

Experimental constraints on Li isotope fractionation during clay formation

Ruth S. Hindshaw^{a,*}, Rebecca Tosca^b, Thomas L. Goût^a, Ian Farnan^a, Nicholas J. Tosca^b, Edward T. Tipper^a

^a*Department of Earth Sciences, University of Cambridge, Downing Street, Cambridge, UK, CB2 3EQ*

^b*Department of Earth Sciences, University of Oxford, South Parks Road, Oxford, UK, OX1 3AN*

Abstract

Knowledge of the lithium (Li) isotope fractionation factor during clay mineral formation is a key parameter for Earth system models. This study refines our understanding of isotope fractionation during clay formation with essential implications for the interpretation of field data and the global geochemical cycle of Li. We synthesised Mg-rich layer silicates (stevensite and saponite) at temperatures relevant for Earth surface processes. The resultant solids were characterised by X-ray diffraction (XRD) and Fourier-transform infrared spectroscopy (FT-IR) to confirm the mineralogy and crystallinity of the product.

Bulk solid samples were treated with ammonium chloride to remove exchangeable Li in order to distinguish the Li isotopic fractionation between these sites and structural (octahedral) sites. Bulk solids, residual solids and exchangeable solutions were all enriched in ⁶Li compared to the initial solution. On average, the exchangeable solutions had $\delta^7\text{Li}$ values 7‰ lower than the initial solution. The average difference between the residue and initial solution $\delta^7\text{Li}$ values ($\Delta^7\text{Li}_{\text{residue-solution}}$) for the synthesised layer silicates was $-16.6 \pm 1.7\text{‰}$ at 20°C, in agreement with modelling studies, extrapolations from high temperature experimental data and field observations. Three bonding environments were identified from ⁷Li-NMR spectra which were present in both bulk and residual solid ⁷Li-NMR spectra, implying that some exchangeable Li remains after treatment with ammonium chloride. The ⁷Li-NMR peaks were assigned to octahedral, outer-sphere (interlayer and adsorbed) and pseudo-hexagonal (ditrigrical cavity) Li. By combining the ⁷Li-NMR data with mass balance constraints we calculated a fractionation factor, based on a Monte Carlo minimum misfit method, for each bonding environment. The calculated values are $-21.5 \pm 1.1\text{‰}$, $-0.2 \pm 1.9\text{‰}$ and $15.0 \pm 12.3\text{‰}$ for the octahedral, outer-sphere and pseudo-hexagonal sites respectively (errors 1 σ). The bulk fractionation factor ($\Delta^7\text{Li}_{\text{bulk-solution}}$) is dependent on the chemistry of the initial solution. The higher the Na concentration in the initial solution the lower the bulk $\delta^7\text{Li}$ value. We suggest this is due to Na outcompeting Li for interlayer sites and as interlayer Li has a high $\delta^7\text{Li}$ value relative to octahedral Li, increased Na serves to lower the bulk $\delta^7\text{Li}$ value. Three experiments conducted at higher pH exhibited lower $\delta^7\text{Li}$ values in the residual solid. This could either be a kinetic effect, resulting from the higher reaction rate at high pH, or an equilibrium effect resulting from reduced Li incorporation in the residual solid and/or a change in Li speciation in solution.

This study highlights the power of ⁷Li-NMR in experimental studies of clay synthesis to target site specific Li isotope fractionation factors which can then be used to provide much needed constraints on field processes.

Keywords: Li isotopes, Clay minerals, Clay synthesis, Biogeochemical cycles

1. Introduction

Lithium (Li) is a trace element which is predominantly found in silicate minerals (e.g. Tardy et al., 1972; Huh et al., 1998). Li has two stable isotopes, ^6Li and ^7Li , and variations in their relative abundance are expressed in standard delta notation as $\delta^7\text{Li}$. The natural range of $\delta^7\text{Li}$ in geological and environmental materials is approximately 60‰ (Tomascak, 2004), with much of this range thought to be generated by fractionation associated with the preferential incorporation of ^6Li into clay minerals and adsorption/exchange onto particle surfaces (e.g. Zhang et al., 1998; Vigier et al., 2008). No appreciable amount of Li is incorporated into the biosphere and fractionation of Li by plants is thought to be negligible (Lemarchand et al., 2010; Clergue et al., 2015). There is no appreciable fractionation of Li isotopes during mineral dissolution at low temperatures $<50^\circ\text{C}$ (Pistiner and Henderson, 2003; Wimpenny et al., 2010a; Verney-Carron et al., 2011). For these reasons, Li isotopes are one of the most promising tracers to emerge in recent decades for the intensity of silicate weathering; defined as the ratio of the total erosion mass flux to the total weathering mass flux (Bouchez et al., 2013). Records of $\delta^7\text{Li}$ in marine carbonates, interpreted to be representative of seawater $\delta^7\text{Li}$ over the Phanerozoic, potentially provide a unique record of changing silicate weathering intensity over time, with implications for the importance of weathering-related feedbacks in the long-term carbon cycle (Froelich and Misra, 2014; Vigier and Godd ris, 2015; Pogge von Strandmann et al., 2017a). The seawater Li isotope record (Hall et al., 2005; Hathorne and James, 2006; Misra and Froelich, 2012; Pogge von Strandmann et al., 2013; Lechler et al., 2015; Pogge von Strandmann et al., 2017a) has been interpreted to reflect 1) variations in the mean riverine $\delta^7\text{Li}$ flux and thus changes in continental silicate weathering processes through time (Hathorne and James, 2006; Misra and Froelich, 2012; Wanner et al., 2014; Vigier and Godd ris, 2015; Pogge von Strandmann et al., 2017a), 2) a change in the balance between low and high temperature sinks of Li from the oceans (Li and West, 2014) and 3) changes in the balance between continental and submarine basalt weathering (for Jurassic ocean anoxic events, Lechler et al., 2015; Pogge von Strandmann et al., 2017a). The mass balance of Li in seawater is controlled by inputs from continental weathering and marine hydrothermal systems, and outputs due to removal by clays (alteration of oceanic crust and reverse weathering). Critically, both the inputs and outputs of Li are affected by Li isotope fractionation associated with the formation of secondary phases (clays) but the magnitude of fractionation lacks experimental constraints, particularly at temperatures relevant to weathering and reverse weathering ($<50^\circ\text{C}$). To date, different studies have assumed different fractionation factors ranging from 13 to 30‰ (Hathorne and James, 2006; Misra and Froelich, 2012; Li and West, 2014; Wanner et al., 2014), which have been compiled/extrapolated from field studies and a limited number of experimental studies (Vigier et al., 2008).

Li isotope fractionation attendant to clay mineral formation likely occurs for a number of reasons. Firstly, Li may be incorporated into octahedral sites within the clay mineral structure. If isotopic equilibrium is attained, the

*corresponding author

Email address: ruth.hindshaw@gmail.com (Ruth S. Hindshaw)

32 magnitude of fractionation should depend on the vibrational frequency or bond strength between Li in octahedral
33 sites and Li in solution (Bigeleisen, 1965). If the isotopic fractionation is kinetic then there may be an additional rate
34 dependence (Schauble, 2004). Secondly, Li may be adsorbed to clay mineral surfaces, held on negatively charged
35 layer surfaces, or be located in pseudo-hexagonal sites within the clay tetrahedral layer. These sites may fractionate
36 Li isotopes if there are differences in bond strength (Pistiner and Henderson, 2003). The bulk fractionation factor in a
37 clay therefore reflects the sum total of the isotope fractionation associated with each bonding environment.

38 Only one study to date has directly synthesised a clay mineral at low (Earth surface) temperature (hectorite, a Li-
39 rich smectite, Vigier et al., 2008). Two studies have investigated incorporation of Li into existing mineral structures.
40 The first investigated the uptake of Li during illitization (Williams and Hervig, 2005) and the second, the intercalation
41 of Li into gibbsite (Wimpenny et al., 2015). Additionally, Li isotope fractionation has been observed during the
42 formation of a solid inferred to be chrysotile as a by-product of forsterite dissolution at high pH (Wimpenny et al.,
43 2010a) and during the formation of a mixture of smectite phases (nontronite, beidellite) resulting from basalt-seawater
44 interaction experiments (Millot et al., 2010b). High-temperature and pressure experiments have investigated Li isotope
45 fractionation during mineral (mica, staurolite, spodumene, serpentine-group) formation (Wunder et al., 2006, 2007,
46 2010) and Li assimilation (Lynton et al., 2005).

47 Adsorption of Li onto several types of mineral (zeolite, vermiculite, kaolinite, smectite, ferrihydrite, gibbsite)
48 has been reported (Taylor and Urey, 1938; Zhang et al., 1998; Pistiner and Henderson, 2003) together with indirect
49 measurements of adsorption onto vernadite and amorphous Fe oxides from Li isotope measurements of bulk Fe-
50 Mn crusts (Chan and Hein, 2007). The resultant fractionation is interpreted to depend on the strength of the Li-
51 surface bond i.e. whether an inner or an outer-sphere complex is formed, with the more strongly-bound inner-sphere
52 complexes resulting in greater Li isotope fractionation relative to the starting solution (Pistiner and Henderson, 2003).
53 In all studies of adsorbed and structurally incorporated Li, ^6Li was preferentially incorporated, consistent with results
54 from density functional theory models which predict that ^7Li is preferentially retained in the aquo-complex (Yamaji
55 et al., 2001; Bogatko et al., 2013).

56 Some studies have explicitly tried to focus on octahedral (structural) Li by treating the solids to remove exchange-
57 able Li and then measuring the Li isotopic composition of the residue (Williams and Hervig, 2005; Vigier et al.,
58 2008), whereas others refer to a bulk phase including octahedral and exchangeable Li (Wimpenny et al., 2015). The
59 term 'exchangeable' is used to encompass all the Li held in loosely bound sites, for example, in the interlayer or
60 adsorbed to negatively charged layer surfaces. It has been suggested that Li held in outer-sphere bonding sites, which
61 is thought to be the case for Li adsorbed to smectites and interlayer Li, is not isotopically fractionated relative to the
62 solution (Pistiner and Henderson, 2003; Chan and Hein, 2007). This implies that bulk phases including a fraction of
63 exchangeable Li will have $\delta^7\text{Li}$ values closer to the initial solution than those where all the exchangeable Li has been
64 removed (Wimpenny et al., 2015).

65 All experimental data considered to be free of non-equilibrium effects were found to be in good agreement with
66 theoretical modelling predictions (Dupuis et al., 2017). In agreement with equilibrium fractionation, increasing the

67 temperature decreased the fractionation factor during hectorite formation (Vigier et al., 2008). In contrast, during
68 chrysotile precipitation, the higher temperature experiments led to a greater difference in $\delta^7\text{Li}$ between the solid
69 formed and the final solution composition (Wimpenny et al., 2010a). Vigier et al. (2008) found that the measured
70 fractionation factor did not depend on the amount of Li incorporated into hectorite or the solution composition whereas
71 Wimpenny et al. (2015) found that fractionation of Li during intercalation of Li into gibbsite was dependent on the
72 availability of free octahedral sites and the identity of anions in solution. Neither the structure nor composition of the
73 clay formed influenced the magnitude of Li isotope fractionation into phyllosilicates (Dupuis et al., 2017). Further
74 studies have investigated the effect of physical controls such as crystal size on the magnitude of fractionation (Williams
75 and Hervig, 2005; Wunder et al., 2010).

76 Collectively, these studies have found that the extent of Li isotope fractionation depends on 1) coordination num-
77 ber (bond strength), 2) temperature and 3) chemistry of solution, including pH. Similar controls on stable isotope
78 fractionation have been inferred for other elements (e.g. Tang et al., 2008; Nielsen et al., 2012; Li et al., 2014).
79 Most previous experimental studies have focussed on high temperature synthesis methods as these typically involve
80 higher crystallisation rates which, in turn, generate larger amounts of crystalline material over shorter experimental
81 timescales (Tosca and Masterson, 2014). However, whether these data are reflective of secondary mineral formation
82 at Earth surface temperatures is less clear. In this study, we apply a low temperature synthesis method to synthesise
83 smectite-group minerals directly from solution and investigate the controls on the resultant Li isotope fractionation.
84 Further, we use a cation exchange procedure to distinguish exchangeable from octahedral Li coupled with ^7Li -NMR
85 to quantify the fraction of Li in each bonding environment, permitting us to estimate site specific fractionation factors.

86 *1.1. Mg-rich layer silicate minerals*

87 We precipitated two layer silicate structures directly from solution: stevensite and saponite. These belong to the
88 Mg-smectite group of minerals $((\text{Na,Ca})_{0.3}(\text{Al,Mg})_2\text{Si}_4\text{O}_{10}(\text{OH})_2 \cdot n\text{H}_2\text{O})$ comprised of layers of octahedra (Mg or Al
89 bonded to O or OH) sandwiched between two layers of Si tetrahedra (Si bonded to O or OH, Fig. 1), thus this structure
90 is also referred to as ‘TOT’ or ‘2:1 layer’. Smectites are found in numerous low temperature environments (marine
91 and continental) and as such they constitute an important reservoir of Mg; a major element which can readily transfer
92 between the crust, mantle and hydrosphere. Li, on the other hand, is a trace element but due the similar ionic radii
93 of Li and Mg, Li can readily substitute for Mg in minerals (Huh et al., 1998; Decarreau et al., 2012) and therefore Li
94 may be able to help constrain the Mg biogeochemical cycle.

95 Stevensite is a trioctahedral layer silicate mineral with a Mg-rich octahedral sheet, a pure Si-rich tetrahedral sheet
96 and octahedral vacancies (Guggenheim, 2016). In stevensite, the spacing between each TOT layer (d_{001} , Fig. 1) is
97 $\sim 14\text{-}15 \text{ \AA}$ depending on whether the sample is in the air-dried, hydrated or ethylene glycol-solvated state. Stevensite
98 can only accommodate trace aluminium in the lattice (Guggenheim, 2016). The octahedral vacancies ($\sim 0.08\text{-}0.10$
99 cation sites per unit) are thought to give rise to an overall layer charge which imparts swelling characteristics and
100 interlayer ion accommodation similar to other members of the smectite group (Brindley et al., 1977; Steudel et al.,

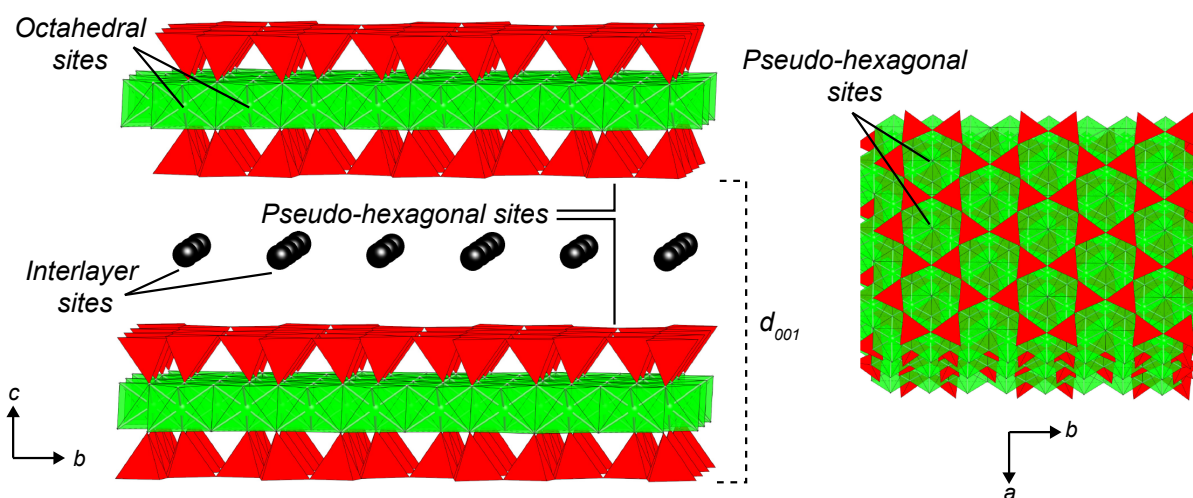


Figure 1: Schematic figure of the overall structure of a 2:1 layer silicate mineral shown from two different perspectives. Each layer consists of an octahedral sheet (green squares) sandwiched between two tetrahedral sheets (red triangles). In between, depending on the basal spacing (d_{001}), which is also related to the overall layer charge and therefore chemical composition of the octahedral sheet, there can be interlayer cations (depicted here without their hydration spheres). Three main crystallographic sites for Li are highlighted: octahedral, interlayer and pseudo-hexagonal sites. Li may also be adsorbed onto edge sites but this process is not depicted in this figure.

