1 2	
3	
4	
5	WEATHERING INTENSITY AND LITHIUM ISOTOPES:
6	A REACTIVE TRANSPORT PERSPECTIVE
/ 0	Matthew I Winnick!* Jannifor I Dryhan? Vata Mahar?
0 9	Matthew J. Winnick , Jennifer L. Drunan , Kate Maner
10	<sup>1</sup> Department of Geosciences
11	University of Massachusetts Amherst
12	Amherst, MA 01003, USA
13	
14	<sup>2</sup> Department of Geology
15	University of Illinois Urbana Champaign,
16	Urbana, IL
17	
18	<sup>3</sup> Department of Earth System Science
19	Stanford University
20	Stanford, CA 94305, USA
21	
22	
23	* Corresponding Author
25	mwinnick@umass.edu
26	627 N Pleasant St.
27	Morrill II Rm 233, Geosciences
28	Amherst, MA 01003, USA
29	
30	
31	
32	This manuscript has been submitted for publication in the American Journal of Science. This
33	version has undergone one round of peer review; however, this manuscript has not yet been
34	accepted for publication. Future versions of this manuscript may have altered content
35	addressing subsequent revisions in response to peer review. If accepted the final version of this
36	manuscript will be available via the DOI link on the right-hand side of this webpage. Please

37 contact the authors with any feedback.

38 ABSTRACT. Lithium isotopes have emerged as a powerful tool to probe the response of 39 global weathering to changes in climate. Due to the preferential incorporation of <sup>6</sup>Li into 40 clay minerals during chemical weathering, the isotope ratio  $\delta^7$ Li may be used to interrogate 41 the balance of primary mineral dissolution and clay precipitation. This balance has been 42 linked to relative rates of chemical and physical denudation, such that dissolved  $\delta^7 Li$ 43  $(\delta^7 \text{Lid}_{\text{iss}})$  is highest at moderate weathering intensities when chemical and physical 44 denudation are comparable. However, we argue that current theory linking  $\delta^7$ Li to 45 weathering regimes through fluid travel times are unable to explain observations of low 46  $\delta^7$ Li and high Li concentrations in rapidly eroding settings. In this study, we re-examine 47 the relationships between  $\delta^7$ Li, Li concentration, and weathering regime by incorporating 48 Li isotopes into simulations of weathering profiles using a reactive transport model 49 (CrunchFlow) that includes advective fluxes of regolith to simulate variable erosion rates in 50 response to uplift. In these simulations, fractionation is implemented through a kinetic 51 fractionation factor during clay precipitation, which allows the  $\delta^7$ Li of dissolved and 52 suspended loads in the model to vary as a function of Li/Al ratios in primary and 53 secondary minerals. When the model is run over a range of infiltration and erosion rates, simulations reproduce observed global patterns of  $\delta^7 \text{Li}_{\text{diss}}$  and suspended load  $\delta^7 \text{Li}$  as a 54 55 function of weathering intensity, controlled primarily by water travel times and mineral 56 residence times in weathered bedrock. We find that reduced water travel times at low 57 weathering intensity, however, are inconsistent with observations of high Li concentrations. 58 As an alternative, we demonstrate how the rapid weathering of soluble, Li-rich minerals 59 such as chlorite under low weathering intensities may resolve this apparent discrepancy 60 between data and theory. We also suggest that observed patterns are consistent with 61 geothermal Li sources under low weathering intensities. This work offers a foundation guiding future studies in testing potential mechanisms underlying global riverine  $\delta^7$ Lidiss. 62 63

- 64
- 65

#### **INTRODUCTION**

66 Chemical weathering, comprising the dissolution of carbonate and silicate minerals along 67 with the oxidation of sulfides and petrogenic carbon, is thought to be a primary regulator of 68 atmospheric composition on geologic timescales. The connections between chemical weathering

and climate are therefore central to our understanding of the evolution of global biogeochemical

- 70 cycles through Earth history. Over the past two decades, lithium isotopes have emerged as a
- 71 powerful tool for characterizing the relationships between silicate weathering and past climates.
- 72 Proxy records of oceanic  $\delta^7$ Li have been used, for example, to characterize connections between
- 73 weathering and Cenozoic cooling (Misra and Froelich, 2012; Li and West, 2014; Caves-
- Rugenstien and others, 2019; Pogge von Strandmann and others, 2020), along with the response
- of weathering to major carbon cycle perturbations including Ocean Anoxic Events (OAE's) 1
- and 2 (Pogge von Strandmann and others, 2013; Lechler and others, 2015), the Permian-Triassic
- boundary (Sun and others, 2018), and global glaciation events (Pogge von Strandmann and
- others, 2017). Interpretations of these proxy records rely on modern observations of lithium
- isotopes in weathering environments; however, even present-day connections between  $\delta^7$ Li and weathering dynamics have proven highly complex.

81 The use of lithium isotopes to probe weathering dynamics is enabled by the fact that the 82 lighter isotope, <sup>6</sup>Li, is preferentially incorporated into secondary minerals relative to the heavy 83 isotope, <sup>7</sup>Li, during the chemical weathering of primary minerals in bedrock. Here,  $\delta^7$ Li is 84 defined in standard delta-notation as,

85

86 
$$\delta^{7}Li = \left(\frac{{}^{7}Li/{}^{6}Li_{sample}}{{}^{7}Li/{}^{6}Li_{L-SVEC}} - 1\right) * 1000\%$$
(1),

87

88 where (<sup>7</sup>Li/<sup>6</sup>Li)<sub>L-SVEC</sub> is the isotope ratio of the L-SVEC standard, 12.2998 (Flesch and others,
89 1973).

90 Isotope fractionation during secondary mineral precipitation represents a combination of 91 effects relating to the incorporation of Li into mineral structural sites (Williams and Hervig, 92 2005; Vigier and others, 2008; Wimpenny and others, 2010) and inner-sphere sorption (Zhang 93 and others, 1998; Anghel and others, 2002; Pistiner and Henderson, 2003; Hindshaw and others, 94 2019) as potentially modified by OH group abundance (Millot and Girard, 2007). Secondary 95 mineral phase  $\delta^7$ Li values are thus offset from primary minerals by -5% to -30%, reflecting this 96 preferential uptake of <sup>6</sup>Li (Vigier and others, 2008; Chan and others, 1992; Williams and Hervig, 97 2005). As a result, global riverine  $\delta^7$ Lidiss values (-2 to >30‰) are significantly enriched relative 98 to average upper continental crust materials (-2-6‰) (Tomascak, 2004; Pogge von Strandmann 99 and others, 2020).

100 When solutes are enriched in <sup>7</sup>Li relative to bedrock, the dissolved  $\delta^7 \text{Li}(\delta^7 \text{Li}_{\text{diss}})$  reflects 101 the balance between Li release from primary mineral dissolution and Li incorporation into 102 secondary minerals. During congruent weathering, the Li released from primary minerals 103 remains in solution and  $\delta^7 \text{Li}_{\text{diss}}$  resembles bedrock values; when weathering is incongruent and a 104 significant amount of Li is incorporated into secondary minerals,  $\delta^7 \text{Li}_{\text{diss}}$  is elevated relative to 105 the minerals. Accordingly, observations of modern river systems show that  $\delta^7 Li_{diss}$  varies as a 106 function of weathering congruency, or the fraction of Li released from primary mineral 107 dissolution that remains in solution and is not incorporated into secondary minerals  $(f_{Li})$ 108 (Dellinger and others, 2015). Notably, the overall pattern of this relationship segregates into two 109 unique relationships, one in upland environments where the degree of  $\delta^7$ Lidiss elevation above 110 bedrock values varies as a linear function  $f_{Li}$ , and the second in floodplain environments where 111  $\delta^7$ Lidiss varies as a power-law or Rayleigh-style function of  $f_{Li}$  potentially due to scavenging of Li 112 by extant phases (Dellinger and others, 2015; Maffre and others, 2020) or the formation of 113 authigenic clays (Zhang and others, 2021). Here, we focus on Li fractionation processes 114 characteristic of upland regolith development.

