases, slow zeolite precipitation kinetics at the temperatures used, relatively high $p\text{CO}_2$ in the experiments, and a relatively high fluid:rock ratio. The data obtained from this study suggest therefore, that zeolites would not be expected in water-conducting, carbonate-lined fractures, and this should be considered when undertaking predictive geochemical modelling of alkaline fluid-fracture assemblages.

Conclusions

Re-investigation of a series of old experiments is providing an unique opportunity to study longer-term fluid-rock reaction processes within alkaline plumes, and this study shows that it is possible to follow alkali fluid-rock reactions within experimental systems for months to years. Some of the geochemical and mineralogical changes are broadly in line with previous experimental, modelling and analogue studies, though others are new. Dissolution of dolomite, and to a lesser extent silicates (probably K-feldspar, but possibly also mica) occurred relatively rapidly at 70°C and were apparent within just a few months. Dolomite dissolution may have been a key factor in reducing pH values, and may be a key mineral in controlling the extent of alkaline disturbed zones. Dissolution was followed by precipitation of

brucite close to dolomite grains, at least two generations of C-S-H phases (which may have contained variable amounts of K, Al and Mg), overgrowths of calcite, small crystals of hydroxyapophyllite and elongate crystals of celestite. Though hydroxyapophyllite was observed (a phase commonly associated with zeolites), there was no evidence for the formation of zeolites in the experiments. Precipitation was more extensive in the experiments involving Na-K-Ca (younger) cement porefluids compared to more evolved (Ca-rich) cement porefluids. With continued fluid-rock reaction however, some of these initially-formed phases had begun to redissolve as continued fluid-rock reaction reduced the pH of the coexisting fluid.

This investigation highlights the importance of running longer-term experiments to study processes that cannot be examined during (the more common) short-term laboratory studies. The relatively simple experimental equipment aids long-duration tests, and a number of identical experiments continue to run. These will be sampled at increasing timescales, and eventually will provide quantitative data on alkaline fluid-rock reaction over several decades.

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