101 2017). Li can fill these octahedral vacancies and also substitute for Mg. A Mg-rich trioctahedral layer silicate closely
 102 related to stevensite is kerolite ($\text{Mg}_3\text{Si}_4\text{O}_{10}(\text{OH})_2 \cdot n\text{H}_2\text{O}$; $n = 0.8-1.2$). Kerolite is non-expandable and is characterised
 103 by H_2O positioned either on edges associated with broken bonds or within the interlayer (Guggenheim, 2016). Kerolite
 104 exhibits a basal spacing commonly measured by powder X-ray diffraction (uncorrected for polarisation effects) of
 105 $\sim 10.1 \text{ \AA}$. Naturally occurring stevensite has been found to contain random mixed layers of kerolite, and the converse
 106 has also been observed (Eberl et al., 1982; Martin de Vidales et al., 1991; Dekov et al., 2008). Thus, as a result
 107 of their closely related structures and thermodynamic stability, kerolite and stevensite often occur together as direct
 108 precipitates from the weathering of ultramafic lithologies and from alkaline saline lakes (Stoessel and Hay, 1978;
 109 Jones, 1986; Tosca, 2016). Saponite is structurally similar to stevensite except Al^{3+} substitution for Si^{4+} in tetrahedral
 110 sites is more extensive. A trioctahedral smectite where Li and Mg occur as a major elements (wt%) is called hectorite.
 111 Stevensite, kerolite and saponite may all contain minor or trace amounts of Li in their structure and there is no formal
 112 division in terms of Li concentration between these structures and hectorite.

113 The Mg-rich trioctahedral layer silicate minerals offer an ideal starting point for investigating low temperature clay
 114 mineral precipitation because they exhibit relatively rapid reaction kinetics yet contain all of the essential structural
 115 characteristics of smectite and other TOT clay minerals.

116 2. Methods

117 2.1. Mg-rich layer silicate synthesis

118 There are established methods for both the high and low temperature synthesis of smectite clay minerals (e.g.
119 Harder, 1972; Vogels et al., 2005). High temperature methods start with a seed or a gel which is then heated at tem-
120 peratures $>90^{\circ}\text{C}$ (e.g. Güven and Carney, 1979; Vogels et al., 2005; Vigier et al., 2008). This method results in highly
121 crystalline materials which are arguably most analogous to clays formed in hydrothermal settings. In contrast, low
122 temperature methods rely on direct precipitation from solution (e.g. Harder, 1972; Tosca and Masterson, 2014). Al-
123 though the quantity of solid formed is lower and the products are less crystalline than the products obtained from high
124 temperature experiments, the experimental conditions of low temperature synthesis are arguably most representative
125 of the majority of low temperature weathering environments (e.g. soils) and processes (e.g. diagenesis). These exper-
126 iments may also be representative of what is occurring in early stage weathering environments where clay phases are
127 often not detected but their presence is inferred from isotope fractionation (Tipper et al., 2012). Low temperature ex-
128 periments are essential for directly determining fractionation factors at similar temperatures to the natural weathering
129 environment, avoiding uncertainty associated with extrapolation from high temperature experiments.

130 We synthesised two different Mg-rich layer silicate minerals (stevensite and saponite) at room temperature (20°C)
131 using two different methods. The first method used 1 L PPE bottles with an electrolyte buffer and the second method
132 used a reaction vessel coupled to an autotitrator. These two setups are hereafter referred to as ‘bottle’ and ‘reaction
133 vessel’ experiments.

134 For stevensite bottle experiments, the solutions contained Mg (from MgCl_2), Li (from LiCl), Na (from NaCl)
135 and Si (from Na_2SiO_3) and the pH was adjusted to the required value using HCl and buffered for the duration of
136 the experiment using TRIS (tris(hydroxymethyl)aminomethane). All experiments were stirred magnetically and the
137 duration of the experiments varied from 9 days (‘010217’ series) to 28 days (‘240417’ series). For the saponite bottle
138 experiment, the initial solution additionally contained Al (from $\text{Al}(\text{NO}_3)_3 \cdot 9\text{H}_2\text{O}$) and was prepared similarly to those
139 for stevensite except that the source of Si was TEOS (tetraethyl orthosilicate), which was allowed to dissociate for
140 24 h before Al and Mg were added to the solution. The solution also contained Ca (from CaCl_2) and the bottle was
141 placed on a shaker table for 26 days. The stevensite reaction vessel experiment was conducted in a 1 L polycarbonate
142 reaction vessel and used the same reagents as the stevensite bottle experiments but as the experiment was connected
143 to an autotitrator which delivered 0.1 mol/kg NaOH , no buffer was needed. This experiment lasted for 14 days. The
144 target initial concentrations of Si, Al, Mg, Na, Ca and Li used in the different experiments are listed in Table 1. A
145 solid precipitate was recovered from all experiments apart from two experiments run at $\text{pH} < 9.3$ and low [Mg]. The
146 pH of the experiments varied from 9.0 to 10.4 (Table 1). In general, the higher the pH, the greater the mass of the
147 recovered precipitate for experiments of the same duration.

148 The elevated pH associated with the onset of nucleation and growth of Mg-rich layer silicate phases stems from
149 the hydration characteristics of the Mg^{2+} ion. Nevertheless, the nucleation and growth processes involved in direct

Table 1: Summary of the target initial solution conditions for the stevensite and saponite synthesis experiments.

Experiment(s)	Type	Clay	pH	Temp °C	[Si] mmol/kg	[Al] mmol/kg	[Na] mmol/kg	[Mg] mmol/kg	[Li] mmol/kg	[Ca] mmol/kg
250117	Rxn vessel	stevensite	10.3	20	2.6	0	5	2	2	0
010217-E	bottle	stevensite	10.0	20	1.7	0	100	2	2	0
010217-F	bottle	stevensite	10.3	20	2.6	0	100	2	2	0
240417A-D, I, J	bottle	stevensite	9.0-10.2	20	2.2	0	500	2	2	0
240417E-H,K,L	bottle	stevensite	9.0-10.4	20	2.2	0	500	20	2	0
20140624F	bottle	saponite	9.5	20	3.7	0.07	1	33	1.2	0.5

precipitation of these phases from solution is thought to closely mirror other clay mineral systems containing, for example, Fe and Al (Tutolo and Tosca, 2018; Tosca, 2016). Alkaline pH is common in environments such as marine pore-waters, subglacial meltwaters and rivers draining basalt (Dessert et al., 2001; Gíslason et al., 1996; Reimers et al., 1996; Pokrovsky et al., 2005; Hindshaw et al., 2014). The canonical natural analogue for reverse weathering is African alkaline soda lakes (pH>9) where smectite-group minerals precipitate (Von Damm and Edmond, 1984). Although the pH of river water and bulk soil pore waters is generally close to neutral (Stumm and Morgan, 1996), the pH of soil pore waters at the nano- to micro-scale, where coupled mineral dissolution and precipitation reactions occur, is higher than that measured in bulk soil solutions (Hochella and Banfield, 1995; Carrado et al., 2006). Therefore the experimental conditions are a good analogue for many, but not all, natural settings.

Aliquots (2 mL) of the initial and final solution were collected from all experiments and filtered at 0.22 μm . Solids from all experiments were recovered by filtration through 0.1 μm nitrocellulose filters. The solid was air-dried and saved for analysis.

Part of the recovered clay sample was saturated with 1 M NH_4Cl in order to extract the Li bound in exchangeable sites (Vigier et al., 2008). 10 mg of the clay was placed in a 2 mL centrifuge tube with 1 mL 1 M NH_4Cl . The mixture was shaken overnight, centrifuged and the supernatant pipetted off. This procedure was performed in triplicate for experiment 20140624F and once for all other experiments. The supernatant was dried down and digested in concentrated HNO_3 . The residue and an untreated bulk aliquot were digested in a mixture of concentrated HF and HNO_3 and then repeatedly dried down and redissolved in 6 M HCl. Once digested, the samples were dissolved in 2% HNO_3 in preparation for concentration analysis.

2.2. Concentration and Li isotope measurements

All Li and Mg concentrations were measured by inductively-coupled plasma optical emission spectrometry (ICP-OES, Agilent Technologies 5100, University of Cambridge) in cool plasma mode to ensure a low Li background and using matrix matched standards. The RSD of water standard TM25.4 (Environment Canada) was 3.2%. Concentrations of the residual and exchangeable fractions are given as mg Li per kg of bulk sample. Li isotopes were measured using a procedure described in Hindshaw et al. (2018). Briefly, a sample containing 15-20 ng Li was dried down and passed through a column containing 3 mL Bio-Rad AG 50W-X12 200-400 mesh resin, eluting with 0.2 M HCl. Lithium isotope ratios were measured on a Neptune Plus multi-collector ICP-MS using an APEX IR sample intro-

177 duction system, a 50 $\mu\text{L}/\text{min}$ PFA nebuliser and nickel H cones. $^7\text{Li}/^6\text{Li}$ ratios were normalised to L-SVEC (NIST
178 RM 8545) using standard-sample bracketing. Analyses were performed on 5 ppb solutions using a 10^{13} Ω resistor
179 on the pre-amplifier for ^6Li (Bohlin et al., 2018). The typical beam-size for samples was around 1 V on ^7Li and
180 the background was typically between 5 and 20 mV. Accuracy and precision were monitored using Li6-N and Li7-N
181 solutions which gave values of $-8.1\pm 0.9\%$ (2SD, n=59) and $+30.2\pm 0.9\%$ (2SD, n=50) respectively. These values are
182 in agreement with previously published values for these standards (Li7-N = $+30.2\%$ and Li6-N = -8.2% , Carignan
183 et al., 2007; Millot et al., 2010c). Long-term reproducibility of L-SVEC was $+0.1\pm 0.5\%$ (2SD, n=72). To ensure
184 there was no fractionation induced during chemical separation, either seawater or a rock standard was processed with
185 every batch of 5 samples. Long-term reproducibility for seawater (OSIL IAPSO batch P157) is $+30.8\pm 1.1\%$ (2SD,
186 n=45) in agreement with the compiled values of $+31.1\%$ (Carignan et al., 2004) and $+30.8\%$ (Rosner et al., 2007).
187 We obtained a value of $+4.9\pm 1.9\%$ (2SD, n = 10) for USGS shale rock standard SGR-1b (Hindshaw et al., 2018),
188 in agreement with previously published values (Phan et al., 2016; Pogge von Strandmann et al., 2017b; Bohlin et al.,
189 2018). The long-term external reproducibility of seawater is applied to the samples measured in this study (1.1%), as
190 this value is greater than the 2SD of individual sample measurements.

Table 2: Li concentration and isotope measurements of the analysed solids and solutions. Concentrations of the residual and exchangeable fractions are given as mg Li per kg of bulk sample.

Experiment	pH	solid g	[Li] _i		δ ⁷ Li _i ¹		[Li] _f		δ ⁷ Li _f ¹		[Li] _e		δ ⁷ Li _e ¹		[Mg] _r		[Li] _r		δ ⁷ Li _r ¹		f _e		f _r		[Li] _b		δ ⁷ Li _b ¹		δ ⁷ Li _b [*]		Δ ⁷ Li _{r, residue-int. s.in²}	
			mg/L	initial solution	‰	‰	mg/L	final solution	‰	‰	mg/kg	exchangeable Li	‰	‰	g/kg	residual Mg and Li	‰	‰	mg/kg	bulk solid	‰	‰	mass fraction	mass fraction	mg/kg	bulk solid	‰	‰	‰	‰	‰	‰
<i>Reaction vessel - stevensite</i>																																
250117	10.3	0.2416	10.4	0.9	11.2	1.4	252.6	-1.4	246.6	56.7	-11.6	0.82	0.18	183.6	309.3	-3.5	-3.9	-12.5														
<i>Bottle - stevensite - low Na series</i>																																
010217E	10.0	0.1000	13.4	1.3	13.1	1.6	47.5	-2.3	120.7	53.5	-14.6	0.47	0.53	101.1	101.0	-8.6	-10.0	-16.0														
010217F	10.3	0.1917	13.9	1.0	14.3	1.1	51.2	-2.3	107.3	33.2	-14.1	0.61	0.39	91.9	84.3	-7.8	-8.6	-15.1														
<i>Bottle - stevensite - low Mg series</i>																																
240417B	9.5	0.0043	15.3	2.1	15.4	1.8	n.m.	n.m.	n.m.	n.m.	n.m.	n.m.	n.m.	97.2	n.m.	-12.7	n.m.	n.m.														
240417J	9.7	0.0393	15.1	2.3	15.1	1.6	19.4	-7.9	87.1	46.8	-14.1	0.29	0.71	65.6	66.2	-12.7	-12.3	-16.3														
240417C	9.9	0.0764	15.1	1.9	15.0	1.3	16.3	-6.9	101.9	48.7	-15.3	0.25	0.75	65.3	65.0	-13.1	-13.2	-17.1														
240417D	10.2	0.0890	16.0	2.1	15.9	1.9	16.2	-5.9	113.5	39.2	-15.8	0.29	0.71	53.8	55.4	-12.6	-12.9	-17.9														
<i>Bottle - stevensite - high Mg series</i>																																
240417E	9.0	0.0287	15.8	2.1	15.6	1.6	7.2	-9.0	67.3	21.1	-14.6	0.25	0.75	27.5	28.2	-13.0	-13.3	-16.7														
240417K	9.3	0.1024	16.4	1.4	16.4	1.0	8.4	-7.1	90.2	17.3	-14.9	0.33	0.67	26.2	25.7	-12.5	-12.3	-16.3														
240417F	9.5	0.1493	16.1	1.8	16.2	1.6	5.5	-4.8	109.8	17.4	-15.8	0.24	0.76	25.2	22.9	-13.4	-13.2	-17.6														
240417G	9.8	0.2545	16.8	1.9	16.7	1.9	4.5	-5.3	142.6	15.5	-18.0	0.23	0.77	17.7	20.0	-14.8	-15.2	-20.0														
240417H	10.0	0.3015	15.8	1.9	15.9	1.4	3.2	-7.4	147.1	6.4	-19.6	0.33	0.67	8.0	9.6	-14.8	-15.6	-21.5														
240417L	10.4	0.2092	16.4	2.1	16.2	1.6	2.6	-9.5	147.7	2.4	-19.5	0.52	0.48	6.7	5.0	-15.2	-14.3	-21.6														
<i>Bottle - saponite</i>																																
20140624F	9.5	0.2372	7.9	13.5	7.8	13.1	20.9	13.9	121.7	15.7	-3.0	0.57	0.43	32.5	36.7	6.0	6.7	-16.5														

n.m. = not measured

* Calculated by mass balance

¹2SD 1.1‰, ²2SD 1.5‰

191 2.3. NMR

192 Room temperature solid-state magic-angle spinning (MAS) ^7Li -NMR (nuclear magnetic resonance) experiments
193 for hectorite (Clay Minerals Society Source Clay SHCa-1), 20140624F and its residue were conducted on a Varian
194 Infinity spectrometer at a magnetic field strength of 9.39 T operating at a spectrometer frequency of 155.47 MHz.
195 The samples were packed into 4.0 mm zirconia rotors and spun at 10 kHz in a double resonance MAS NMR probe.
196 Spectra were secondary referenced to spodumene at -1.0 ppm on a ppm scale where 0.5 M LiCl (in DI water) is 0.0
197 ppm, and were acquired using a single-pulse experiment with a $\pi/6$ pulse (3 μs) and a 16 s recycle delay. Similarly,
198 ^7Li MAS NMR experiments for hectorite, the empty rotors, 010217F and its residue, and a rerun of 20140624F took
199 place on a Varian Infinity-plus spectrometer operating at a spectrometer frequency of 194.21 MHz at a magnetic field
200 strength of 11.74 T. The same 4.0 mm rotors as in the previous NMR experiment were used but with a triple resonance
201 MAS NMR probe. Again, samples were spun at 10 kHz and spectra were acquired in a single pulse experiment with
202 a $\pi/6$ pulse and recycle delay of 16 s. As the ceramic rotors and probe components could contain Li impurities at
203 concentrations exceeding those of the samples studied, the empty rotors were analysed under the same conditions as
204 the samples. Spectra acquired for the empty rotors were indistinguishable from noise and so were not subtracted from
205 the sample spectra. Peak fitting was performed using the multi-peak fit package in Igor Pro (WaveMetrics) which
206 uses a non-linear least squares approach to fit Gaussian and Lorentzian peaks to characteristic features observed in
207 the NMR spectra. The percentage of Li removed from each site as a result of treatment with NH_4Cl was calculated
208 based on the difference in number of Li atoms between the bulk and residue spectra which was calculated as follows:
209 the areas fitted to each peak were normalised to the number of scans acquired for each sample and the mass of Li
210 analysed in each site was calculated using a quantified spectrum of a known mass of hectorite. The mass of Li in each
211 site was then normalised to the mass of sample analysed to obtain the concentrations of Li in each site (mg/kg) and
212 subsequently the number of Li atoms in each site.