115 Extending the relationship between  $\delta^7 Li$  and weathering congruence to a metric for weathering fluxes has proven more difficult. For example,  $\delta^7 \text{Li}_{\text{diss}}$  has been empirically linked to 116 117 both chemical weathering fluxes and riverine Li concentrations within specific regions; however, 118 these relationships may be contradictory in both sign and magnitude when compared across 119 regions (Huh and others, 2001; Vigier and others, 2009; Millot and others, 2010). More recent 120 work, both theoretical and observational (Bouchez and others, 2013; Dellinger and others, 2015, 121 2017), has contended that  $\delta^7$ Lidiss reflects weathering intensity (WI), or the ratio of chemical 122 weathering (W) to total denudation, D ( $\sim$ E + W). At steady state regolith thickness, regolith production (RP) must equal total denudation (RP = E + W), hence we define WI = W/RP, as 123 124 outlined in figure 1. Within this framework, weathering congruence and subsequent  $\delta^7 Li_{diss}$  are 125 controlled by weathering intensity, which in turn dictates the combination of mineral residence-126 and fluid travel times in the regolith.

In figure 1, Point A represents a condition of moderate WI where physical and chemical erosion are comparable. Under these conditions, regolith profiles reflect the balance between regolith production and denudation rates, as minerals are supplied via uplift and are partially depleted towards the surface through chemical denudation before they are eroded. As a result of

131 these well-developed regolith profiles, fluid travel times during infiltration are sufficient to 132 support secondary mineral precipitation, mineral residence times are sufficient to support 133 secondary mineral accumulation, and weathering occurs incongruently. This situation results in maximum  $\delta^7$ Li<sub>diss</sub> values and moderate Li concentrations ([Li<sup>+</sup>]; ~10<sup>2</sup> nM; Dellinger and others, 134 2015). When chemical denudation rates are much greater than erosion or regolith production 135 136 (high WI, Point B), chemical weathering depletes primary minerals from soils (i.e. supply-137 limited weathering), and dissolved solutes represent the dissolution of low-solubility secondary 138 mineral phases, characteristic of wet tropical environments (von Blanckenburg and others, 2004). 139 Due to this low solubility and the preferential uptake of <sup>6</sup>Li during initial clay precipitation, clay 140 dissolution results in both low [Li<sup>+</sup>] (<10 nM) and low  $\delta^7$ Lidiss.

141 Finally, under low WI conditions when physical erosion greatly outpaces chemical 142 weathering (Point C), mineral residence times in regolith are short such that secondary minerals 143 are unable to accumulate in significant amounts (i.e. kinetic-limited weathering). Steady-state 144 weathering models that describe this behavior predict that under low WI, clay precipitation rates 145 are reduced relative to primary mineral dissolution rates (Chamberlain and others, 2005; Ferrier and Kirchner, 2008), resulting in congruent weathering and low, bedrock-like  $\delta^7 \text{Li}_{\text{diss}}$  values 146 147 (Bouchez and others, 2013). This conceptual model of congruent weathering under low WI 148 implicitly encapsulates fluid travel times (via their dissolution/precipitation rate 149 parameterizations), suggesting that infiltration rates are too short to support active clay 150 precipitation, possibly due to thin soil mantles. Examples of fluid travel time controls on the 151 extent of  $\delta^7 Li_{diss}$  have also been demonstrated by previous reactive transport studies (Wanner and 152 others, 2014; Bohlin and Bickle, 2019; Golla et al. 2021). We note, however, that observations of 153 low clay content in low WI environments may arise from reduced timescales for accumulation 154 due to short regolith residence times, rather than slow clay precipitation rates. Here, we show 155 that fluid travel time controls lead to a notable discrepancy between theories that predict low 156  $[Li^+]$  at low WI due to short fluid travel times, and observations that show the highest  $[Li^+]$  in 157 low WI systems. To our knowledge, this discrepancy has not previously been articulated. 158 Specifically, observations of [Li<sup>+</sup>] in global rivers show the highest concentrations on the 159 order of  $>10^3$  nM in low WI and rapidly eroding environments (Dellinger and others, 2015; 160 Pogge von Strandmann and Henderson, 2015), leading to a parabolic relationship between  $\delta^7$ Lidiss and [Li<sup>+</sup>], similar to that of  $\delta^7$ Lidiss and WI (fig. 1). However, short fluid travel times and 161

162 reduced clay precipitation should theoretically limit overall primary mineral dissolution and the 163 subsequent accumulation of Li in solution, as balanced by reduced clay Li uptake. In fact, some 164 field sites do show the co-limitation of clay precipitation and associated primary mineral 165 dissolution under short fluid travel times, wherein high river discharge results in diluted [Li<sup>+</sup>] 166 and low  $\delta^7$ Lidiss values relative to baseflow (Lemarchand and others, 2010; Manaka and others, 167 2017). 168 Herein, we explore this apparent paradox and the links between weathering intensity and 169 congruence using a reactive transport model of lithium isotopes. Through the incorporation of 170 lithium isotope dynamics into the CrunchFlow reactive transport code (Steefel and others 2015), 171 we expand on previous observational and theoretical formulations through the contemporaneous 172 representation of fluid flow (Maher 2010), multi-component mineral dissolution and 173 precipitation reactions (Steefel and others, 2015), erosional forcing (Maher and Chamberlain, 174 2014), and isotope fractionation (Druhan and others, 2013; Wanner and others, 2014; Maher and 175 von Blanckenburg, 2016; Golla et al. 2021). Simulations across a range of weathering intensity 176 scenarios are evaluated against observations of riverine  $\delta^7 Li_{diss}$ , making use of an update to the 177 CrunchFlow software facilitating fractionating precipitation and non-fractionating dissolution of 178 a given mineral phase. We also explore the transient evolution of weathering profiles under 179 varying weathering intensities, along with the potential role for changes in weathering zone 180 mineralogy to influence observed lithium isotope dynamics. While we acknowledge the broad 181 and seminal applications of  $\delta^7$ Li to study weathering processes across the globe in many other 182 studies, we restrict our data-model comparison to data presented in Dellinger and others (2014, 183 2015, 2017) based on their framework for interpreting  $\delta^7$ Li signals as a function of Li/Al ratios. 184 This normalization allows comparison across a diversity of bedrock types. 185 186 METHODS 187 188 Mineralogy and Geochemistry 189 190 In order to probe the relationships between weathering intensity, congruence, and  $\delta^7 L_i$ , 191 we simulated upland weathering profiles using the reactive transport software CrunchFlow, 192 making use of the isotope features developed in prior studies and a software update necessary to

193 treat isotopes in systems of high weathering intensity. In CrunchFlow, mineral

194 dissolution/precipitation rates are represented by Transition State Theory (TST)-style rate laws

(cf. Lasaga, 1998) with a dependence on reaction affinity that allows dissolution/precipitation to
 occur only when solutes are under/over-saturated with respect to that given mineral phase, as

198 
$$R_{net} = k_d * SA * \left(\frac{Q}{K_{eq}} - 1\right)$$
(2),

199

where  $R_{net}$  is the net reaction rate for a specified mineral,  $k_d$  is a temperature-dependent intrinsic rate constant, *SA* is mineral surface area, *Q* is the ion activity product, and  $K_{eq}$  is the mineral equilibrium constant, and the  $(Q/K_{eq} - I)$  term represents reaction affinity. The composition of each solid phase is described as a stoichiometric and charge balanced assemblage of primary species, thus assuring mass balance between co-evolving dissolved solutes and solids.