213 3. Results

214 3.1. Solid characterisation

215 Fourier Transform Infrared (FT-IR) spectroscopy and X-ray diffraction (XRD) were used to characterise the nature
216 of the synthetic materials formed. Transmission FT-IR spectra of all products were characterised by principal Si-O
217 stretches ($\sim 1000\text{-}1020\text{ cm}^{-1}$, Fig. 2a), consistent with a 2:1 layer structure and Mg occupancy in trioctahedral sheets
218 ($\text{Mg}_3\text{-OH}$ stretch at 3680 cm^{-1} , Fig. 2a), as well as corresponding lattice vibration features at 544 and 464 cm^{-1}
219 (Wilkins and Ito, 1967; Farmer, 1974; Russell and Fraser, 1994; Tosca and Masterson, 2014). The products exhibit
220 clear hkl peaks at 10-15, 4.47, 3.51, 2.56 and $\sim 1.52\text{ \AA}$ (Fig. 2b) in powder XRD data (random mounts). These
221 are each produced from the overlapping of multiple reflections and peak broadening produced by a relatively small
222 coherent scattering domain and therefore small particle size (a few to one hundred nanometers). In addition, all
223 diffraction peaks are significantly broadened relative to phases that possess long-range crystalline order, indicating

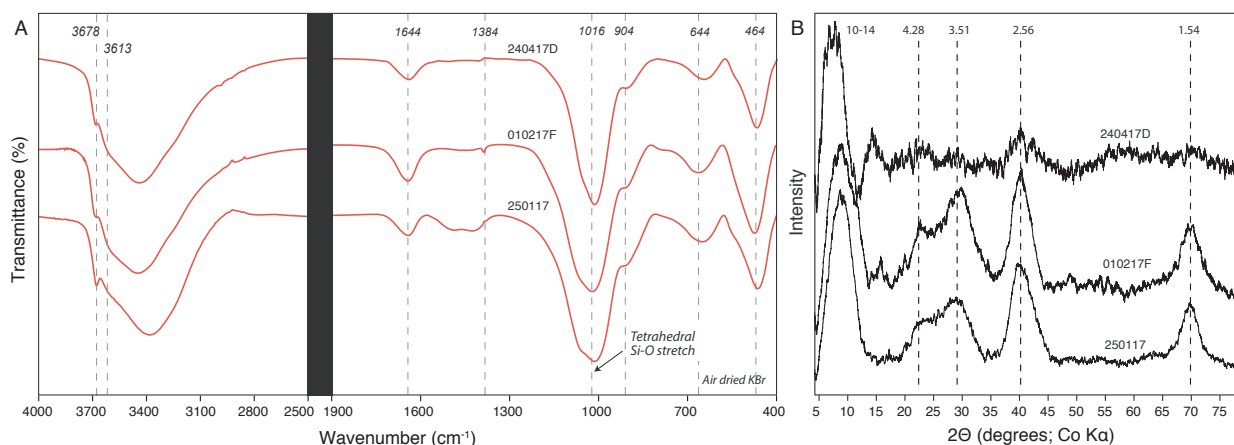


Figure 2: Three representative FT-IR (A) and powder XRD (random mounts) (B) spectra for the bulk solids formed. The key peak positions indicating the formation of a 2:1 layer trioctahedral Mg layer silicate mineral are indicated.

224 that, in general, the products are poorly crystalline. The peak positioned at $\sim 1.52\text{-}1.54 \text{ \AA}$ is particularly informative
 225 because it provides independent yet complimentary evidence to FT-IR (i.e. the $\text{Mg}_3\text{-OH}$ stretch at $\sim 3680 \text{ cm}^{-1}$) that
 226 the synthetic materials have developed an ordered trioctahedral layer. Where present, the diffraction peak at $\sim 10\text{-}15$
 227 \AA provides critical information as to the nature of the precipitate. Typical powder XRD peak positions (uncorrected
 228 for polarisation effects) are commonly located at either $\sim 10 \text{ \AA}$ or $\sim 14 \text{ \AA}$. As discussed in section 1.1, these peak
 229 positions are consistent with kerolite-type and stevensite-type structures, respectively. However, XRD data from
 230 oriented clay mounts suggest that this peak is largely unresponsive to ethylene glycol treatment, even after several
 231 weeks of exposure to ethylene glycol vapour at 60°C . This indicates that although the development of layer stacking
 232 order is evident along the 001 direction, ethylene glycol absorption does not take place. This may be related to strong
 233 hydration of Mg-silicate nuclei interlayers and edges (cf. TGA-DTA data in Tosca and Masterson, 2014) which may
 234 inhibit ethylene glycol absorption.

235 3.2. Initial, final and exchangeable solutions

236 Due to variable $\delta^7\text{Li}$ in the starting solutions, all data is reported as the difference between solid and initial solution
 237 $\delta^7\text{Li}$ values ($\Delta^7\text{Li}_{x\text{-solution}}$). The fraction of Li from the initial solution removed into clay is ≤ 0.005 (Table 2) and
 238 therefore the observed variations in Li isotope fractionation are not due to variations in the fraction of Li uptake. The
 239 experiments were terminated once successive concentration measurements were within measurement error of each
 240 other, i.e. they had reached steady-state with respect to concentration measurements (Figure S1).

241 The concentration of exchangeable Li varied from 3 to 51 mg kg^{-1} (excluding the reaction vessel experiment,
 242 section 3.5). Exchangeable $\delta^7\text{Li}$ values were higher than the corresponding bulk $\delta^7\text{Li}$ value but were lower than the
 243 starting solution $\delta^7\text{Li}$ value by an average of 7‰ (Fig. 3, Table 2).

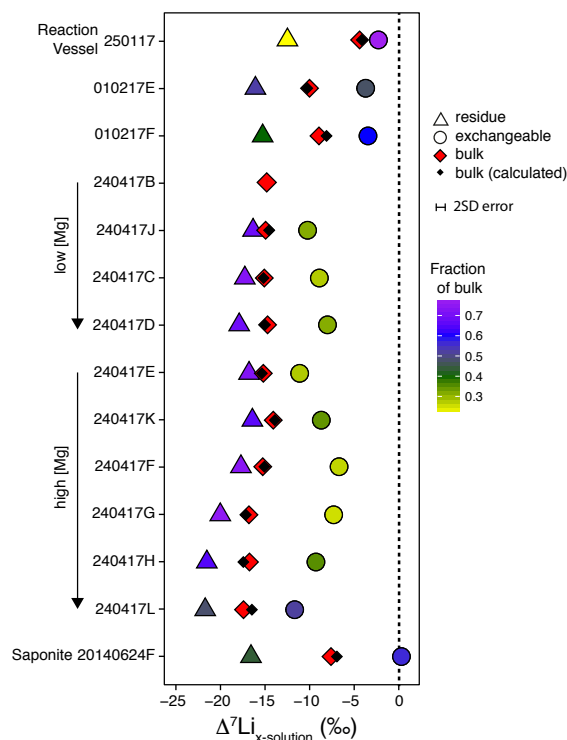


Figure 3: Summary of the Li isotopic compositions for the exchangeable, residual and bulk phases for each of the experiments. The calculated $\delta^7\text{Li}$ derived from mass balance is shown for comparison. The colour gradient of the exchangeable and structural (residue) Li pools reflects the fractional contribution of those pools to the bulk. All Li isotopic compositions are plotted relative to the initial solution which is at 0‰ in this plot and highlighted by the dotted line. The arrows for the high and low [Mg] series of experiments indicate increasing pH towards the head of the arrow.

244 3.3. NMR to assess the efficiency of the cation exchange procedure

245 The efficiency of the removal of exchangeable ions by NH_4Cl was checked using ^7Li -NMR. Due to the differing
 246 bonding environments, the induced magnetic field experienced by structural (octahedral) Li is expected to be different
 247 compared to exchangeable Li and therefore have a different chemical shift. We observe three peaks in the ^7Li -NMR
 248 spectra (Fig. 4) of the solid recovered from both a saponite (20140624F) and a stevensite (010217F) experiment
 249 demonstrating that three bonding environments are present. Based on a comparison with the spectra of the hectorite
 250 standard SHCa-1, which only contains octahedral Li, we conclude that the largest peak (peak 3), with the most
 251 negative shift, is the octahedral site (Fig. 4). This peak is also least affected by the exchange procedure (Fig. 4, Table
 252 3), consistent with Li incorporated into the clay structure. The assignment of the remaining two peaks is discussed in
 253 section 4.1.

254 The peak area is proportional to the number of Li atoms in each bonding environment and the results of the peak
 255 fitting are used to calculate the relative proportion of Li in each (Table 3). For the 20140624F sample, addition of
 256 NH_4Cl resulted in $81\pm 6\%$, $58\pm 16\%$ and $3\pm 5\%$ loss of Li from peaks 1, 2 and 3 respectively and for the 010217F
 257 sample the equivalent values were $66\pm 3\%$, $62\pm 9\%$ and $18\pm 2\%$ (Table 3). The significant loss of Li from peaks 1

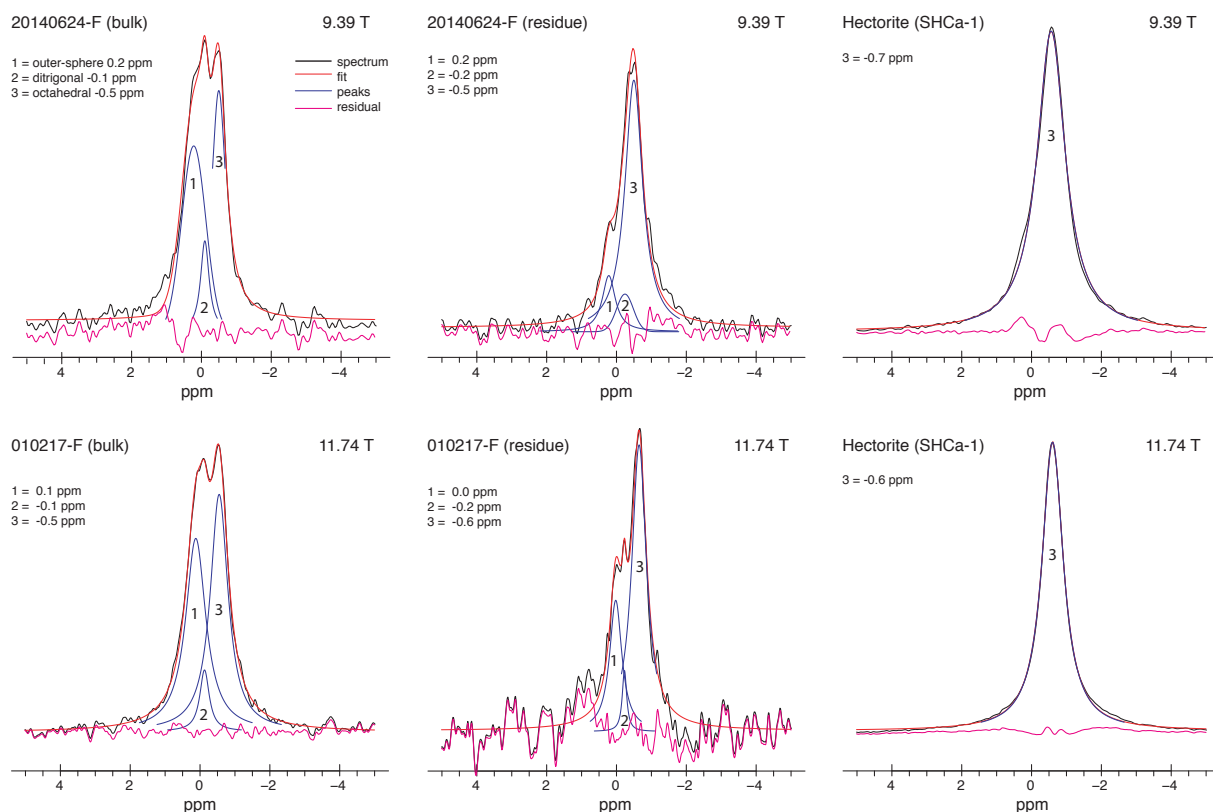


Figure 4: NMR spectra. Top row spectra were collected at 9.39 T (400 MHz) whereas bottom row spectra were collected at 11.74 T (500 MHz). Note the decrease in both the outer-sphere and pseudo-hexagonal peaks (peaks 1 and 2) in the residual (after treatment with NH_4Cl) sample spectra compared to the bulk sample spectra. The resolution (FWHM - full width at half maximum) in peak position is ± 0.1 ppm.

258 and 2 supports the contention that these sites represent exchangeable Li. Although a significant proportion of Li from
 259 peaks 1 and 2 is removed as a result of the exchange procedure, 13-26% peak 1 Li and 4-7% peak 2 Li remains in
 260 the residual solid (Table 3). Assuming these samples are representative, the exchange procedure, although effective,
 261 has not removed all of the exchangeable Li and some remains in the residual solid. Equally, some octahedral Li was
 262 lost during the exchange procedure. The total fraction of exchangeable Li calculated from NMR ($40\pm 7\%$ 20140624F,
 263 $43\pm 3\%$ 010217F) is in agreement with that calculated from concentration measurements ($57\pm 4\%$ 20140624F, $61\pm 3\%$
 264 010217F (f_e in Table 2), Table 3).

265 3.4. Bulk and residual solids

266 All solids were enriched in ^6Li compared to the starting solution (Fig. 3, Table 2). The bulk solids had $\delta^7\text{Li}$
 267 values 4.4 to 17.3‰ lower than the initial solution and the $\delta^7\text{Li}$ values of the residual solids were lower than the
 268 corresponding bulk $\delta^7\text{Li}$ value by 1.4 to 9.0‰ (Fig. 3, Table 2). The bulk solids contained 7 to 184 mg kg^{-1} Li whilst
 269 the concentration of Li in the residual phase varied from 2 to 57 mg kg^{-1} (Table 2).

270 There is no relationship between pH and $\Delta^7\text{Li}_{\text{bulk-solution}}$ values (Fig. 5a), even when pH was varied systematically
 271 ('240417' series). However, covariation between pH and $\Delta^7\text{Li}_{\text{residue-solution}}$ values (exchangeable Li removed) is

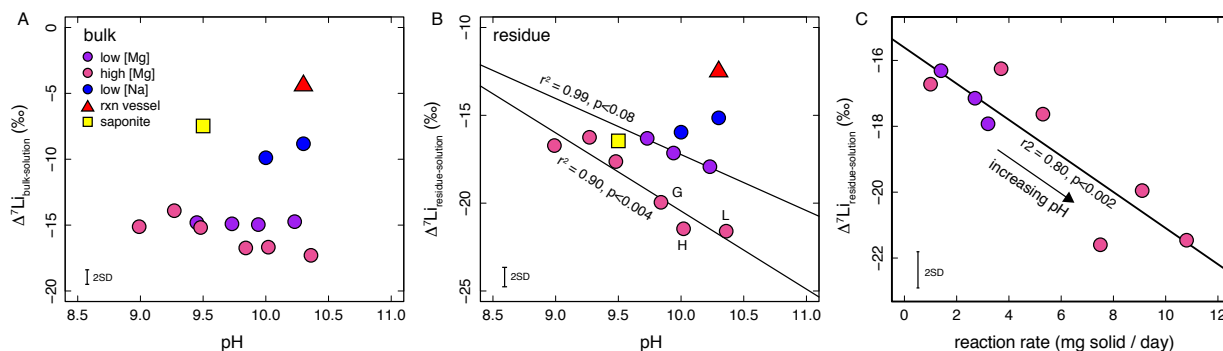


Figure 5: Dependence of $\delta^7\text{Li}$ values on pH in the bulk (A) and residual (B) solids. In the high [Mg] series of experiments there is a trend of decreasing $\Delta^7\text{Li}_{\text{residue-solution}}$ values with increasing pH. (C) Higher reaction rates (mass of solid produced per day) are associated with lower $\Delta^7\text{Li}_{\text{residue-solution}}$ values.