205 One important effect of this affinity dependence is that it directly links the rate of primary 206 mineral dissolution to the rate of secondary mineral precipitation. A common example is the 207 precipitation of Al and Si to form secondary phases, which maintains undersaturated solute 208 concentrations with respect to primary minerals bearing these elements (Zhu and others, 2005; 209 Maher and others, 2006). Hence, the removal of weathering products into clay minerals allows 210 mineral dissolution to continue by increasing the driving force for dissolution via the reaction 211 affinity. This affinity dependence is not included in the regolith models that have primarily 212 informed theoretical considerations of Li isotopes in previous studies (Chamberlain and others, 213 2005; Ferrier and Kirchner, 2008; Bouchez and others, 2013). In these models, variation in 214 primary mineral dissolution occurs independent of clay precipitation, where the latter is 215 parameterized as a constant rate. We note that these models were originally developed to track 216 the evolution of solid-phase weathering zone mineralogy (Chamberlain and others, 2005; Ferrier 217 and Kirchner, 2008). However, dissolved Li isotope compositions in these models are controlled 218 by the relative kinetic rates of primary mineral dissolution and clay precipitation and are thus 219 uniquely sensitive to rate parameterizations (Bouchez and others, 2013). More recently, a 220 CrunchFlow model was developed for Li isotope fractionation associated with an upland shale 221 weathering profile (Golla et al. 2021). This study demonstrated the functionality of explicit

222	coupling between primary mineral dissolution and secondary mineral precipitation and
223	successfully reproduced fluid $\delta^7$ Li profiles through the bedrock vadose zone and groundwater.
224	Here we generalize the approach taken in Golla et al. (2021) using a representative
225	granitic weathering profile simulated as in Maher (2011), with bedrock modeled as a
226	combination of 60% volume quartz, 30% plagioclase feldspar (20% anorthite, 80% albite), and
227	2% halloysite, leaving ~8% protolith porosity. Equilibrium constants were used primarily from
228	the EQ3/EQ6 thermodynamic database (Wolery and others, 1990), supplemented with
229	plagioclase equilibrium constants from Arnorsson and Stefansson (1999). Intrinsic rate constants
230	and activation energies were taken from a compilation by Palandri and Kharaka (2004). Mineral
231	reactions, kinetic rate constants $(k_d)$ , and equilibrium constants $(K_{eq})$ are shown in Table 1, and
232	we note that as in equation (2), affinity dependence is assumed to be linear for simplicity.
233	Additional simulations were conducted including a high solubility chlorite mineral at
234	7.5% volume with quartz reduced to 52.5% to maintain porosity. We include these simulations
235	as Li is often preferentially concentrated in ferro-magnesian minerals, substituting for $Mg^{2\scriptscriptstyle +}$ and
236	Fe <sup>2+</sup> due to similar ionic radii (Tardy and others, 1972; Penniston-Dorland and others, 2017). We
237	use the reaction stoichiometry and equilibrium constants for chlorite and intermediate weathering
238	product vermiculite from Heidari and others (2017) based on constrained reactive transport
239	simulations of weathering in the Shale Hills Critical Zone Observatory. We amend mineral
240	stoichiometry such that $Li^+$ substitutes for $Fe^{2+}$ . Additionally, we assume no fractionation during
241	vermiculitization by enforcing the same <sup>7</sup> Li: <sup>6</sup> Li ratios in chlorite and vermiculite. We note that
242	the choice of chlorite rather than other Li-rich soluble minerals such as lithium-bearing alkali
243	amphiboles is primarily based on the existence well-established CrunchFlow reaction networks
244	constrained by field studies (Heidari and others, 2017). However, the relationship between
245	chlorite lithium dynamics and weathering intensity investigated here are theoretically
246	transferrable to other highly soluble, lithium-rich minerals.
247	
248	Lithium Isotope Parameterizations
249	
250	Lithium isotopes were incorporated into these simulations by defining <sup>6</sup> Li and <sup>7</sup> Li as
251	separate primary chemical species. To capture typical bedrock compositions, Li was
252	incorporated into plagioclase minerals as a solid solution with Na, which offers a simple means

of maintaining charge balance. The <sup>6</sup>Li and <sup>7</sup>Li values are set such that  $\delta^{7}$ Libedrock is 0‰ and total 253 254 concentration in our granitic mineral assemblage is ~40 ppm, within error of average upper 255 continental crust (Teng and others, 2004). The inclusion of Li-rich chlorite in our mineral 256 assemblage raises bedrock Li concentrations to ~60 ppm, more characteristic of shale concentrations (Holland, 1984) and bedrock within the low WI watersheds of the Beni river 257 258 (Dellinger and others, 2015). Based on observations of stoichiometric dissolution of <sup>6</sup>Li and <sup>7</sup>Li 259 from primary silicate minerals, we assume no fractionation during dissolution, and <sup>6</sup>Li and <sup>7</sup>Li 260 are incorporated into a single mineral with a single  $K_{eq}$  and  $k_d$  value as shown in Table 1.

To simulate fractionation during clay precipitation, we employ a solid-solution approach that consists of two isotopic end-member phases: <sup>6</sup>Li-Halloysite and <sup>7</sup>Li-Halloysite (Druhan and others, 2013). The coupled rate expressions for precipitation of these phases are,

264

265 
$$R_{6Li-H} = \frac{[^{6}Li^{+}]}{[^{6}Li^{+}+^{7}Li^{+}]} * k_{6Li-H} * SA_{H} * \left(\frac{Q_{cc}}{K_{eq}} - 1\right), \quad (3) \quad \text{and}$$

266

267 
$$R_{7Li-H} = \frac{[^{7}Li^{+}]}{[^{6}Li^{+}]} * \frac{[^{6}Li^{+}]}{[^{6}Li^{+}+^{7}Li^{+}]} * \alpha_{k} * k_{6Li-H} * SA_{H} * \left(\frac{Q_{cc}}{K_{eq}} - 1\right),$$
(4)

268

269 where  $Q_{cc}$  is the total ion activity product (including <sup>6</sup>Li and <sup>7</sup>Li) with respect to the combined 270 halloysite phases,  $K_{eq}$  is the equilibrium constant (same for both phases), and  $\alpha_k$  is the kinetic 271 fractionation factor (*k*<sub>7*Li*-*H*</sub>/*k*<sub>6*Li*-*H*</sub>) after Druhan and others (2013) and Steefel and others (2014). 272 As described in Steefel and others (2014), this formulation assumes that the isotopic composition 273 of the solid phase does not play a role in fractionation during precipitation through the mineral 274 isotopologue activities. Fractionation occurs solely as a result of specified differences in intrinsic 275 rate constants between isotopologue clay phases. Thus, under appropriately simplified 276 conditions, this model would create Rayleigh distillation-style fractionation in the  $\delta^7$ Lidiss as [Li<sup>+</sup>] 277 is lost from solution due to clay precipitation. In other words, removal of Li from solution due to 278 clay precipitation in the absence of primary mineral Li inputs would act as an open-system 279 process with a constant fractionation at far-from-equilibrium conditions, similar to Rayleigh 280 distillation.