272 observed in the ‘240417’ series of experiments (Fig. 5b). Whilst reaction rate could be one factor affecting the
 273 $\Delta^7\text{Li}_{\text{residue-solution}}$ values (Fig. 5c), another factor could be the solution chemistry. The stevensite experiments were
 274 conducted at constant Li concentration in the initial solution but with varying Na, Mg, and H^+ concentrations (Table
 275 1). For similar pH (10.0 - 10.4), higher [Na] concentrations in the initial solution were associated with a greater
 276 difference in $\delta^7\text{Li}$ values between the bulk solid and the initial solution (Fig. 6a). For pairs of experiments conducted
 277 at similar pH, higher [Mg] is linked to lower $\Delta^7\text{Li}_{\text{residue-solution}}$ and lower Li/Mg values in both bulk and residual
 278 solids (Table 2). The Li/Mg ratios of the residual solids ranged between 1.6×10^{-5} and 5.3×10^{-4} $\text{mg kg}^{-1}/\text{mg kg}^{-1}$ and
 279 there is a relationship between the Li/Mg ratios of the residual solids and $\Delta^7\text{Li}_{\text{residue-solution}}$ values in experiments with
 280 constant Mg concentration (Fig. 7).

281 3.5. Mass balance

282 The exchangeable and residual Li concentrations (calculated relative to the bulk sample) were summed and compared
 283 to the measured bulk concentrations. Calculated bulk Li concentrations are within 15% of measured bulk Li
 284 values except for 240417H (18%), 240417L (25%) and the reaction vessel experiment (72%). The poor agreement
 285 between calculated and measured bulk values for the reaction vessel experiment implies the results from this experi-
 286 ment should be treated cautiously. The percentage of Li in the exchangeable and residual pools was calculated relative
 287 to the calculated bulk value and these were used to calculate the isotopic composition of the bulk which agreed with
 288 the measured value to within 1.5‰ (Fig. 3), demonstrating the self-consistency of the methodology.

289 4. Discussion

290 Our experiments confirm the results of previous laboratory experiments (Vigier et al., 2008) which found that hec-
 291 torite preferentially incorporates ^6Li and are also in agreement with the direction of Li isotope fractionation inferred
 292 from field studies (e.g. Huh et al., 2001) and modelling studies (Dupuis et al., 2017). There is considerable variation

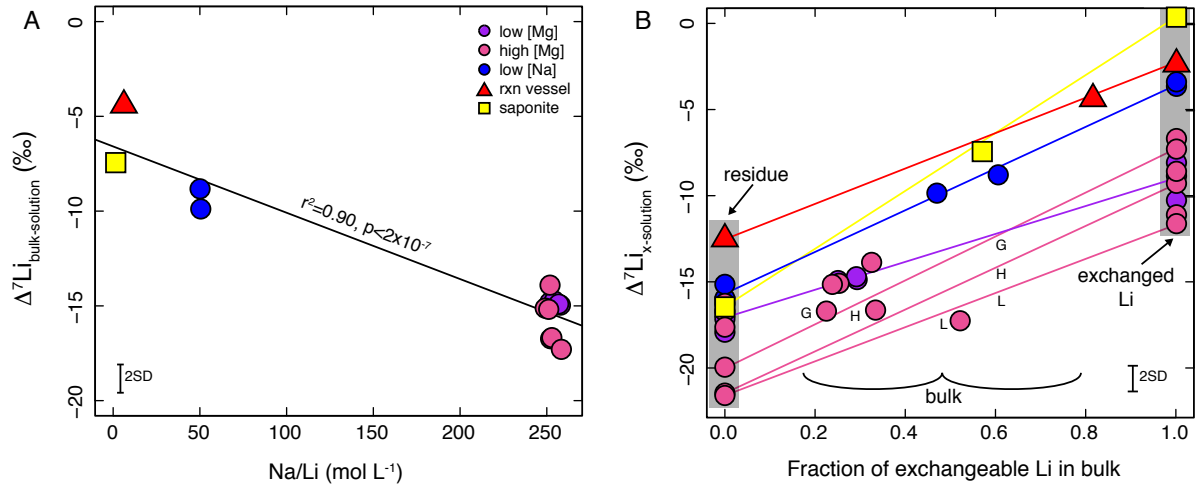


Figure 6: (A) Plot of $\Delta^7\text{Li}_{\text{bulk-solution}}$ vs Na/Li in the initial solution. The more Na, the lower the $\Delta^7\text{Li}_{\text{bulk-solution}}$ value consistent with less outer-sphere Li. (B) Plot of $\Delta^7\text{Li}$ versus the fraction of exchangeable Li. The greater the fraction of exchangeable Li, the higher the $\Delta^7\text{Li}_{\text{bulk-solution}}$ value consistent with less competition from Na. $\Delta^7\text{Li}_{\text{bulk-solution}}$ values are a mixture between residual Li (octahedral \gg outer-sphere $>$ pseudo-hexagonal) and exchangeable Li (outer-sphere \gg pseudo-hexagonal \approx octahedral). The $\Delta^7\text{Li}_{\text{bulk-solution}}$ values are within error of the lines connecting these two end-members. For the low [Na] (blue circles) and the [Mg] experiments (pink/purple circles, excluding G, H and L), the average for each end-member (residual and exchangeable) is used. Variations in the positions and slopes of the lines are likely due to varying proportions of the three Li sites (octahedral, pseudo-hexagonal and outer-sphere) in each of the end-members. For example, 20140624F (yellow square) contains more outer-sphere Li ($\delta^7\text{Li} \sim 0\text{‰}$, Section 4.1) than 010217F (blue circle) in the exchangeable fraction but less outer-sphere Li in the residue (Table 3) resulting in a mixing line with a steeper slope.

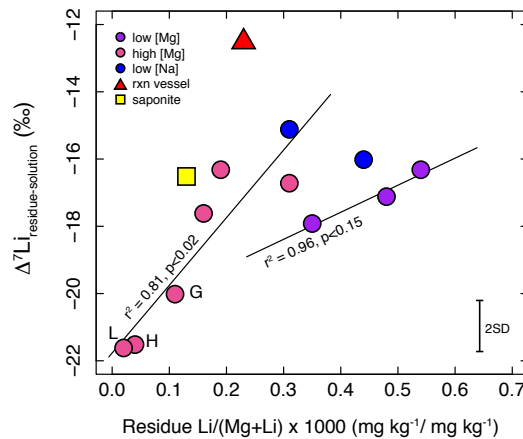


Figure 7: Dependence of $\delta^7\text{Li}$ values in the residual phase on solid composition. In sets of experiments where only pH varies (pink and purple points), $\Delta^7\text{Li}_{\text{residue-solution}}$ values increase with increasing Li/(Mg+Li).

Table 3: Results from ^7Li -NMR peak fitting. Peak numbers refer to Fig. 4.

Experiment	Sample	Octahedral	Pseudo-hexagonal	Outer-sphere	Li removed		$\Delta^7\text{Li}_{\text{bulk-solution}}$
		Peak 3 %	Peak 2 %	Peak 1 %	NMR (%)	Conc ¹ (%)	‰
20140624F	Bulk	49±1	11±1	40±1			-7.5
	Residue	80±1	7±1	13±1			-16.5
	Exchangeable ²	4±6	15±5	81±15			0.4
	Removed ³	3±5	58±16	81±6	40±7	57±4	
010217F	Bulk	48±0	7±0	45±0			-8.8
	Residue	69±1	4±1	26±1			-15.1
	Exchangeable ²	21±3	10±2	70±5			-3.3
	Removed ³	18±2	62±9	66±3	43±3	61±4	

¹Calculated by mass balance based on measured Li concentrations (Table 2).

²Calculated based on loss of Li from bulk.

³Percentage of Li removed from a given bulk spectra peak as a result of the exchange procedure.

293 in the residual and bulk $\Delta^7\text{Li}$ values (Fig. 3) and in the following sections we will discuss some experimental fac-
 294 tors, including the bonding environment and chemistry of the initial solution (including pH), which could cause this
 295 variation.

296 4.1. Li fractionation associated with different bonding environments

297 In a layer silicate structure Li is located in octahedral sites. Due to its small size Li can also be located in pseudo-
 298 hexagonal sites (ditrigronal cavity) in the tetrahedral sheet, interlayer sites between TOT layers and be adsorbed onto
 299 surfaces (Fig. 1, Meunier, 2005). The combination of isotopic analyses of exchangeable solutions, bulk and residual
 300 solids, coupled to ^7Li -NMR provides a unique way to deconvolve the relative Li isotope fractionation between distinct
 301 bonding environments. Three peaks are observed in the ^7Li -NMR spectra and two of these decrease markedly in area
 302 after saturation with NH_4 (peaks 1 and 2, Fig. 4), confirming that the exchange procedure was effective at removing
 303 exchangeable Li but that some remained in the residual solid (Table 3). If the uptake of Li into a non-octahedral
 304 site has a unique fractionation factor then the presence of Li in these sites in the residual phase would mean that the
 305 $\delta^7\text{Li}$ value of the residual phase can no longer be assumed to represent fractionation into octahedral sites only. Below
 306 we discuss what is known about the bonding environments of Li to assign each ^7Li -NMR peak to a specific bonding
 307 environment and assess the magnitude of Li isotope fractionation in the three environments relative to each other.

308 Peaks 1 and 2 in the ^7Li -NMR spectra correspond to exchangeable Li. These were assigned based on coordination
 309 number as chemical shift is dependent on coordination number, with a lower coordination number resulting in less
 310 shielding and a more positive chemical shift (Xu and Stebbins, 1995). Chemical shift ranges for Li with coordination
 311 numbers 4 and 6 taken from ^6Li -NMR spectra (same chemical shift as ^7Li -NMR) are 0 to 1 ppm and -0.3 to -1.3
 312 ppm, respectively (Xu and Stebbins, 1995). For smectites, both interlayer and adsorbed Li are thought to exist as
 313 outer-sphere complexes with a coordination number of 4 (hydrated Li, Pistiner and Henderson, 2003; Chan and Hein,
 314 2007) and are therefore expected to experience the same chemical shift. In the following discussion these two sites are
 315 therefore grouped together as ‘outer-sphere’ sites. In contrast, pseudo-hexagonal Li exists in an inner-sphere complex
 316 with O^{2-} atoms and, due to distortions, has an effective coordination number of 6 (Meunier, 2005). Allowing for the

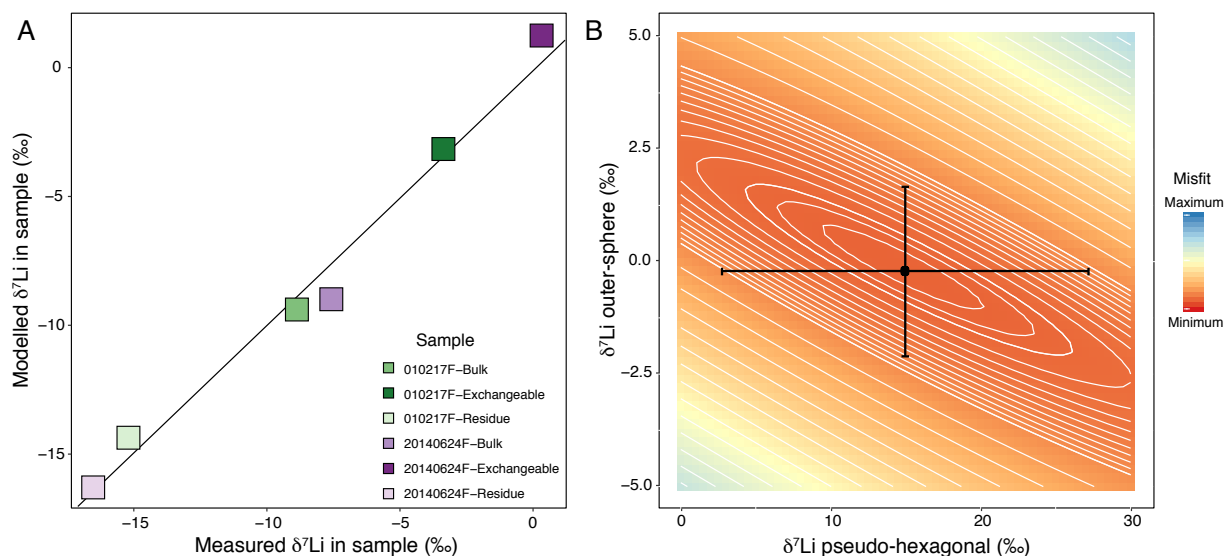


Figure 8: (a) Comparison of the measured $\delta^7\text{Li}$ values (Table 2) with the modelled $\delta^7\text{Li}$ values of the samples, calculated using the fractionation factors obtained through a Monte Carlo minimum misfit method and the proportions of Li in each bonding environment calculated from ^7Li -NMR (Table 3). The black line is the 1:1 line. (b) Example of a misfit contour plot in 2-D space (octahedral $\delta^7\text{Li}$ value set at the minimum misfit value of -21.5‰ in order to make a 2-D plot). The calculation of the misfit is described in the supplementary information. The black point indicates the optimum (smallest misfit) values for outer-sphere and pseudo-hexagonal bonding environments. 1σ error bars are also shown.

317 fact that ^7Li -NMR peaks are broader than ^6Li -NMR peaks, the chemical shift of the most positive peak (peak 1: 0.0 to
 318 0.2 ppm) is in agreement with 4-fold coordinated outer-sphere Li and the negative chemical shift of the middle peak
 319 (peak 2: -0.1 to -0.2), 6-fold coordinated pseudo-hexagonal Li. The assignment of peak 2 to pseudo-hexagonal Li
 320 is consistent with this site having a more similar bonding environment (same coordination number) to octahedral Li
 321 compared to outer-sphere Li and therefore its chemical shift is closer to the octahedral peak (peak 3). The contrasting
 322 $\delta^7\text{Li}$ values between exchange solutions and residual solids implies that each bonding environment has its own isotope
 323 fractionation factor. Our data cannot directly constrain the three fractionation factors but a system of linear equations
 324 can be solved based on the parameters in Table 3 (6 equations, 3 unknowns). A Monte Carlo minimum misfit method
 325 (Supplementary Information) was employed. The fractionation factors for the three bonding environments determined
 326 using this method were: octahedral $-21.5 \pm 1.1\text{‰}$, outer-sphere $-0.2 \pm 1.9\text{‰}$ and pseudo-hexagonal $15.0 \pm 12.3\text{‰}$ (errors
 327 1σ , Fig. 8).

328 Previous studies have suggested that both interlayer Li and Li adsorbed to smectites have the same isotopic com-
 329 position as the starting solution since Li in both cases is held in weakly bonded outer-sphere complexes (Pistiner
 330 and Henderson, 2003; Chan and Hein, 2007). Interlayer Li should in theory retain its 4-fold coordinated hydration
 331 sphere (and therefore not be fractionated), however it has been suggested that in addition to outer-sphere complexes,
 332 inner-sphere (dehydrated) complexes may also form in the interlayer where the solvation number can vary from 2 to
 333 4 (Greathouse and Sposito, 1998). If the coordination number of Li in the interlayer does decrease below 4, then this
 334 Li will be enriched in ^7Li relative to the initial solution (Wimpenny et al., 2015). Our fitting results suggest that the

335 fractionation factor for outer-sphere complexes is within error of zero, supporting the former hypothesis; that inter-
336 layer Li is unfractionated with respect to the starting solution (Pistiner and Henderson, 2003; Vigier et al., 2008), and
337 therefore likely remains in a 4-fold coordination.