281 In our simulations, we assume a kinetic fractionation factor of  $\alpha_k = 0.982$  for 282 precipitation of the halloysite solid solution. This value is based on calculated fractionation

283 factors from upland environments in the Dellinger and others (2015) dataset and falls roughly in 284 line with experimental values (Vigier and others, 2008). As in previous modeling studies, this 285 fractionation is meant to encompass a range of potential processes including structural 286 substitution, interlayer exchange, and surface sorption (Bouchez and others, 2013; Wanner and 287 others, 2014). Notably, in our treatment, Li uptake by clay is simulated via charge balance with 288 Al (Table 1), which is necessary within CrunchFlow. Rather than the direct structural 289 substitution of Li for Al, this parameterization is meant to reflect effective Li uptake by clays 290 that includes mechanisms such as sorption, interlayer exchange, and structural substitution.

291 The magnitude of observed fractionation in solutes is also controlled by the partition 292 coefficient for Li into clays, which reflects the ratio of lithium dissolved from primary minerals 293 that is incorporated into clays  $(P_{Li})$  versus left in solution  $(f_{Li})$  at equilibrium. Under this 294 formulation,  $P_{Li} = (1 - f_{Li})$ . The result is that isotopic signals in the dissolved phase are maximized 295 when almost all the lithium released via dissolution is incorporated into secondary precipitates. 296 To examine the sensitivity of fractionation to  $P_{Li}$ , we manually vary the concentration of clay Li 297 such that  $P_{Li}$  ranges from 0.01 to 0.99 (x values in Table 1). Under this formulation,  $P_{Li}$  is equal 298 to the ratio of Li/Al in Halloysite to Plag(An<sub>20</sub>) (i.e.  $P_{Li} = (x/2)/(0.0026/1.2)$ ). This range of  $P_{Li}$ 299 values is based on field observations of *f*<sub>Li</sub> across South American watersheds, calculated from 300 ratios of Li/Al in suspended sediments to bedrock (Dellinger et al., 2015).

301 We chose to simulate a simple 1:1 clay phase (halloysite) rather than a suite of competing 302 clay phases including more complex 2:1 clays in order to maintain constant  $P_{Li}$  values within a 303 given model run. This reasoning also underlies our choice to represent an 'effective' clay uptake 304 rather than explicit representations of sorption, interlayer exchanges, and structural substitution. 305 Under these more complex representations,  $P_{Li}$  would necessarily vary within an individual 306 model run through time and with depth as the weathering reaction network evolved or Li actively 307 exchanged with clay phases. While these dynamics certainly impact Li during weathering, they 308 have not been adequately linked to mineral residence or fluid travel times in natural 309 environments, which are the primary focus of our analyses. Thus, any encapsulated links 310 between PLi and erosion and flow rates would not be justified by previous literature. Our 311 simplified representation allows us to isolate the impacts of erosion and flow rates across a range 312 of set  $P_{Li}$  values and facilitates the direct comparison to Dellinger and others (2015, 2017) 313 observations. As described above, these are based on Li/Al ratios in bedrock versus dissolved

and suspended loads and thus also represent a composite suite of 'effective' uptake mechanisms. We note that x is the same within any given simulation for <sup>6</sup>Li-Halloysite and <sup>7</sup>Li-Halloysite such that fractionation is only induced by the kinetic fractionation factor. For simplicity, we assume

- 317 that Li substitution at ppm levels does not impact  $K_{eq}$  values (Wang and Xu, 2001).
- 318

314

315

316

319320

### Software Update for Isotope Partitioning at High Weathering Intensity

321 As discussed above, the high weathering intensity regime requires that isotopically 322 distinct secondary clays are ultimately re-dissolved into the fluid phase. However, equations (3) and (4) are not applicable under conditions where  $\frac{Q_{cc}}{K_{eq}} < 1$  for the following reasons. First, the 323 324 fractionation factor ( $\alpha_k = 0.982$ ) would result in a more rapid reaction rate for the <sup>6</sup>Li endmember relative to <sup>7</sup>Li during dissolution. This would mean that the <sup>6</sup>Li-bearing halloysite (eq. 325 326 3) would dissolve faster than its counterpart, leading to spurious fractionation over the timescales 327 considered here. Second, equations (3) and (4) contain no reference to the isotopic composition 328 of the solid phase, and thus there is no ability to incorporate the isotope ratio of the halloysite 329 that is being solubilized into the fluid phase within this framework. To address these issues, we 330 use the version of the isotopic solid solution model originally developed in CrunchFlow (Druhan 331 and others, 2013) where the solid solution ratio is based on the isotopic mole fractions of the 332 mineral phase:

333

334

$$R_{6Li-H} = Y_{6Li-H} * k_{6Li-H} * SA_H * \left(\frac{Q_{cc}}{Y_{6Li-H} * K_{eq}} - 1\right), \quad (5) \text{ and}$$

335

336 
$$R_{7Li-H} = Y_{7Li-H} * \alpha_k * k_{6Li-H} * SA_H * \left(\frac{[^{7}Li^+]}{[^{6}Li^+]} \frac{Q_{cc}}{Y_{7Li-H} * K_{eq}} - 1\right), \quad (6)$$

337

Where *Y* is the isotopic mole fraction or the number of moles of a given isotope of lithium pertotal number of moles of lithium in the halloysite phase.

340 To stoichiometrically dissolve halloysite that was originally formed following the 341 coupled rate laws given in equations (3) and (4), we modify the original formulation of equations 342 (5) and (6) such that the kinetic fractionation factor  $\alpha_k$  is set to 1.0 (no fractionation) and thus the 343 two minerals dissolve at the same relative rates. The result is that the isotope ratio of the neo-

344 formed halloysite mineral will be delivered into the fluid phase, ensuring stoichiometric

345 dissolution without spurious fractionation:

346

347 
$$R_{6Li-H} = \begin{cases} eq. 3, \ \frac{Q_{cc}}{\kappa_{eq}} \ge 1\\ eq. 5, \ \frac{Q_{cc}}{\kappa_{eq}} < 1 \end{cases}$$
(7a) and

348 
$$R_{7Li-H} = \begin{cases} eq. 4, \alpha_k = 0.982, \frac{Q_{cc}}{K_{eq}} \ge 1\\ eq. 6, \alpha_k = 1.000, \frac{Q_{cc}}{K_{eq}} < 1 \end{cases}$$
(7b)

A similar approach was utilized in Golla et al. (2021) for the development of a weathering
profile model incorporating Li isotopes specifically to reproduce a measured dataset of vadose
zone and groundwater samples.

Weathering Scenarios

352 353

We simulate four separate weathering scenarios, sequentially increasing in complexity, to test the controls of weathering regimes on Li systematics as outlined in Table 2.

- Scenario 1- Batch Reactor: we simulate batch reactor weathering reactions, to evaluate
   temporal trends in solute concentrations and to analyze the sensitivity of Li fractionation
   to Li congruence.
- Scenario 2- Chronosequence: we simulate a 1-D weathering profile chronosequence with
   no uplift/erosion to evaluate temporal trends in mineralogical and solute profiles.
- Scenario 3- Uplift/Erosion: we simulate 1-D weathering profiles with specified regolith
   production rates to test the controls of weathering intensity.



365 Model set-up and boundary conditions are described in detail in the following subsections.

- 367 Scenario 1 Batch reactors: To examine the general characteristics of our Li isotope
- 368 parameterizations, we first simulate a simple batch reactor with no transport. These are
- 369 initialized with an un-weathered primary mineral composition, dilute waters, a temperature of
- $15^{\circ}$  C, and fixed open-system pCO<sub>2</sub> of 5000 ppm, setting an initial pH of 5.0 that increases with

371 progressive mineral dissolution/precipitation reactions. This  $pCO_2$  choice is meant to reflect 372 typical elevated values in soil environments due to oxidation of organic matter. We note that 373 increased (decreased) pCO<sub>2</sub> leads to increased (decreased) equilibrium solute concentrations 374 (Winnick and Maher, 2018), but does not influence isotopic evolution. Within these batch 375 simulations, we vary the partition coefficient  $P_{Li}$  from 0.01-0.99 via x values as defined in 376 Lithium Isotope Parameterizations and shown in Table 1 for the halloysite phase. In each 377 simulation, reaction progress is monitored for 5 kyr. We also demonstrate how the simulation of 378 a 2:1 clay (Ca-beidellite) results in the same lithium fractionation dynamics as a function of  $P_{Li}$ 379 as halloysite in Appendix A.

380

381 Scenario 2 - Chronosequence: Next, we simulate 1-D weathering profiles with no erosion to 382 characterize the temporal evolution of solid and aqueous geochemical profiles. Weathering 383 domains are set to 10 m with a grid cell size of 5 cm (200 cells). Initial porosity is set to 7%, and weathering occurs under 75% water saturation and a large prescribed gas flux of CO<sub>2</sub> into the 384 385 domain to maintain open-system weathering conditions (Winnick and Maher, 2018). As in the 386 batch simulations, profiles are initialized with an un-weathered primary mineral composition, 387 dilute waters, a temperature of  $15^{\circ}$  C, and pCO<sub>2</sub> of 5000 ppm. Water infiltrates through the 388 domain via Darcy flux with a prescribed rate of 0.5 m/yr. Simulations are run for 1 Myr, and 389 changes in solid and aqueous geochemistry are analyzed at multiple timesteps.

390

391 *Scenario 3 - Uplift/Erosion:* We analyze the effects of variable weathering intensity by

392 simulating 1-D weathering profiles with prescribed volumetric uplift/erosional forcing.

393 CrunchFlow incorporates uplift/erosion through the specification of a volumetric uplift rate at the

bottom of the domain equivalent to a regolith production rate. Material is removed from the top

395 of the domain at the same volumetric rate; however, mass erosion rates are reduced relative to

regolith production due to chemical denudation of material before it is uplifted and removed at

397 the surface (fig. 1). Weathering domains, water saturation, and  $pCO_2$  are set as in

398 chronosequence simulations. To examine the relationship between lithium isotope dynamics and

399 reactive transport, we simulated a suite of weathering regimes by varying infiltration and erosion

400 rates. Infiltration (Darcy flux) was varied between 0.1-5 m/yr, and regolith production

401 (volumetric erosion) was varied between 0.005-0.2 mm/yr, corresponding to erosion rates of 1.4

402 - 485 t/km²/yr after correcting for porosity changes during weathering (Table 3), resulting in 30
403 unique weathering intensity regimes. Each weathering simulation was run for 0.5 Myr, resulting
404 in either the development of a steady-state profile (constant aqueous and solid geochemical depth
405 profiles) or transient profiles (time-dependent geochemical depth profiles characterized by the
406 progressive depletion of primary minerals). Model output was analyzed at multiple timesteps to
407 characterize the temporal evolution of mineral and solute profiles.

408 We define the regolith weathering zone within each time-step as model cells with >101% 409 initial bedrock porosity. This depth is meant to represent a conservative estimate of the 410 weathering front based on observable changes in bulk density (Maher and von Blanckenburg, 411 2016). We note that other modeling studies evaluate weathering depths based on solute 412 equilibrium length scales (Lebedeva and others, 2010), which in our simulations always occur 413 below the porosity-based weathering front. Our definition of weathering zone is meant to capture 414 potential contrasts in permeability that may affect subsurface flowpaths and travel times. 415 Specifically, due to the model setup which specifies constant Darcy flow, simulated water fluxes 416 do not vary within the profile domain based on porosity changes. To capture short fluid travel 417 times associated with preferential and interflow caused by thin regolith and sharp permeability contrasts between weathered and fresh bedrock, we evaluate solute concentrations and  $\delta^7 Li_{diss}$  at 418 419 the base of this regolith weathering zone in addition to the base of the model domain. In the 420 simulations with low regolith production rates that result in weathering of the entire domain, we 421 evaluate solute concentrations at 9 m depth to avoid boundary condition effects associated with 422 the bedrock boundary condition at 10 m.

423 Model outputs and metric calculations are shown in Table 3. Mass erosion rates are 424 calculated by multiplying the specified regolith production rate by bulk density in the top model 425 cell. Riverine suspended material is taken as the mineralogical composition of the eroding top 426 cell of the domain. Chemical denudation rate is calculated as the sum of major dissolved weathering products ( $[Ca^{2+}]+[Mg^{2+}]+[Na^{+}]+[SiO_{2(aq)}]$ ) multiplied by the prescribed 427 428 infiltration rate at both the domain boundary and at the bottom of the regolith weathering zone. 429 Finally, a metric of weathering incongruence (I) is calculated based on the ratio of the 430 instantaneous rate of hallovsite precipitation to plagioclase dissolution ( $I = -R_{hallovsite}/R_{plagioclase}$ ; 431 as per equation (2), precipitation is a positive rate, and dissolution is a negative rate. This metric 432 is similar to qualitative descriptions of weathering incongruence under conditions of plagioclase

dissolution and halloysite precipitation, in which incongruence represents the molar ratio of clay
formation to primary mineral dissolution. When halloysite dissolves, however, incongruence is
calculated as a negative number.

436

437 Scenario 4 – Uplift/Erosion with chlorite.— To address the potential role of soluble, Li-rich 438 ferro-magnesian minerals such as chlorite in driving global  $\delta^7$ Lidiss patterns, we conduct a series 439 of simulations including an amended chlorite mineral as shown in Table 1. The weathering of 440 chlorite minerals involves a multi-step reaction series by which, first, cations in the octahedral coordination layer ( $Mg^{2+}$ ,  $Fe^{2+}$ ) are leached in a process referred to as supergene 441 'vermiculitization' (Proust and others, 1986). In general, this weathering step is not well-442 443 characterized in field environments due to analytical challenges involved in distinguishing 444 unaltered chlorite and vermiculite. Recent studies, however, have made use of ratios of 7Å to 14 445 Å XRD peaks along with stoichiometric characterizations to demonstrate vermiculitization in 446 deep regolith profiles (Sullivan and others, 2016; Gu and others, 2020). Following 447 vermiculitization, which may involve intermediate products of interstratified 448 chlorite/vermiculite, vermiculite products are further weathered to kaolinite-type clays. 449 To represent this multi-step process, we have included an intermediate, lower solubility 450 vermiculite mineral in addition to chlorite following the methodology of Heidari and others 451 (2017) (Table 1). In terms of Li<sup>+</sup>, we assume no fractionation during 'vermiculitization' by incorporating equivalent <sup>7</sup>Li<sup>+</sup>:<sup>6</sup>Li<sup>+</sup> ratios in mineral stoichiometry. Further, we assume that Li<sup>+</sup> 452 leaching relative to  $Al^{3+}$  falls intermediately between simulated leaching of  $Mg^{2+}$  and  $Fe^{2+}$  in 453 order to maintain realistic dissolved Li concentrations ( $<10^4$  nM). 454 455 In these chlorite simulations, bedrock is initialized as 52.5% guartz, 30% plagioclase 456 (An20), 7.5% 'chlorite' (6% chlorite, 1.5% vermiculite), and 2% halloysite. We note that these 457 mineral constituents are similar to quartz diorite in the Luquillo Critical Zone Observatory 458 (Murphy and others, 1997), whereas weathering fluxes from shale environments are more likely

460 others, 2020). We conduct 1-D uplift/erosion weathering profiles under the same conditions as
461 described above.

to be dominated by chlorite, other clay minerals, with smaller amounts of plagioclase (Gu and