338 Whilst outer-sphere Li is not fractionated significantly relative to the starting solution, octahedral Li is, with a value
339 of $-21.5 \pm 1.1\%$. This finding is consistent with isotope equilibrium theory where strong bonds with a high vibrational
340 frequency will always favour the heavy isotope because the vibrational frequency of the bond scales with the square
341 root of the bond strength over reduced mass (Bigeleisen, 1965). Typically, bond strength scales with bond length,
342 such that shorter bonds are stronger. Li in octahedral sites is bonded to oxygen atoms in a six-fold coordination with
343 a bond length of around 2.08 \AA (sodium fluorohectorite, Kalo et al., 2012) whereas Li in the aqueous phase exists
344 predominantly as the $[\text{Li}(\text{H}_2\text{O})_4]^+$ species with a bond length of around 1.9 \AA (Rudolph et al., 1995; Mähler and
345 Persson, 2012). Therefore octahedral Li, with a longer, weaker bond will be enriched in ^6Li compared to the initial
346 solution as calculated, and predicted by both density functional theory (DFT) calculations (Dupuis et al., 2017) and
347 experimental data (Fig. 3, Vigier et al., 2008).

348 Pseudo-hexagonal Li has been assumed to have the same fractionation factor as the octahedral site (Williams and
349 Hervig, 2005) based on both sites having a coordination number of 6 (Meunier, 2005). Nevertheless, the bonding
350 environment of the pseudo-hexagonal and octahedral sites is different as evinced by the different chemical shifts in
351 ^7Li -NMR spectra (Fig. 4, Theng et al., 1997) and could therefore be expected to have different fractionation factors.
352 The looser bound pseudo-hexagonal Li may have longer Li–O bonds compared to the octahedral sites, and this is
353 supported by a DFT calculation demonstrating that the Li sits asymmetrically in the pseudo-hexagonal site with Li–O
354 bond lengths varying from 2.01 to 3.91 \AA (Wungu et al., 2011). This would result in a longer average bond length
355 compared to octahedral Li and we would therefore predict a lower $\delta^7\text{Li}$ value. However, the results from the Monte
356 Carlo calculation suggest that pseudo-hexagonal Li is enriched in ^7Li compared to the initial solution (Fig. 8a),
357 which is inconsistent with the ^7Li -NMR peak position. The bonding environment with lowest coordination number
358 (outer-sphere Li) should have the highest chemical shift (Xu and Stebbins, 1995) and therefore the highest $\delta^7\text{Li}$ value.
359 However, we note that the calculated isotope fractionation for the pseudo-hexagonal site is subject to large uncertainty
360 which strongly co-varies with the $\delta^7\text{Li}$ value of outer-sphere Li (Fig. 8b). If the pseudo-hexagonal value was reduced
361 to 0% , this would shift the outer-sphere $\delta^7\text{Li}$ value by $<2.5\%$ (Fig. 8b). As more synthetic clay samples are measured
362 by ^7Li -NMR and for $\delta^7\text{Li}$, the uncertainty in the pseudo-hexagonal $\delta^7\text{Li}$ value will reduce.

363 In summary, there are unique Li isotope fractionation factors for the three bonding environments observed in ^7Li -
364 NMR spectra (octahedral, outer-sphere and pseudo-hexagonal) and bulk, exchangeable and residual phases contain
365 variable proportions of all three, leading to the observed variability in $\delta^7\text{Li}$ values in these phases (Fig. 3). Therefore
366 it is essential that the potential for remaining interlayer, adsorbed and pseudo-hexagonal Li is taken into account when
367 using the residual phase to infer a fractionation factor for Li uptake into octahedral sites.

368 4.2. pH dependence

369 The main experimental variable which changed between the different experiments was pH. Increasing pH increases
370 nucleation rate, growth rate and promotes better crystallisation and therefore reaction rates scale with pH (Table 2).
371 There is no correlation between pH and $\Delta^7\text{Li}_{\text{bulk-solution}}$ and this is most likely due to the variable contribution of
372 exchangeable Li, with a distinct fractionation factor, to the bulk Li isotopic composition (Fig. 5a). However, there
373 is a strong relationship between pH and $\Delta^7\text{Li}_{\text{residue-solution}}$ (Fig. 5b). It is possible that the trend of more negative
374 $\Delta^7\text{Li}_{\text{residue-solution}}$ values with increasing pH observed in the '240417' series of experiments (low and high [Mg],
375 Fig. 5b) is the result of a kinetic isotope fractionation effect. Indeed, for these experiments, the $\Delta^7\text{Li}_{\text{residue-solution}}$
376 values decreased as reaction rate (mg solid formed per day) increased (Fig. 5c), consistent with increased kinetic
377 fractionation at higher pH as observed for Ca isotope fractionation during calcite synthesis (Tang et al., 2008; Nielsen
378 et al., 2012).

379 However, pH can also affect the Li speciation in solution (Bogatko et al., 2013) and a recent study attributed
380 a relationship between $\delta^7\text{Li}$ and pH in foraminifera to changes in Li speciation (Roberts et al., 2018). The aqueous
381 speciation of Li is not well defined but Li can be considered to be a very weak acidic cation with aqua (e.g. $\text{Li}(\text{H}_2\text{O})_x^+$)
382 and hydroxo forms (e.g. $\text{Li}(\text{OH})(\text{H}_2\text{O})_{x-1}$), most likely with a secondary hydration shell (Bogatko et al., 2013). These
383 two forms have different desolvation energies and as the isotope exchange equilibrium constant is greater than 1,
384 the isotopic composition of a solid precipitating from solution will depend on the relative proportion of these two
385 forms in solution. Bogatko et al. (2013) calculated that for the precipitation of carbonate from solution at 25°C
386 the $\Delta^7\text{Li}_{\text{CO}_3^{2-}-\text{H}_2\text{O}}$ value decreased when, depending in the exact model parameters used, pH increased above ~9.
387 Therefore the lower $\Delta^7\text{Li}_{\text{residue-solution}}$ values we observe in this study when increasing pH from 9.0 to 10.4 (Fig. 5b)
388 would be consistent with a change in aqueous Li speciation. As pH increases the Mg/Li ratio of the residual solid
389 also increases which may additionally impact on the $\Delta^7\text{Li}_{\text{residue-solution}}$ value (Section 4.3). Further experimental work
390 would be required to assess whether equilibrium or kinetic fractionation is dominant.

391 4.3. Composition dependence

392 The initial solution composition will likely affect the crystallisation process and our experiments were conducted
393 at a range of Si, Na, Mg and Al concentrations (Table 1). For the bulk solids, lower Na concentrations in the initial
394 solutions are associated with smaller $\Delta^7\text{Li}_{\text{bulk-solution}}$ values (Fig. 6a) and the lower the Na concentration, the greater
395 the fraction of exchangeable Li in the bulk solid (Fig. 6b). These observations may be a result of competition between
396 Na and Li for interlayer sites (Tutolo and Tosca, 2018). If it is assumed that interlayer Li has an isotopic composition
397 similar to the initial solution, an increased fraction of interlayer Li in the bulk solid would increase the bulk Li isotopic
398 composition. On the other hand, in the experiments with high Na concentrations, Na outcompetes Li, reducing the
399 fraction of interlayer Li in the bulk solid. After treatment to remove exchangeable cations, we do not find significant
400 differences in $\Delta^7\text{Li}_{\text{residue-solution}}$ values between experiments with different initial Na concentrations (Fig. 6b), if the
401 reaction vessel and three high pH, high [Mg] experiments are excluded due to additional factors related to pH (Section

402 4.2). The lack of control of initial solution compositions on the residual solid Li isotopic composition is in agreement
403 with previous studies (Vigier et al., 2008; Wunder et al., 2006).

404 The composition of the solid itself appears to have a negligible effect on $\Delta^7\text{Li}_{\text{residue-solution}}$ values: the synthetic
405 saponite has a similar value to the low [Mg] stevensite experiments (Fig. 3). This is in agreement with a modelling
406 study which predicted only a minor difference ($\leq 1.0\text{‰}$) between a Li-Mg and a Li-Al phyllosilicate (Dupuis et al.,
407 2017). However, there is a hint in the data that the balance between Mg and Li in the residual solids could affect
408 the $\Delta^7\text{Li}_{\text{residue-solution}}$ value, with lower Li/Mg ratios associated with lower $\Delta^7\text{Li}_{\text{residue-solution}}$ values (Fig. 7). Even at
409 equilibrium, a change in element ratio is predicted to affect isotope fractionation due to changes in bond lengths as
410 the crystal structure distorts to accommodate an ion with a different charge and/or radius (Perdikatsis and Burzlaff,
411 1981; Brigatti et al., 2000; Laurora et al., 2011; Michalski et al., 2015; Wang et al., 2017a,b). A decrease in the Li
412 content of the residual solid could therefore cause the Li-O bond length to change, impacting the fractionation factor
413 for the octahedral site. It is also important to consider exactly how the Li ends up in an octahedral site. A study on
414 Mg-Fe substitution in dioctahedral clay minerals found that entering into a vacant octahedral site or substituting into
415 an already filled octahedral site had opposite effects on bond length (Michalski et al., 2015). Further modelling and
416 experimental data would be required to see if changes in bond length could explain the relationship between Li/Mg
417 and $\Delta^7\text{Li}_{\text{residue-solution}}$ values (Fig. 7).

418 However, bond length arguments assume equilibrium isotope fractionation is occurring and it also assumes that
419 all Li in the residue is in octahedral sites. As we discussed in section 4.1, variable proportions of exchangeable Li
420 remaining in the residual solid could impact on the Li isotopic composition of the residue. Based on the fact that
421 the $\Delta^7\text{Li}_{\text{residue-solution}}$ values of H and L (-21.5 and -21.6‰) are within error of the fractionation calculated for the
422 octahedral site (-21.5‰), these samples may contain a negligible proportion of exchangeable Li. The higher reaction
423 rate of these experiments (Fig. 5c) could have resulted in larger particles decreasing the proportion of exchangeable
424 Li held on surface and edge sites. The high pH of experiments G, H and L may also change the speciation of Li in
425 solution (section 4.2) which could contribute to the trend observed in Fig. 7.

426 4.4. Temperature dependence

427 In equilibrium isotope fractionation there is often a linear relationship between the fractionation factor and the
428 reciprocal of temperature squared, with greater fractionation at lower temperatures (Schauble, 2004). Vigier et al.
429 (2008) argued that the good reproducibility of several experiments conducted at a single temperature under different
430 conditions indicated the lack of kinetic isotope effects and that the synthesised clays had reached isotopic equilibrium.
431 Our experiments reach chemical equilibrium (Figure S1) and the majority of the isotope data also reproduce very
432 well (Fig. 9). The overall average $\Delta^7\text{Li}_{\text{residue-solution}}$ is $-17.3 \pm 5.0\text{‰}$ (2SD), where this value represents a mixture of
433 octahedral and exchangeable Li. The lower $\Delta^7\text{Li}_{\text{residue-solution}}$ values of samples G, H and L and the higher value
434 for the reaction vessel experiment compared to the other samples could relate to kinetic or equilibrium controls as
435 discussed in sections 4.2 and 4.3. If these values are excluded, the average $\Delta^7\text{Li}_{\text{residue-solution}}$ is $-16.6 \pm 1.7\text{‰}$ (2SD).

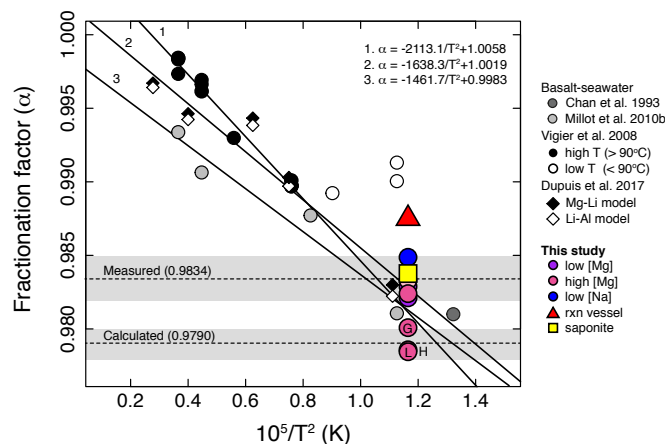


Figure 9: Fractionation factor dependence on temperature. Equilibrium fractionation factors are calculated from the Li isotopic compositions of the initial solution and residual solid using the relationship $\alpha = (1000 - \delta^7\text{Li}_r) / (1000 - \delta^7\text{Li}_i)$. Literature data were compiled from synthesis experiments (Vigier et al., 2008), modelling (Dupuis et al., 2017) and basalt-seawater interaction experiments (Millot et al., 2010b; Chan et al., 1993). In the latter, solids were not treated to remove exchangeable Li. We do not include experiments conducted at high pressure (Wunder et al., 2006, 2007, 2010). Regression lines for each set of literature data are plotted. Line 1 only includes high temperature experiments (see Vigier et al. (2008) for details) and the line for basalt-seawater (line 3) only includes data from Millot et al. (2010b). Note that in the Millot et al. (2010b) study, the solids were not measured directly and were shown to contain a mixture of phases including Li from seawater and the original basalt. The dotted line marked ‘measured’ indicates the average α at 20°C for the measured residual fractions (assuming complete removal of exchangeable Li), excluding the reaction vessel and G, H and L (see text for details). The dotted line marked ‘calculated’ indicates the octahedral fractionation factor calculated using mass balance and ^7Li -NMR data (Section 4.1, Supplementary Information). The grey shaded areas indicate the 2SD error for the ‘measured’ line and the 1 σ error for the ‘calculated’ line.

436 As the fraction of Li lost from solution is ≤ 0.005 , we assume that $\Delta^7\text{Li}_{\text{residue-solution}}$ is equivalent to α with a value
 437 of 0.9834 ± 0.0017 . This value is in good agreement with fractionation factors predicted for 20°C based on linear
 438 regression of previously published experimental data (0.9812 (Vigier et al., 2008) and 0.9813 (Millot et al., 2010b)).
 439 Nevertheless, we know from ^7Li -NMR data that the residual phase contains some exchangeable Li and therefore the
 440 ‘true’ fractionation factor into the octahedral site will be lower than 0.9834, and is calculated to be 0.9790 ± 0.0011
 441 (Section 4.1). For comparison, a value of 0.9828 at 20°C was obtained from a linear regression of results from a
 442 TI-PIMD (thermodynamic integration path-integral based molecular dynamics) (Fig. 9, Dupuis et al., 2017).

443 Vigier et al. (2008) also conducted low temperature experiments but these had $\delta^7\text{Li}$ values approximately 7‰
 444 higher than those predicted based on a regression line defined by the high temperature samples. It was postulated
 445 that this may have been due to low crystallinity and a high proportion of edge octahedra in the solids formed at low
 446 temperature, resulting in additional kinetic fractionation effects. Similar processes may also be impacting our reaction
 447 vessel experiment (red triangle, Fig. 9) where the actual nucleation rate may have been higher as a consequence of
 448 increasing the pH of a weakly acidic solution of Mg and SiO_2 , potentially producing smaller particles with more
 449 defects and a greater proportion of edge octahedra sites. Alternatively, a consequence of the low crystallinity may
 450 be that a greater proportion of exchangeable Li remained in the residual solid after exchange with the preferred NH_4
 451 cation.

452 4.5. Implications for field and modelling studies

453 The $\Delta^7\text{Li}_{\text{solid-solution}}$ value is a critical parameter for modelling weathering processes, in particular the relative
454 balance between dissolution and precipitation reactions (e.g. Bouchez et al., 2013; Dellinger et al., 2015; Pogge von
455 Strandmann and Henderson, 2015). Importantly, as we show in this study, the $\Delta^7\text{Li}_{\text{bulk-solution}}$ value is comprised
456 of three separate fractionation factors corresponding to the three bonding environments visible in ^7Li -NMR spectra:
457 octahedral, pseudo-hexagonal and outer-sphere. The value of $\Delta^7\text{Li}_{\text{solid-solution}}$ in natural systems is thus influenced
458 by the relative proportions of Li in each of these three environments and the consequent variation in $\Delta^7\text{Li}_{\text{solid-solution}}$
459 could have implications for the global Li budget of seawater as illustrated below in a very simplistic manner.