462

459

463

### RESULTS

464	
465	Batch Reactor Results (Scenario 1)
466	
467	At the start of batch reactor simulations (Scenario 1), dilute waters are undersaturated
468	with respect to both plagioclase and halloysite, and primary mineral dissolution and secondary
469	mineral precipitation proceed as solute concentration profiles stabilize over roughly 10 <sup>2</sup> years
470	(fig. 2). When the simulation begins, plagioclase dissolution is at its maximum rate, which is
471	sustained for the first ~6 months of simulation, representing kinetic controls. As solute
472	concentrations increase and waters approach saturation, a quasi-steady state develops in the
473	aqueous profiles and the affinity term in equation (2) becomes the limiting factor on dissolution
474	rates, which subsequently approach zero. We note that in these batch reactor scenarios,
475	plagioclase depletion is minimal, such that changes in surface area do not impact dissolution
476	rates through time (Eq. 2). Secondary mineral rates at the start of simulations denote net
477	dissolution, though at rates orders of magnitude slower than plagioclase dissolution, for the first
478	few days of simulation time as dilute solutes are undersaturated with respect to halloysite. As
479	$Al^{3+}$ and $SiO_2(aq)$ activities increase due to plagioclase dissolution, halloysite transitions to net
480	precipitation; however, due to the low intrinsic rate constant, net precipitation rates are initially
481	subdued. Precipitation rates increase to a maximum value roughly 2.5 months into the simulation
482	as solutes become orders of magnitude oversaturated with respect to halloysite and then
483	subsequently decrease over the rest of the simulation.
484	Instantaneous weathering incongruence is shown as the ratio of rates of halloysite
485	precipitation to plagioclase dissolution in figure 2b. At the start of the simulation, weathering is

485 precipitation to plagioclase dissolution in figure 2b. At the start of the simulation, weathering is 486 fully congruent (ratio of zero) until the initiation of halloysite precipitation. Increasing halloysite 487 precipitation results in an overshoot in weathering incongruence, whereby more halloysite 488 precipitates than predicted based on equilibrium reaction stoichiometry to compensate for initial 489 congruent weathering. Following this overshoot, the ratio of halloysite precipitation to 490 plagioclase dissolution approaches the stoichiometric balance of ~0.6 from the equilibrium 491 reaction,

493 
$$1.66Plag(An20) + 2CO_2(g) + 3H_2O \leftrightarrow Halloysite + 0.33Ca^{2+} + 1.33Na^+ + 2.66SiO_2(aq) + 2HCO_3^-$$
 (8).

495

- We note that even as dissolution/precipitation rates decrease through time as the systemapproaches thermodynamic equilibrium, this stoichiometric ratio is maintained.
- Simulated  $Ca^{2+}$  concentrations (fig. 2c) are influenced only by plagioclase dissolution, as 498 499  $Ca^{2+}$  is not incorporated into halloysite. Concentrations rise rapidly over the first ~6 months of 500 simulation, after which increases attenuate as dissolution rates decrease. Maximum 501 concentrations of ~45 µM are reached at roughly 500 years of simulation as the system 502 approaches equilibrium, and this final concentration is dependent on the specified open-system 503  $pCO_2$  through its control on initial and equilibrium pH (5.0 and 8.9, respectively) (Winnick and Maher, 2018). The evolution of Li<sup>+</sup> differs from Ca<sup>2+</sup> due to the incorporation of Li<sup>+</sup> into 504 505 precipitating halloysite. Concentrations are therefore strongly dependent on the partition 506 coefficient ( $P_{Li}$ ) as shown in fig. 2d. In the initial stages of weathering, Li<sup>+</sup> increases similarly 507 across the range of partition coefficients, and after hallovsite precipitation initiates, partition 508 coefficient simulations diverge. Simulations with low partition coefficients in which relatively 509 little of the Li<sup>+</sup> released during plagioclase dissolution is incorporated into hallovsite result in rapidly increasing concentrations, similar to  $Ca^{2+}$ . In contrast, simulations with high partition 510 511 coefficients result in a transient decrease in Li<sup>+</sup> reflecting the overshoot in weathering 512 incongruence, followed by slowly increasing values as the system approaches equilibrium.

513 Partition coefficients also control the evolution of  $\delta^7 \text{Li}$  (fig. 3). When the partition 514 coefficient is low and little Li<sup>+</sup> is incorporated into clays, the extent of fractionation is reduced, 515 and  $\delta^7 \text{Li}_{\text{diss}}$  is similar to  $\delta^7 \text{Li}_{\text{plag}}$ . When the partition coefficient is large,  $\delta^7 \text{Li}_{\text{diss}}$  reflects the full 516 18‰ fractionation factor. Additionally, the transient weathering incongruence overshoot results 517 in transiently elevated  $\delta^7 Li_{diss}$  values, which subsequently approach steady-state values. We note 518 that, similar to instantaneous weathering incongruence,  $\delta^7 \text{Li}_{\text{diss}}$  reach steady-state values at 519 roughly 6 months, which is long before the system as a whole approaches equilibrium. Rather 520 than overall reaction progress,  $\delta^7$ Lidiss reflects the approach to stoichiometric balance between 521 plagioclase dissolution and hallovsite precipitation.

In figure 3b, we show the relationship between  $\delta^7 \text{Li}_{\text{diss}}$  and  $f_{Li}$ , which in this case reflects the weathering congruence of Li<sup>+</sup> (combining both congruence changes in the approach to stoichiometric balance and  $P_{Li}$ ). Steady-state  $\delta^7 \text{Li}_{\text{diss}}$  values (colored circles reflecting plateau values in fig. 3a) are linearly related to  $f_{Li}$ , similar to the 'batch-reactor' relationship described by

526	Bouchez and others (2013) and Dellinger and others (2015). The approach to steady-state $\delta^7 Li_{diss}$
527	values within each $P_{Li}$ simulation (colored lines) involves a stronger dependence on $f_{Li}$ than
528	steady-state values, which is due to the previously described overshoot in halloysite precipitation
529	rates and weathering incongruence. Global riverine $\delta^7 Li_{diss}$ observations from Dellinger and
530	others (2015) are also shown and fall roughly in line with CrunchFlow simulations, with the
531	exception of flood-plain dominated environments as described above, which require sustained
532	open-system clay uptake as in Maffre and others (2020).
533	
534	Weathering Profiles – Chronosequence (Scenario 2)
535	
536	Reactive transport simulations of weathering profiles have been presented in a number of
537	previous studies (Maher et al., 2009; Maher, 2010, 2011; Moore et al., 2012; Lebedeva and
538	Brantley, 2013; Lawrence et al., 2014; Pandey and Rajaram, 2016; Heidari and others, 2017;
539	Winnick and Maher, 2018; Jung and Navarre-Sitchler, 2018; Perez-Fodich and Derry, 2019;
540	Golla et al. 2021). We limit this discussion to a brief overview of general weathering
541	characteristics and refer readers to these references for further details. Results from our
542	simulations are shown in figure 4 with a $P_{Li}$ value of 0.9, where colored lines are used to
543	represent the evolution of solid and aqueous profiles through time.
544	At the beginning of the simulation, the solid profile represents fresh, unweathered
545	bedrock. For this choice of initial conditions, the spatial evolution of aqueous solutes and mineral
546	reaction rates generally resemble the temporal evolution of the batch reactor experiments
547	presented above. Plagioclase weathering begins immediately at the surface as dilute, weakly
548	acidic (pH = 5.0) waters infiltrate, releasing $Ca^{2+}$ and $Li^+$ (along with other reaction products not
549	shown) into solution. Halloysite precipitation begins at ~20 cm depth, reflecting the water travel
550	times necessary to significantly oversaturate solutes with respect to halloysite, as in figure 2a.
551	This balance of dissolution/precipitation is reflected in weathering incongruence (fig. 4c), where
552	incongruence increases at the depth where halloysite precipitation becomes significant, reaches a
553	maximum at $\sim 0.75$ m, and approaches the stoichiometric balance of plagioclase dissolution to
554	halloysite precipitation (0.6) at roughly 2 m. As with the batch reactor experiments (fig. 3a),
555	$\delta^7$ Li <sub>diss</sub> tracks weathering incongruence, approaching enriched steady-state values at ~1 m (fig.
556	4f).