460 The current best estimate of the $\delta^7\text{Li}$ input to the world's oceans is a flux weighted mean from large rivers with
461 a value of $+23\text{‰}$ (Huh et al., 1998), which has been widely adopted in modelling studies (e.g. Li and West, 2014).
462 At a global scale, the average continental rock drained is approximately -0.9‰ (Fig. 10), in agreement with the av-
463 erage $\delta^7\text{Li}$ in the upper continental crust ($0\pm 2\text{‰}$, Teng et al., 2004), and hence $\Delta^7\text{Li}_{\text{solid-solution}}$ has an average value
464 of $-23.9\pm 3.1\text{‰}$ (1σ error). This value is not a fractionation factor, but is a global difference between large river
465 and rock/sediment $\delta^7\text{Li}$ values. Interestingly, this value is within uncertainty of the octahedral fractionation factor
466 calculated in the present study ($-21.5\pm 1.1\text{‰}$). Thus it could be argued that Li incorporation into the octahedral site
467 dominates fractionation at a global scale and that, to a first order, rivers are in equilibrium with their sedimentary
468 products. However, we note that it is unlikely that natural clays will only contain octahedral Li. Exchangeable Li will
469 very likely be present and may be derived from a different fluid than octahedral Li, due to changes in solution com-
470 position during mineral growth (Williams and Hervig, 2005). Therefore an assessment of equilibrium fractionation in
471 field data would first require the removal of exchangeable Li.

472 A further observation is that the difference between seawater ($+30.8\text{‰}$, Rosner et al., 2007) and the riverine input
473 to the oceans based on large rivers ($+23\text{‰}$) is 7.8‰ . The difference between this value and the $\Delta^7\text{Li}_{\text{solid-solution}}$
474 value (-24‰) has been interpreted as a balance between high temperature formation of clays with a low fractionation
475 factor and low temperature clays with a high fractionation factor (Li and West, 2014). Recognising that bulk solids
476 likely contain a fraction of exchangeable Li, another potential explanation could be that, compared to the continental
477 environment, the proportion of Li in exchangeable sites is greater in the marine environment, reducing $\Delta^7\text{Li}_{\text{solid-solution}}$.
478 However, this would be inconsistent with our results which suggest that the higher Na concentrations in seawater
479 would lead to a reduced fraction of exchangeable Li in bulk solids (Fig. 6).

480 When the $\delta^7\text{Li}$ of small rivers are considered in addition to the large rivers (compiled as a histogram in Fig. 10), the
481 riverine input value is $+16.5\text{‰}$ (defined by the mode of the density function, Fig. 10), significantly lower than the flux
482 weighted estimate from large rivers ($+23\text{‰}$, Huh et al., 1998). The difference between seawater and the river input
483 would then be 14.3‰ and if this difference was only caused by the precipitation of marine clays then this apparent
484 fractionation factor would be similar to the difference between rivers and sediments (17.3‰ , Fig. 10). These values
485 are similar to the average $\Delta^7\text{Li}_{\text{residue-solution}}$ value ($-16.6\pm 1.7\text{‰}$, Fig. 9). If the $\delta^7\text{Li}$ value for the riverine input to
486 seawater is closer to the modal value of smaller rivers rather than large rivers then it could imply a greater role of low

487 temperature reverse weathering processes on the sea-floor as being an important sink for Li (cf. Li and West, 2014).
488 Alternatively it could imply that regardless of the type of clay or initial solution conditions (marine vs continental),
489 the clays formed contain roughly a similar proportion of Li in the different bonding environments, supporting previous
490 suggestions that $\Delta^7\text{Li}_{\text{solid-solution}}$ is neither affected by the initial solution composition (Vigier et al., 2008) nor the type
491 of clay mineral forming (Dupuis et al., 2017; Pogge von Strandmann et al., 2017a).

492 These conflicting possibilities highlight the importance of understanding isotope fractionation processes at an
493 atomistic scale in order to decipher global processes, and reinforces the need for further experimental isotopic and
494 material characterisation of mineral phases. The potential implications discussed above should be treated with caution
495 not only because of the large uncertainty, particularly on the pseudo-hexagonal fractionation factor, but also because
496 of our limited understanding of field data and kinetics. For example, a fractionation factor ($\Delta^7\text{Li}_{\text{solid-solution}}$) of -
497 30‰ for the uptake of Li into secondary minerals during weathering was calculated by Bouchez et al. (2013) from
498 a steady-state mass-balance weathering model and the authors suggested that this could indicate kinetic processes
499 superimposed on equilibrium fractionation. In summary, additional experimental work is essential to provide a better
500 quantitative constraint on field data.

501 **5. Conclusions**

502 We have synthesised Mg-rich layer silicate minerals at temperatures relevant for Earth surface processes, charac-
503 terising the mineralogy, structure, bonding environment and Li isotope composition. ^7Li -NMR data reveals that the
504 composition of bulk clays is a mixture Li located in three bonding environments: octahedral, outer-sphere complexes
505 (interlayer and adsorbed) and pseudo-hexagonal. Following cation exchange with ammonium chloride to remove ex-
506 changeable Li (outer-sphere and pseudo-hexagonal), the measured $\Delta^7\text{Li}_{\text{residue-solution}}$ value was $-16.6 \pm 1.7\text{‰}$ at 20°C ,
507 which agrees well with the apparent fractionation factor obtained from a global compilation of field data (-17.1‰)
508 and is in agreement with previous experimental estimates for this temperature (-18.8‰ , Vigier et al., 2008). However,
509 ^7Li -NMR data shows that, although significantly reduced, some exchangeable Li remains in the residual solid. Once
510 the proportions of Li in the three bonding environments are accounted for, our calculated estimate of the fraction-
511 ation factor into octahedral sites is $-21.5 \pm 1.1\text{‰}$, slightly lower than a previous modelled value based on a density
512 functional theory (DFT) calculation (-17.2‰ , Dupuis et al., 2017). Our calculated fractionation factors for Li in-
513 corporation into outer-sphere (interlayer and adsorbed) and pseudo-hexagonal sites are $-0.2 \pm 1.9\text{‰}$ and $15.0 \pm 12.3\text{‰}$
514 respectively. However, caution should be exercised when applying these values to natural clay samples as exchange-
515 able and octahedral Li may be derived from different fluids if the isotopic composition of the fluid has evolved over
516 time.

517 Our data also suggest that controls on the magnitude of the $\Delta^7\text{Li}_{\text{residue-solution}}$ value, which are also relevant to
518 natural clays in field settings, relate to both pH, perhaps via a change in aqueous speciation or a kinetic rate control, and
519 to the composition of the final solid (in particular the Mg/Li ratio). Combining experimental approaches with detailed

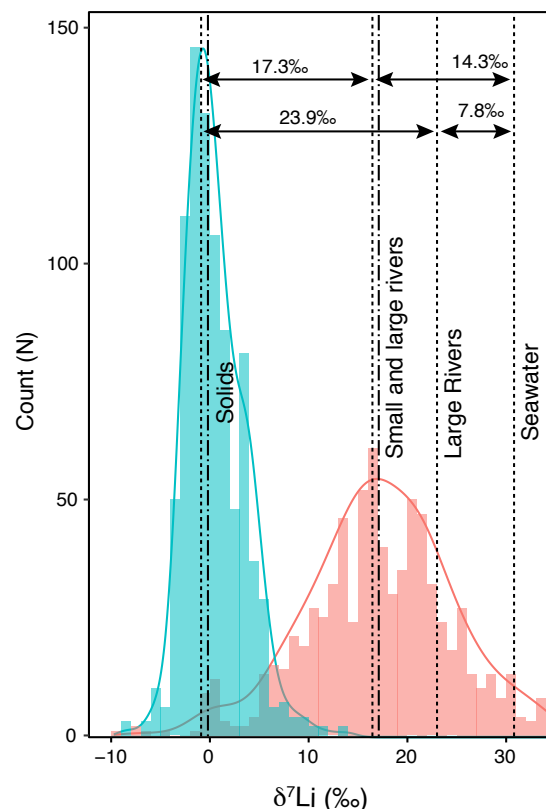


Figure 10: Histogram of a literature compilation of $\delta^7\text{Li}$ field data. Bin size is 1‰, and the density function for all the data is shown overlaid with a bandwidth of 1.5. Literature sources: Bagard et al. (2015); Clergue et al. (2015); Dellinger et al. (2014, 2015); Henchiri et al. (2016); Hindshaw et al. (2018); Huh et al. (1998, 2001); Kısakürek et al. (2004, 2005); Lemarchand et al. (2010); Liu et al. (2013, 2015); Manaka et al. (2017); Millot et al. (2010a,c, 2011, 2012); Murphy et al. (2019); Négrel et al. (2010); Négrel et al. (2012); Pogge von Strandmann et al. (2006, 2010, 2014, 2017b); Pogge von Strandmann and Henderson (2015); Qiu et al. (2011); Ryu et al. (2014); Sauzéat et al. (2015); Teng et al. (2004); Tsai et al. (2014); Vigier et al. (2009); Wang et al. (2015); Weynell et al. (2017); Wimpenny et al. (2010b); Xiao et al. (2011). The ‘Large Rivers’ line is from Huh et al. (1998) and the seawater line is from Rosner et al. (2007). Dashed lines indicate the mode of the density functions whilst the dash-dot lines indicate the median values. The calculated differences are based on the dashed lines.

520 solid characterisation (including ^7Li -NMR) and DFT calculations offers a promising pathway to elucidate the factors
 521 controlling Li isotope fractionation into clay minerals: an essential pre-requisite for a quantitative understanding of
 522 the global Li geochemical cycle.

523 6. Acknowledgments

524 This project was funded by NERC Standard Grant NE/M001865/1, NERC New Investigators Grants NE/K000705/1
 525 and NE/K000705/2, and a Marie Curie Intra-European Fellowship (PIEF-GA-2012-331501). We thank Daniel Daw-
 526 son (University of St. Andrews) for analysing pilot ^7Li -NMR data and Nathalie Vigier for providing the Li7-N and
 527 Li6-N standard solutions. Lynda Williams, two anonymous reviewers and associate editor Horst Marschall are thanked
 528 for their constructive reviews.

529 7. References

- 530 Bagard, M.-L., West, A. J., Newman, K., Basu, A. R., 2015. Lithium isotope fractionation in the Ganges-Brahmaputra floodplain and implications
531 for groundwater impact on seawater isotopic composition. *Earth Planet. Sci. Lett.* 432, 404–414.
- 532 Bigeleisen, J., 1965. Chemistry of isotopes. *Science* 147, 463–471.
- 533 Bogatko, S., Claeys, P., De Proft, F., Geerlings, P., 2013. Li⁺ speciation and the use of ⁷Li/⁶Li isotope ratios for ancient climate monitoring. *Chem.*
534 *Geol.* 357, 1–7.
- 535 Bohlin, M. S., Misra, S., Lloyd, N., Elderfield, H., Bickle, M. J., 2018. High-precision determination of lithium and magnesium isotopes utilising
536 single column separation and multi-collector inductively coupled plasma mass spectrometry. *Rapid Commun. Mass Spectrom.* 32, 93–104.
- 537 Bouchez, J., von Blanckenburg, F., Schuessler, J. A., 2013. Modeling novel stable isotope ratios in the weathering zone. *Am. J. Sci.* 313, 267–308.
- 538 Brigatti, M. F., Lugli, C., Poppi, L., Foord, E. E., Kile, D. E., 2000. Crystal chemical variations in Li- and Fe-rich micas from Pikes Peak batholith
539 (central Colorado). *Am. Mineral.* 85, 1275–1286.
- 540 Brindley, G. W., Bish, D. L., Wan, H.-M., 1977. The nature of kerolite, its relation to talc and stevensite. *Miner. Mag.* 41, 443–452.
- 541 Carignan, J., Cardinal, D., Eisenhauer, A., Galy, A., Rehkämper, M., Wombacher, F., Vigier, N., 2004. A reflection on Mg, Cd, Ca, Li and Si
542 isotopic measurements and related reference materials. *Geostand. Geoanal. Res.* 28, 139–148.
- 543 Carignan, J., Vigier, N., Millot, R., 2007. Three secondary reference materials for lithium isotope measurements: Li7-N, Li6-N and LiCl-N
544 solutions. *Geostand. Geoanal. Res.* 31, 7–12.
- 545 Carrado, K. A., Decarreau, A., Petit, S., Bergaya, F., Lagaly, G., 2006. Synthetic clay minerals and purification of natural clays. In: Bergaya,
546 F., Theng, B. K. G., Lagaly, G. (Eds.), *Handbook of Clay Science*. Vol. 1 of *Developments in Clay Science*. Elsevier, Amsterdam, Ch. 4, pp.
547 115–139.
- 548 Chan, L.-H., Edmond, J. M., Thompson, G., 1993. A lithium isotope study of hot springs and metabasalts from mid-ocean ridge hydrothermal
549 systems. *J. Geophys. Res.* 98, 9653–9659.
- 550 Chan, L.-H., Hein, J. R., 2007. Lithium contents and isotopic compositions of ferromanganese deposits from the global ocean. *Deep-Sea Res. II*
551 54, 1147–1162.
- 552 Clergue, C., Dellinger, M., Buss, H. L., Gaillardet, J., Benedetti, M. F., Dessert, C., 2015. Influence of atmospheric deposits and secondary minerals
553 on Li isotopes budget in a highly weathered catchment, Guadeloupe (Lesser Antilles). *Chem. Geol.* 414, 28–41.
- 554 Decarreau, A., Vigier, N., Pálková, H., Petit, S., Vieillard, P., Fontaine, C., 2012. Partitioning of lithium between smectite and solution: An
555 experimental approach. *Geochim. Cosmochim. Acta* 85, 314–325.
- 556 Dekov, V. M., Cuadros, J., Shanks, W. C., Koski, R. A., 2008. Deposition of talc – kerolite-smectite – smectite at seafloor hydrothermal vent fields:
557 Evidence from mineralogical, geochemical and oxygen isotope studies. *Chem. Geol.* 247, 171–194.
- 558 Dellinger, M., Gaillardet, J., Bouchez, J., Calmels, D., Galy, V., Hilton, R. G., Louvat, P., France-Lanord, C., 2014. Lithium isotopes in large rivers
559 reveal the cannibalistic nature of modern continental weathering and erosion. *Earth Planet. Sci. Lett.* 401, 359–372.
- 560 Dellinger, M., Gaillardet, J., Bouchez, J., Calmels, D., Louvat, P., Dosseto, A., Gorge, C., Alanoca, L., Maurice, L., 2015. Riverine Li isotope
561 fractionation in the Amazon River basin controlled by the weathering regimes. *Geochim. Cosmochim. Acta* 164, 71–93.
- 562 Dessert, C., Dupré, B., François, L. M., Schott, J., Gaillardet, J., Chakrapani, G., Bajpai, S., 2001. Erosion of Deccan Traps determined by river
563 geochemistry: impact on the global climate and the ⁸⁷Sr/⁸⁶Sr ratio of seawater. *Earth Planet. Sci. Lett.* 188, 459–474.
- 564 Dupuis, R., Benoit, M., Tuckerman, M. E., Méheut, M., 2017. Importance of a fully anharmonic treatment of equilibrium isotope fractionation
565 properties of dissolved ionic species as evidenced by Li⁺(aq). *Acc. Chem. Res.* 50, 1597–1605.
- 566 Eberl, D. D., Jones, B. F., Khoury, H. N., 1982. Mixed-layer kerolite/stevensite from the Amargosa Desert, Nevada. *Clay Clay Miner.* 30, 321–326.
- 567 Farmer, V. C., 1974. The layer silicates. In: *The Infrared Spectra of Minerals*. Vol. 4. Mineralogical Society, London, pp. 331–363.
- 568 Froelich, F., Misra, S., 2014. Was the late Paleocene-early Eocene hot because the Earth was flat? An ocean lithium isotope view of mountain
569 building, continental weathering, carbon dioxide, and Earth's Cenozoic climate. *Oceanography* 27, 36–49.
- 570 Gíslason, S. R., Arnórsson, S., Ármannsson, H., 1996. Chemical weathering of basalt in southwest Iceland: Effects of runoff, age of rocks and
571 vegetative/glacial cover. *Am. J. Sci.* 296, 837–907.

572 Greathouse, J., Sposito, G., 1998. Monte Carlo and molecular dynamics studies of interlayer structure in $\text{Li}(\text{H}_2\text{O})_3$ -smectites. *J. Phys. Chem. B*
573 102, 2406–2414.