557 As the simulation moves through time, plagioclase is progressively depleted from the 558 shallow depths, with this pattern propagating downward through the profile. Plagioclase is 559 largely depleted from the 10 m weathering profile by the end of the simulation at  $10^6$  yrs; 560 however, small amounts remain as the surface area term in equation (2) limits kinetic rates of 561 dissolution. Halloysite builds up in the profile from top to bottom due to progressive 562 precipitation, though, precipitation is limited in the top 20 cm due to slow kinetics. Additionally, 563 halloysite transitions to dissolution when plagioclase is sufficiently depleted at the top of the 564 profile. Solute profiles of  $[Ca^{2+}]$  and  $[Li^{+}]$  generally track the evolution of mineral profiles on 565 these long timescales. Specifically, decreases in  $[Ca^{2+}]$  and  $[Li^{+}]$  at a given depth through time 566 match plagioclase depletion and are related to reduced mineral surface area and subsequent 567 decreasing dissolution rates.

Weathering incongruence and subsequent  $\delta^7 \text{Lid}_{\text{idiss}}$  approach steady-state values at 568 569 increasing depths through time. This is tied to reduced plagioclase dissolution rates in the upper 570 profile as primary mineral is depleted, resulting in greater travel times needed to initiate 571 hallovsite precipitation and approach  $\delta^7 \text{Li}_{\text{diss}}$  steady-state values. Despite these longer  $\delta^7 \text{Li}_{\text{diss}}$ 572 length scales, steady-state values are approached by the bottom of the 10 m profile until plagioclase is depleted throughout the profile at  $10^6$  yrs. When plagioclase has been depleted, the 573 574 weathering regime transitions to net clay dissolution throughout the profile, and  $\delta^7 Li_{diss}$  values 575 become more negative, resembling the clav values that were precipitated earlier in the 576 simulation. However, through time, very small rates of plagioclase dissolution cause  $\delta^7 Li_{diss}$  to 577 evolve during transport and fall between bedrock and clay values by the bottom of the profile. 578 This last timestep also coincides with low [Li<sup>+</sup>] due to slow dissolution kinetics of halloysite. 579 These results are consistent with some behavior shown in tropical weathering environments 580 (Lara and others, 2022) in which  $\delta^7$ Lidiss are extremely low in weathered soils (though absolute values of measured pore waters and clay  $\delta^7 Li$  would require extremely low  $P_{Li}$  values and large 581 582 clay fractionation) and are disconnected from stream environments as deeper weathering and 583 higher [Li<sup>+</sup>] from primary mineral dissolution seemingly overwhelm clay dissolution. 584

585

*Uplift/Erosion (Scenario 3)* 

As described previously, we simulate a range of both uplift/erosion and flow rates to capture a broad range of weathering intensity regimes. Across this suite, 30 unique WI regimes were tested, varying from  $2.7 \times 10^{-5}$  to 0.93. Illustrative examples of low weathering intensity (high uplift/erosion, low flow), moderate weathering intensity (comparable uplift/erosion and flow), and high weathering intensity (low uplift/erosion, high flow) environments are compared in figure 5. We note that in figure 5, only the low weathering intensity simulation reaches steadystate with respect to the mineral abundances and fluid compositions, as discussed below.

594 Under low weathering intensity, the supply of fresh bedrock overwhelms chemical 595 denudation, and weathering profile solid compositions reflect bedrock values throughout the 596 simulation (fig. 5a-c). In the illustrative example, plagioclase is very slightly depleted within the 597 top <1 m. Based on our definition of the regolith weathering zone (porosity >101% bedrock 598 porosity), this profile corresponds to a regolith depth of 5 cm, and solute concentrations are taken 599 from this depth when comparing between weathering intensity simulations. Similar to solid 600 composition, weathering incongruence and  $\delta^7 \text{Li}_{\text{diss}}$  profiles do not vary significantly with depth 601 or through the course of the simulation. As water infiltrates through the 1D un-weathered 602 bedrock profile, the behavior is largely comparable to the batch reactors; however, the approach 603 to stoichiometric weathering incongruence and steady-state  $\delta^7 \text{Lid}_{\text{id}\text{iss}}$  values occur at depths below 604 the defined regolith weathering zone. As a result,  $\delta^7 \text{Li}_{\text{diss}}$  values from this simulation are ~0% at 605 the weathering zone depth of 5 cm.

606 When erosion rates and chemical denudation are comparable under moderate weathering 607 intensity, weathering profiles evolve with depth throughout the duration of the simulation. 608 Through time, plagioclase is depleted and halloysite is precipitated in the upper portion of the 609 profile. In contrast to the no-erosion simulation in figure 4, plagioclase reaction fronts are 610 condensed, reflecting the additional uplift flux of bedrock. Similarly, the erosion of material 611 from the surface eliminates the diminished accumulation of halloysite in the top few cm's of the 612 profile as compared to figure 4. In figure 5e-h, regolith depth reaches a maximum of  $\sim 8$  m in 10<sup>6</sup> 613 vrs. As with the no-erosion case, weathering incongruence and  $\delta^7 L_{idiss}$  track the evolution of the 614 solid profile, with an increasing length needed to approach stoichiometric incongruence and steady-state  $\delta^7$ Lidiss values. Throughout the simulation, both incongruence and  $\delta^7$ Lidiss approach 615 616 steady-state maximum values at the base of the weathered regolith profile. At 10<sup>6</sup> years when 617 plagioclase is completely depleted from the top of the profile, clay dissolution occurs over the

618 top ~1 m during infiltration; however, the associated low  $\delta^7$ Lidiss values are quickly overwhelmed 619 by the influx of lithium when waters reach the plagioclase dissolution front.

620 Under high weathering intensity, the lack of uplift/erosional forcing results in the 621 depletion of primary minerals and accumulation of secondary minerals as shown in figure 5i-l. We note that while patterns resemble those presented in figure 4, a few differences exist largely 622 623 related to differences in infiltration rate (5 m/yr in fig. 5 v. 0.5 m/yr in fig. 4). As a result of these 624 high infiltration rates, overall weathering intensity increases, causing plagioclase depletion 625 almost uniformly throughout the profile rather than in a discrete reaction front. Another 626 important difference is the approach to bedrock conditions at 10 m depth, which reflects the 627 imposed uplift/erosion boundary condition. Similar to figure 4, weathering incongruence profiles 628 track the solid profile composition. At early time, weathering incongruence remains low over the 629 first ~1 m of infiltration before approaching stoichiometric incongruence coincident with the approach to steady-state  $\delta^7$ Lidiss values; this is due to fast infiltration rates which result in reduced 630 water travel times in comparison to figure 4. At simulation time  $10^5$  yrs, very little plagioclase 631 632 remains throughout the profile. As a result, while weathering incongruence reaches 633 stoichiometric balance by the bottom of the profile, overall plagioclase dissolution rates and 634 subsequent Li release from primary minerals are so low that  $\delta^7 \text{Li}_{\text{diss}}$  reflects a mixture of 635 halloysite dissolution at the top of the profile (negative incongruence) and plagioclase 636 dissolution/hallovsite precipitation at the bottom. Following the complete denudation of plagioclase from the upper profile by 2.5x10<sup>5</sup> yrs, weathering incongruence and  $\delta^7$ Li<sub>diss</sub> remain 637 638 low throughout the profile.