574 Guggenheim, S., 2016. Introduction to Mg-rich clay minerals: Structure and composition. In: Pozo, M., Galán, E. (Eds.), *Magnesian clays:*
575 *Characterization, origins and applications*. No. 2 in AIPEA Educational Series. AIPEA, Bari, Italy, pp. 1–62.

576 Güven, N., Carney, L. L., 1979. The hydrothermal transformation of sepiolite to stevensite and the effect of added chlorides and hydroxides. *Clay*
577 *Clay Miner.* 27, 253–260.

578 Hall, J. M., Chan, L.-H., McDonough, W. F., Turekian, K. K., 2005. Determination of the lithium isotopic composition of planktic foraminifera
579 and its application as a paleo-seawater proxy. *Marine Geology* 217, 255–265.

580 Harder, H., 1972. The role of magnesium in the formation of smectite minerals. *Chem. Geol.* 10, 31–39.

581 Hathorne, E. C., James, R. H., 2006. Temporal record of lithium in seawater: A tracer for silicate weathering? *Earth Planet. Sci. Lett.* 246, 393–406.

582 Henchiri, S., Gaillardet, J., Dellinger, M., Bouchez, J., Spencer, R. G. M., 2016. Riverine dissolved lithium isotopic signatures in low-relief central
583 Africa and their link to weathering regimes. *Geophys. Res. Lett.* 43, 4391–4399.

584 Hindshaw, R. S., Aciego, S. M., Tipper, E. T., 2018. Li and U isotopes as a potential tool for monitoring active layer deepening in permafrost
585 dominated catchments. *Front. Earth Sci.* 6, 102.

586 Hindshaw, R. S., Rickli, J., Leuthold, J., Wadham, J., Bourdon, B., 2014. Identifying weathering sources and processes in an outlet glacier of the
587 Greenland Ice Sheet using Ca and Sr isotope ratios. *Geochim. Cosmochim. Acta* 145, 50–71.

588 Hochella, Jr, M. F., Banfield, J. F., 1995. Chemical weathering of silicates in nature: A microscopic perspective with theoretical considerations.
589 In: White, A. F., Brantley, S. L., Ribbe, P. H. (Eds.), *Chemical weathering rates of silicate minerals*. Vol. 31 of *Reviews in Mineralogy and*
590 *Geochemistry*. Mineralogical Society of America, Washington, DC, Ch. 8, pp. 353–406.

591 Huh, Y., Chan, L.-H., Edmond, J. M., 2001. Lithium isotopes as a probe of weathering processes: Orinoco river. *Earth Planet. Sci. Lett.* 194,
592 189–199.

593 Huh, Y., Chan, L.-H., Zhang, L., Edmond, J. M., 1998. Lithium and its isotopes in major world rivers: Implications for weathering and the oceanic
594 budget. *Geochim. Cosmochim. Acta* 62, 2039–2051.

595 Jones, B. F., 1986. Clay mineral diagenesis in lacustrine sediments. In: Mumpton, F. A. (Ed.), *Studies in diagenesis*. Vol. 1578. U.S. Geological
596 Survey, Washington, DC, pp. 291–300.

597 Kalo, H., Milius, W., Breu, J., 2012. Single crystal structure refinement of one- and two-layer hydrates of sodium fluorohectorite. *RSC Adv.* 2,
598 8452–8459.

599 Kısakürek, B., James, R. H., Harris, N. B. W., 2005. Li and $\delta^7\text{Li}$ in Himalayan rivers: Proxies for silicate weathering? *Earth Planet. Sci. Lett.* 237,
600 387–401.

601 Kısakürek, B., Widdowson, M., James, R. H., 2004. Behaviour of Li isotopes during continental weathering: the Bidar laterite profile, India. *Chem.*
602 *Geol.* 212, 27–44.

603 Laurora, A., Brigatti, M. F., Malferrari, D., Galli, E., Rossi, A., 2011. The crystal chemistry of lizardite-1T from northern Apennines ophiolites
604 near Modena, Italy. *The Canad. Mineral.* 49, 1045–1054.

605 Lechler, M., Pogge von Strandmann, P. A. E., Jenkyns, H. C., Prosser, G., Parente, M., 2015. Lithium-isotope evidence for enhanced silicate
606 weathering during OAE 1a (Early Aptian Selli event). *Earth Planet. Sci. Lett.* 432, 210–222.

607 Lemarchand, E., Chabaux, F., Vigier, N., Millot, R., Pierret, M.-C., 2010. Lithium isotope systematics in a forested granitic catchment (Strengbach,
608 Vosges Mountains, France). *Geochim. Cosmochim. Acta* 74, 4612–4628.

609 Li, G., West, A. J., 2014. Evolution of Cenozoic seawater lithium isotopes: Coupling of global denudation regime and shifting seawater sinks.
610 *Earth Planet. Sci. Lett.* 401, 284–293.

611 Li, W., Beard, B. L., Li, C., Johnson, C. M., 2014. Magnesium isotope fractionation between brucite $[\text{Mg}(\text{OH})_2]$ and Mg aqueous species:
612 Implications for silicate weathering and biogeochemical processes. *Earth Planet. Sci. Lett.* 394, 82–93.

613 Liu, X.-M., Rudnick, R. L., McDonough, W. F., Cummings, M. L., 2013. Influence of chemical weathering on the composition of the continental
614 crust: Insights from Li and Nd isotopes in bauxite profiles developed on Columbia River Basalts. *Geochim. Cosmochim. Acta* 115, 73–91.

- 615 Liu, X.-M., Wanner, C., Rudnick, R. L., McDonough, W. F., 2015. Processes controlling $\delta^7\text{Li}$ in rivers illuminated by study of streams and
616 groundwaters draining basalts. *Earth Planet. Sci. Lett.* 409, 212–224.
- 617 Lynton, S. J., Walker, R. J., Candela, P. A., 2005. Lithium isotopes in the system Qz-Ms-fluid: An experimental study. *Geochim. Cosmochim. Acta*
618 69, 3337–3347.
- 619 Mähler, J., Persson, I., 2012. A study of the hydration of the alkali metal ions in aqueous solution. *Inorg. Chem.* 51, 425–438.
- 620 Manaka, T., Araoka, D., Yoshimura, T., Hossain, H. M. Z., Nishio, Y., Suzuki, A., Kawahata, H., 2017. Downstream and seasonal changes of
621 lithium isotope ratios in the Ganges-Brahmaputra river system. *Geochem. Geophys. Geosys.* 18, 3003–3015.
- 622 Martin de Vidales, J. L., Pozo, M., Alia, J. M., Garcia-Navarro, F., Rull, F., 1991. Kerolite-stevensite mixed-layers from the Madrid Basin, Central
623 Spain. *Clay Miner.* 26, 329–342.
- 624 Meunier, A., 2005. *Clays*. Springer-Verlag, Berlin.
- 625 Michalski, J. R., Cuadros, J., Bishop, J. L., Dyar, M. D., Dekov, V., Fiore, S., 2015. Constraints on the crystal-chemistry of Fe/Mg-rich smectitic
626 clays on Mars and links to global alteration trends. *Earth Planet. Sci. Lett.* 427, 215–225.
- 627 Millot, R., Guerrot, C., Innocent, C., Négrel, P., Sanjuan, B., 2011. Chemical, multi-isotopic (Li–B–Sr–U–H–O) and thermal characterization of
628 Triassic formation waters from the Paris Basin. *Chem. Geol.* 283, 226–241.
- 629 Millot, R., Hegan, A., Négrel, P., 2012. Geothermal waters from the Taupo Volcanic Zone, New Zealand: Li, B and Sr isotopes characterization.
630 *Appl. Geochem.* 27, 677–688.
- 631 Millot, R., Petelet-Giraud, E., Guerrot, C., Négrel, P., 2010a. Multi-isotopic composition ($\delta^7\text{Li}$ – $\delta^{11}\text{B}$ – δD – $\delta^{18}\text{O}$) of rainwaters in France: Origin
632 and spatio-temporal characterization. *Appl. Geochem.* 25, 1510–1524.
- 633 Millot, R., Scaillet, B., Sanjuan, B., 2010b. Lithium isotopes in island arc geothermal systems: Guadeloupe, Martinique (French West Indies) and
634 experimental approach. *Geochim. Cosmochim. Acta* 74, 1852–1871.
- 635 Millot, R., Vigier, N., Gaillardet, J., 2010c. Behaviour of lithium and its isotopes during weathering in the Mackenzie Basin, Canada. *Geochim.*
636 *Cosmochim. Acta* 74, 3897–3912.
- 637 Misra, S., Froelich, P. N., 2012. Lithium isotope history of Cenozoic seawater: changes in silicate weathering and reverse weathering. *Science* 335,
638 818–823.
- 639 Murphy, M. J., Porcelli, D., Pogge von Strandmann, P. A. E., Hirst, C. A., Kutscher, L., Katchinoff, J. A., Mörth, C.-M., Maximov, T., Ander-
640 sson, P. S., 2019. Tracing silicate weathering processes in the permafrost-dominated Lena River watershed using lithium isotopes. *Geochim.*
641 *Cosmochim. Acta* 245, 154–171.
- 642 Négrel, P., Millot, R., Brenot, A., Bertin, C., 2010. Lithium isotopes as tracers of groundwater circulation in a peat land. *Chem. Geol.* 276, 119–127.
- 643 Négrel, P., Millot, R., Guerrot, C., Petelet-Giraud, E., Brenot, A., Malcuit, E., 2012. Heterogeneities and interconnections in groundwaters: Coupled
644 B, Li and stable-isotope variations in a large aquifer system (Eocene Sand aquifer, Southwestern France). *Chem. Geol.* 296–297, 83–95.
- 645 Nielsen, L. C., DePaolo, D. J., De Yoreo, J. J., 2012. Self-consistent ion-by-ion growth model for kinetic isotopic fractionation during calcite
646 precipitation. *Geochim. Cosmochim. Acta* 86, 166–181.
- 647 Perdikatsis, B., Burzlaff, H., 1981. Strukturverfeinerung am Talk $\text{Mg}_3[(\text{OH})_2\text{Si}_4\text{O}_{10}]$. *Z. Kristall.* 156, 177–186.
- 648 Phan, T. T., Capo, R. C., Stewart, B. W., Macpherson, G. L., Rowan, E. L., Hammack, R. W., 2016. Factors controlling Li concentration and
649 isotopic composition in formation waters and host rocks of Marcellus Shale, Appalachian Basin. *Chem. Geol.* 420, 162–179.
- 650 Pistiner, J. S., Henderson, G. M., 2003. Lithium-isotope fractionation during continental weathering processes. *Earth Planet. Sci. Lett.* 214, 327–
651 339.
- 652 Pogge von Strandmann, P. A. E., Burton, K. W., James, R. H., van Calsteren, P., Gíslason, S. R., 2010. Assessing the role of climate on uranium
653 and lithium isotope behaviour in rivers draining a basaltic terrain. *Chem. Geol.* 270, 227–239.
- 654 Pogge von Strandmann, P. A. E., Burton, K. W., James, R. H., van Calsteren, P., Gíslason, S. R., Mokadem, F., 2006. Riverine behaviour of uranium
655 and lithium isotopes in an actively glaciated basaltic terrain. *Earth Planet. Sci. Lett.* 251, 134–147.
- 656 Pogge von Strandmann, P. A. E., Desrochers, A., Murphy, M. J., Finlay, A. J., Selby, D., Lenton, T. M., 2017a. Global climate stabilisation by
657 chemical weathering during the Hirnantian glaciation. *Geochem. Persp. Lett.* 3, 230–237.

- 658 Pogge von Strandmann, P. A. E., Frings, P. J., Murphy, M. J., 2017b. Lithium isotope behaviour during weathering in the Ganges Alluvial Plain.
659 *Geochim. Cosmochim. Acta* 198, 17–31.
- 660 Pogge von Strandmann, P. A. E., Henderson, G. M., 2015. The Li isotope response to mountain uplift. *Geology* 43, 67–70.
- 661 Pogge von Strandmann, P. A. E., Jenkyns, H. C., Woodfine, R. G., 2013. Lithium isotope evidence for enhanced weathering during Ocean Anoxic
662 Event 2. *Nat. Geosci.* 6, 668–672.
- 663 Pogge von Strandmann, P. A. E., Porcelli, D., James, R. H., van Calsteren, P., Schaefer, B., Cartwright, I., Reynolds, B. C., Burton, K. W., 2014.
664 Chemical weathering processes in the Great Artesian Basin: Evidence from lithium and silicon isotopes. *Earth Planet. Sci. Lett.* 406, 24–36.
- 665 Pokrovsky, O. S., Schott, J., Kudryavtzev, D. I., Dupré, B., 2005. Basalt weathering in Central Siberia under permafrost conditions. *Geochim.*
666 *Cosmochim. Acta* 69, 5659–5680.
- 667 Qiu, L., Rudnick, R. L., McDonough, W. F., Bea, F., 2011. The behavior of lithium in amphibolite- to granulite-facies rocks of the Ivrea–Verbano
668 Zone, NW Italy. *Chem. Geol.* 289, 76–85.
- 669 Reimers, C. E., Ruttenger, K. C., Canfield, D. E., Christiansen, M. B., Martin, J. B., 1996. Porewater pH and authigenic phases formed in the
670 uppermost sediments of the Santa Barbara Basin. *Geochim. Cosmochim. Acta* 60, 4037–4057.
- 671 Roberts, J., Kaczmarek, K., Langer, G., Skinner, L. C., Bijma, J., Bradbury, H., Turchyn, A. V., Lamy, F., Misra, S., 2018. Lithium isotopic
672 composition of benthic foraminifera: A new proxy for paleo-pH reconstruction. *Geochim. Cosmochim. Acta* 236, 336–350.
- 673 Rosner, M., Ball, L., Peucker-Ehrenbrink, B., Blusztajn, J., Bach, W., Erzinger, J., 2007. A simplified, accurate and fast method for lithium isotope
674 analysis of rocks and fluids, and $\delta^7\text{Li}$ values of seawater and rock reference materials. *Geostand. Geoanal. Res.* 31, 77–88.
- 675 Rudolph, W., Brooker, M. H., Pye, C. C., 1995. Hydration of lithium ion in aqueous solution. *J. Phys. Chem.* 99, 3793–3797.
- 676 Russell, J. D., Fraser, A. R., 1994. Infrared methods. In: Wilson, M. J. (Ed.), *Clay mineralogy: Spectroscopic and chemical determinative methods.*
677 Vol. 23. Chapman & Hall, London, pp. 11–67.
- 678 Ryu, J.-S., Vigier, N., Lee, S.-W., Lee, K.-S., Chadwick, O. A., 2014. Variation of lithium isotope geochemistry during basalt weathering and
679 secondary mineral transformations in Hawaii. *Geochim. Cosmochim. Acta* 145, 103–115.
- 680 Sauzéat, L., Rudnick, R. L., Chauvel, C., Garçon, M., Tang, M., 2015. New perspectives on the Li isotopic composition of the upper continental
681 crust and its weathering signature. *Earth Planet. Sci. Lett.* 428, 181–192.
- 682 Schauble, E. A., 2004. Applying stable isotope fractionation theory to new systems. In: Johnson, C. M., Beard, B. L., Albarède, F. (Eds.), *Geochem-*
683 *istry of Non-traditional Stable Isotopes. Vol. 55 of Reviews in Mineralogy & Geochemistry.* Mineralogical Society of America, Washington,
684 DC, pp. 65–111.
- 685 Studel, A., Friedrich, F., Schuhmann, R., Ruf, F., Sohling, U., Emmerich, K., 2017. Characterization of a fine-grained interstratification of
686 turbostatic talc and saponite. *Minerals* 7, 5.
- 687 Stoessell, R. K., Hay, R. L., 1978. The geochemical origin of sepiolite and kerolite at Amboseli, Kenya. *Contrib. Mineral. Petrol.* 65, 255–267.
- 688 Stumm, W., Morgan, J. J., 1996. *Aquatic Chemistry: chemical equilibria and rates in natural waters*, 3rd Edition. Wiley, New York, NY.
- 689 Tang, J., Dietzel, M., Böhm, F., Köhler, S. J., Eisenhauer, A., 2008. $\text{Sr}^{2+}/\text{Ca}^{2+}$ and $^{44}\text{Ca}/^{40}\text{Ca}$ fractionation during inorganic calcite formation: II.
690 Ca isotopes. *Geochim. Cosmochim. Acta* 72, 3733–3745.
- 691 Tardy, Y., Krempp, G., Trauth, N., 1972. Le lithium dans les minéraux argileux des sédiments et des sols. *Geochim. Cosmochim. Acta* 36, 397–412.
- 692 Taylor, T. I., Urey, H. C., 1938. Fractionation of the lithium and potassium isotopes by chemical exchange with zeolites. *J. Chem. Phys.* 6, 429–438.
- 693 Teng, F.-Z., McDonough, W. F., Rudnick, R. L., Dalpé, C., Tomascak, P. B., Chappell, B. W., Gao, S., 2004. Lithium isotopic composition and
694 concentration of the upper continental crust. *Geochim. Cosmochim. Acta* 68, 4167–4178.
- 695 Theng, B. K. G., Hayashi, S., Soma, M., Seyama, H., 1997. Nuclear magnetic resonance and X-ray photoelectron spectroscopic investigation of
696 lithium migration in montmorillonite. *Clay. Clay Miner.* 45, 718–723.
- 697 Tipper, E. T., Lemarchand, E., Hindshaw, R. S., Reynolds, B. C., Bourdon, B., 2012. Seasonal sensitivity of weathering processes: Hints from
698 magnesium isotopes in a glacial stream. *Chem. Geol.* 312–313, 80–92.
- 699 Tomascak, P. B., 2004. Developments in the understanding and application of lithium isotopes in the Earth and planetary sciences. In: Johnson,
700 C. M., Beard, B. L., Albarède, F. (Eds.), *Geochemistry of Non-traditional Stable Isotopes. Vol. 55 of Reviews in Mineralogy & Geochemistry.*

701 Mineralogical Society of America, Washington, DC, pp. 153–195.

702 Tosca, N., Masterson, A. L., 2014. Chemical controls on incipient Mg-silicate crystallization at 25°C: Implications for early and late diagenesis.
703 *Clay Miner.* 49, 195–224.

704 Tosca, N. J., 2016. Geochemical pathways to Mg-silicate formation. In: Pozo, M., Gálan, E. (Eds.), *Magnesian clays: Characterization, origins and
705 applications*. No. 2 in AIPEA Educational Series. AIPEA, Bari, Italy, pp. 283–330.

706 Tsai, P.-H., You, C.-F., Huang, K.-F., Chung, C.-H., Sun, Y.-B., 2014. Lithium distribution and isotopic fractionation during chemical weathering
707 and soil formation in a loess profile. *J. Asian Earth Sci.* 87, 1–10.

708 Tutolo, B. M., Tosca, N. J., 2018. Experimental examination of the Mg-silicate-carbonate system at ambient temperature: Implications for alkaline
709 chemical sedimentation and lacustrine carbonate formation. *Geochim. Cosmochim. Acta* 225, 80–101.

710 Verney-Carron, A., Vigier, N., Millot, R., 2011. Experimental determination of the role of diffusion on Li isotope fractionation during basaltic glass
711 weathering. *Geochim. Cosmochim. Acta* 75, 3452–3468.

712 Vigier, N., Decarreau, A., Millot, R., Carignan, J., Petit, S., France-Lanord, C., 2008. Quantifying Li isotope fractionation during smectite formation
713 and implications for the Li cycle. *Geochim. Cosmochim. Acta* 72, 780–792.

714 Vigier, N., Gislason, S. R., Burton, K. W., Millot, R., Mokadem, F., 2009. The relationship between riverine lithium isotope composition and
715 silicate weathering rates in Iceland. *Earth Planet. Sci. Lett.* 287, 434–441.

716 Vigier, N., Godd eris, Y., 2015. A new approach for modeling Cenozoic ocean lithium isotope paleo-variations: the key role of climate. *Clim. Past.*
717 11, 635–645.

718 Vogels, R. J. M. J., Klopogge, J. T., Geus, J. W., 2005. Synthesis and characterization of saponite clays. *Am. Mineral.* 90, 931–944.

719 Von Damm, K. L., Edmond, J. M., 1984. Reverse weathering in the closed-basin lakes of the Ethiopian Rift. *Am. J. Sci.* 284, 835–862.

720 Wang, Q.-L., Chetelat, B., Zhao, Z.-Q., Ding, H., Li, S.-L., Wang, B.-L., Li, J., Liu, X.-L., 2015. Behavior of lithium isotopes in the Changjiang
721 River system: Sources effects and response to weathering and erosion. *Geochim. Cosmochim. Acta* 151, 117–132.

722 Wang, W., Qin, T., Zhou, C., Huang, S., Wu, Z., Huang, F., 2017a. Concentration effect on equilibrium fractionation of Mg-Ca isotopes in carbonate
723 minerals: Insights from first-principles calculations. *Geochim. Cosmochim. Acta* 208, 185–197.

724 Wang, W., Zhou, C., Qin, T., Kang, J.-T., Huang, S., Wu, Z., Huang, F., 2017b. Effect of Ca content on equilibrium Ca isotope fractionation between
725 orthopyroxene and clinopyroxene. *Geochim. Cosmochim. Acta* 219, 44–56.

726 Wanner, C., Sonnenthal, E. L., Liu, X.-M., 2014. Seawater $\delta^7\text{Li}$: A direct proxy for global CO₂ consumption by continental silicate weathering?
727 *Chem. Geol.* 381, 154–167.

728 Weynell, M., Wiechert, U., Schuessler, J. A., 2017. Lithium isotopes and implications on chemical weathering in the catchment of Lake Donggi
729 Cona, northeastern Tibetan Plateau. *Geochim. Cosmochim. Acta* 213, 155–177.

730 Wilkins, R. W. T., Ito, J., 1967. Infrared spectra of some synthetic talcs. *Am. Mineral.* 52, 1649–1661.

731 Williams, L. B., Hervig, R. L., 2005. Lithium and boron isotopes in illite-smectite: the importance of crystal size. *Geochim. Cosmochim. Acta* 69,
732 5705–5716.

733 Wimpenny, J., Colla, C. A., Yu, P., Yin, Q.-Z., Rustad, J. R., Casey, W. H., 2015. Lithium isotope fractionation during uptake by gibbsite. *Geochim.*
734 *Cosmochim. Acta* 168, 133–150.

735 Wimpenny, J., Gislason, S. R., James, R. H., Gannoun, A., Pogge von Strandmann, P. A. E., Burton, K. W., 2010a. The behaviour of Li and Mg
736 isotopes during primary phase dissolution and secondary mineral formation in basalt. *Geochim. Cosmochim. Acta* 74, 5259–5279.

737 Wimpenny, J., James, R. H., Burton, K. W., Gannoun, A., Mokadem, F., Gislason, S. R., 2010b. Glacial effects on weathering processes: New
738 insights from the elemental and lithium isotopic composition of West Greenland rivers. *Earth Planet. Sci. Lett.* 290, 427–437.

739 Wunder, B., Deschamps, F., Watenphul, A., Guillot, S., Meixner, A., Romer, R. L., Wirth, R., 2010. The effect of chrysotile nanotubes on the
740 serpentine-fluid Li-isotopic fractionation. *Contrib. Mineral. Petrol.* 159, 781–790.

741 Wunder, B., Meixner, A., Romer, R. L., Feenstra, A., Schettler, G., Heinrich, W., 2007. Lithium isotope fractionation between Li-bearing staurolite,
742 Li-mica and aqueous fluids: An experimental study. *Chem. Geol.* 238, 277–290.

743 Wunder, B., Meixner, A., Romer, R. L., Heinrich, W., 2006. Temperature-dependent isotopic fractionation of lithium between clinopyroxene and

- 744 high-pressure hydrous fluids. *Contrib. Mineral. Petrol.* 151, 112–120.
- 745 Wungu, T. D. K., Diño, W. A., Dipojono, H. K., Kasai, H., 2011. Effect of lithium absorption at tetrahedral site and isomorphic substitution on
746 montmorillonite properties: A density functional theory study. *Jap. J. Appl. Phys.* 50, 055701.
- 747 Xiao, Y., Hoefs, J., Hou, Z., Simon, K., Zhang, Z., 2011. Fluid/rock interaction and mass transfer in continental subduction zones: constraints from
748 trace elements and isotopes (Li, B, O, Sr, Nd, Pb) in UHP rocks from the Chinese Continental Scientific Drilling Program, Sulu, East China.
749 *Contrib. Mineral. Petrol.* 162, 797–819.
- 750 Xu, Z., Stebbins, J. F., 1995. ⁶Li nuclear magnetic resonance chemical shifts, coordination number and relaxation in crystalline and glassy silicates.
751 *Solid State Nucl. Magn. Reson.* 5, 103–112.
- 752 Yamaji, K., Makita, Y., Watanabe, H., Sonoda, A., Kanoh, H., Hirotsu, T., Ooi, K., 2001. Theoretical estimation of lithium isotopic reduced
753 partition function ratio for lithium ions in aqueous solution. *J. Phys. Chem. A* 105, 602–613.
- 754 Zhang, L., Chan, L.-H., Gieskes, J. M., 1998. Lithium isotope geochemistry of pore waters from Ocean Drilling Program Sites 918 and 919,
755 Irminger Basin. *Geochim. Cosmochim. Acta* 62, 2437–2450.

756 **8. Supplementary Material**

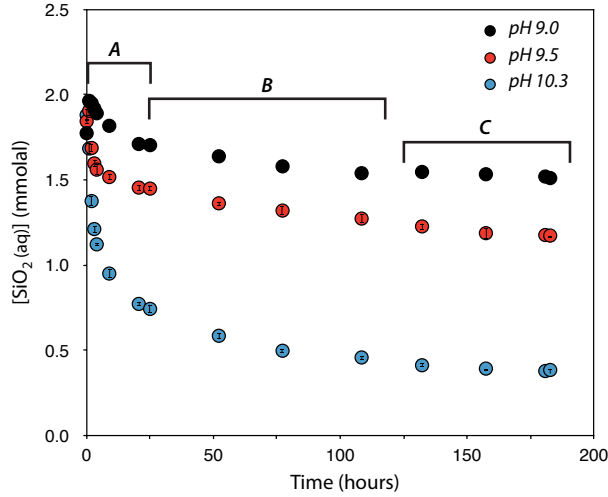


Figure S1: Evolution of SiO₂ concentrations in solution with time for additional experiments conducted with identical initial solution compositions to the '240417' series of experiments. Three phases are illustrated. **A** is a nucleation-dominated phase which lasts several hours and is characterised by rapid removal of constituents from solution. **B** is a growth-dominated phase which lasts several days and is characterised by slower removal of constituents from solution, reflecting growth of newly precipitated nuclei. **C** is steady-state behaviour where the element concentrations no longer change within experimental error.

757 **8.1. Monte Carlo minimum misfit method**

758 The “total” $\delta^7\text{Li}$ of each sample, be it a bulk, residue or exchange solution sample is given by the mass weighted
 759 sum of $\delta^7\text{Li}$ associated with each bonding environment (octahedral, pseudo-hexagonal and outer-sphere) as identi-
 760 fied by ^7Li -NMR. Given the difference between the exchange solution data and the residue data there are at least
 761 two different $\delta^7\text{Li}$ values corresponding to individual bonding environments. Since the mass fraction of Li in each
 762 environment is constrained by ^7Li -NMR, it is possible to write down a mass balance equation for each sample:

$$\delta^7\text{Li}_{total} = f_{oct} \cdot \delta^7\text{Li}_{oct} + f_{ph} \cdot \delta^7\text{Li}_{ph} + f_{os} \cdot \delta^7\text{Li}_{os} \quad (1)$$

763 where the subscripts *oct*, *ph* and *os* refer to the octahedral, pseudo-hexagonal and outer-sphere environments respec-
 764 tively and *f* is the mass fraction in each environment.

765 Since $\delta^7\text{Li}_{total}$ and the mass fractions were determined for six samples (bulk, residue and exchange) and there are
 766 three unknowns in this equation, $\delta^7\text{Li}_{oct}$, $\delta^7\text{Li}_{ph}$ and $\delta^7\text{Li}_{os}$, the problem is over determined and it is possible to solve
 767 for each of the unknowns. This was done using a misfit approach where the misfit is a measure of the difference
 768 between the measured and calculated $\delta^7\text{Li}$ of the sample:

$$misfit = \sqrt{\sum_k (\delta^7\text{Li}_{total-measured} - f_{oct} \cdot \delta^7\text{Li}_{oct} - f_{ph} \cdot \delta^7\text{Li}_{ph} - f_{os} \cdot \delta^7\text{Li}_{os})^2} \quad (2)$$

769 where k is the number of samples (six), the fractions are determined from ^7Li -NMR spectra and $\delta^7\text{Li}_{oct}$, $\delta^7\text{Li}_{ph}$, $\delta^7\text{Li}_{os}$
770 are variables, allowed to vary over a plausible range of parameter space.

771 The minimum misfit was found as a unique minimum corresponding to optimum (or best fit) values of $\delta^7\text{Li}_{oct}$,
772 $\delta^7\text{Li}_{ph}$ and $\delta^7\text{Li}_{os}$. The uncertainty on these values was determined by using a Monte-Carlo approach, creating 10000
773 synthetic data sets using a gaussian distribution of errors based on the uncertainties associated with apportioning the
774 mass fractions in each crystallographic site and the measurement uncertainties of $\delta^7\text{Li}$ on the Neptune.

775 The minimisation has been represented visually by plotting the mean of the misfit generated by the Monte-Carlo
776 simulation over an appropriate range of parameter space (Fig. S2). The misfit is represented by the colour gradient
777 and the contours. Since there are three variables, a 2D representation has been given by plotting a slice in the $\delta^7\text{Li}_{oct}$,
778 $\delta^7\text{Li}_{ph}$ and $\delta^7\text{Li}_{os}$ parameter space at the best fit value in the third dimension. Note that whilst a single solution has
779 been determined, the optimum $\delta^7\text{Li}_{ph}$ value shows a broad dependency in particular on $\delta^7\text{Li}_{os}$ value which is why the
780 uncertainty is large, whereas the $\delta^7\text{Li}_{oct}$ and $\delta^7\text{Li}_{os}$ values are relatively well constrained.

781 Finally, a visual check on the quality of these calculated fractionation factors is given by plotting the modelled vs
782 the measured $\delta^7\text{Li}_{total}$ (Fig. 8a). The data scatter very closely around the 1:1 line indicating the robustness of the fit.

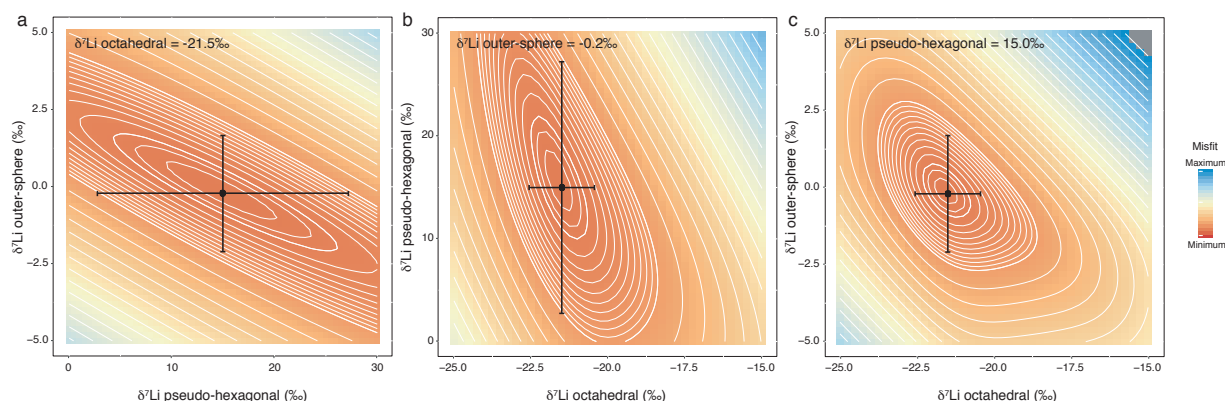


Figure S2: Contour maps indicating the misfit between measured $\delta^7\text{Li}$ values and calculated values when varying two of the fractionation factors. The third fractionation factor is held constant at its optimum value for the purposes of plotting. The black points indicate the optimum (smallest misfit) values for the two fractionation factors plotted and 1 sigma error bars are also shown.