- 639
- 640

### Erosion/Uplift with Chlorite (Scenario 4)

641

The introduction of soluble chlorite minerals changes lithium weathering dynamics within our simulations. In these simulations, the rapid vermiculitization of chlorite releases Li congruently. As a result, dissolved Li concentrations are high (~1  $\mu$ M), and  $\delta^7$ Li<sub>diss</sub> values are close to bedrock values when chlorite is present. Plagioclase dissolution and subsequent halloysite precipitation continue to a reduced degree in the presence of chlorite but have little effect on Li concentrations and isotopes due to the relatively small amount of Li released and removed in comparison to vermiculitization.

649 The temporal evolution of 1-D weathering profiles with chlorite can be seen in the high 650 weathering intensity panels in figure 6. Due to its high solubility, chlorite is rapidly weathered to vermiculite, involving the leaching of Mg<sup>2+</sup> and Li<sup>+</sup> throughout the weathering profile over 651 <1000 years of simulation. When chlorite is present in the profile, Li concentrations are high and 652 653  $\delta^7$ Lidiss resemble bedrock values. However, after the rapid depletion of chlorite, the high 654 weathering intensity simulations behave similar to those in figure 5i-l. Plagioclase is depleted 655 from the profile over timescales of 10<sup>5</sup> yrs, after which clay dissolution is the primary contributor 656 to Li concentrations and  $\delta^7 Li_{diss.}$ 

Similar dynamics are observed under moderate weathering intensity (fig. 6 e-h). Despite elevated uplift/erosion rates relative to the high weathering intensity simulation, protolith chlorite is still depleted from the profile in  $<10^5$  years. As before, once chlorite is depleted, the temporal evolution of the profile behaves as in figure 5 e-h, in which incongruent plagioclase weathering leads to high  $\delta^7$ Lidiss values and Li concentrations on the order of  $10^2$  nM, both of which are dependent on  $P_{Li}$ .

In the low weathering intensity scenario, uplift/erosion rates are sufficiently high that significant chlorite mineral concentrations remain within the weathering zone throughout the simulation (fig. 6 a-d). In other words, under high enough uplift/erosion and low enough water infiltration rates, chlorite becomes kinetic- rather than supply limited. As a result, Li concentrations remain high (~10<sup>3</sup> nM) and  $\delta^7$ Lidiss reflect bedrock values throughout 1 Myr of simulation within the shallow weathering zone.

These results are broadly consistent with field studies demonstrating deep chlorite 'vermiculitazation' or the leaching and oxidative weathering of octahedral cations ( $Mg^{2+}$ , Fe<sup>2+</sup>) relative to plagioclase depletion depths under low to moderate erosion rates, along with the presence of fresh chlorite in shallow soils under high erosion rates (Sullivan and others, 2016; Gu and others, 2020). Mineralogic profiles in tropical weathering environments have also demonstrated the complete depletion of chlorite in weathered saprolite and soil (Murphy and others, 1998).

676

677

*Weathering Intensity,*  $\delta^7 Li$ , and  $[Li^+]$ 

679 In figure 7, we compare the relationships among  $\delta^7 \text{Li}_{\text{diss}}$  and both weathering intensity 680 and [Li<sup>+</sup>] from Scenarios 3 and 4 and global observations from Dellinger and others (2015) and 681 references therein. Figure 7 shows model output at the 500,000-year timestep for simulations 682 with a  $P_{Li}$  value of 0.9 (maximum isotopic offset of ~17‰). We also discuss the sensitivity of model results to variable PLi values in Appendix A. This timestep choice allows for the 683 684 development of deeply weathered profiles under low erosion conditions, characteristic of tropical 685 craton environments; at earlier timesteps, many high WI simulations have not fully transitioned 686 to clay dissolution. In both our simulations without and with the inclusion of chlorite,  $\delta^7 L_{idiss}$ 687 follows a parabolic pattern with low (bedrock-like) values at both high and low weathering 688 intensity. In agreement with observations and as previously theorized, low  $\delta^7 \text{Li}_{\text{diss}}$  at high 689 weathering intensity is due to long mineral residence times that result in the depletion of primary 690 minerals and the dominance of secondary mineral dissolution across both classes of simulation. 691 However, despite similar  $\delta^7 \text{Li}_{\text{diss}}$  values at low weathering intensity, the mechanisms underlying 692 this behavior vary between simulations. When chlorite is not included in our mineral 693 assemblage, low weathering intensity  $\delta^7 Li_{diss}$  values are due to rapid fluid travel times through 694 the shallow weathering zone that limit secondary mineral precipitation. In contrast, low 695 weathering intensity  $\delta^7 \text{Li}_{\text{diss}}$  values with the inclusion of chlorite are caused by the active 696 weathering of soluble, Li-rich minerals which overcome clay precipitation uptake and are 697 sustained only through the rapid uplift supply of fresh bedrock.

698 As a result of these differing mechanisms, the relationships between  $\delta^7 \text{Li}_{\text{diss}}$  and [Li<sup>+</sup>] 699 differ between the two classes of simulation. When chlorite is not included,  $\delta^7 \text{Li}_{\text{diss}}$  is low at low 700 [Li<sup>+</sup>] and high at high [Li<sup>+</sup>]. Under high weathering intensities when clay dissolution dominates 701 Li release, low clay solubilities and lack of a precipitating phase result in low [Li<sup>+</sup>] and  $\delta^7 Li_{diss.}$ 702 Under low weathering intensities, fluid travel time limitations on secondary mineral precipitation 703 also limit primary mineral dissolution, and  $[Li^+]$  is unable to accumulate. Only under moderate 704 weathering intensities when incongruent weathering (eq. 8) can occur within the weathering zone 705 does  $[Li^+]$  approach equilibrium concentrations: however, we note that even these  $[Li^+]$  values 706 are roughly an order of magnitude less than maximum riverine [Li<sup>+</sup>] observations. When chlorite 707 is included, simulations produce the observed parabolic relationship between  $\delta^7 \text{Li}_{\text{diss}}$  and [Li<sup>+</sup>], as 708 the dissolution of chlorite at low weathering intensities allows for [Li<sup>+</sup>] to accumulate in solution 709 and approach the range of maximum observed  $[Li^+]$  values (10<sup>3</sup> nM).

710	Lastly, figure 8 shows simulated $\delta^7 Li_{sus}$ , taken as the isotopic composition of lithium in		
711	plagioclase and halloysite (and chlorite when included) in the eroding uppermost model domain		
712	cell (0.025 m) at the 250 ka time-step, in comparison to values from Dellinger and others (2017).		
713	We exclude low-Li quartz eroded from the surface in our calculation, as Dellinger and others		
714	(2017) report values from the fine fraction of sediment. Figure 8 shows results from both classes		
715	of simulation (with and without chlorite), between which the overall pattern does not vary. At		
716	low weathering intensity, the erosion of minimally-weathered bedrock from the profile surface		
717	results in bedrock-like $\delta^7$ Li <sub>sus</sub> values that vary across our simulations based on initialized $\delta^7$ Li		
718	values of halloysite in bedrock. Above weathering intensities of ~0.01, plagioclase (and chlorite)		
719	is increasingly depleted from eroding surfaces in our simulations. As a result, $\delta^7 Li_{sus}$ values		
720	decrease, approaching $\delta^7$ Li values of precipitating clay, which are dependent on $P_{Li}$ .		
721	Observational data follow this general trend as well, with low $\delta^7 Li_{sus}$ values at high weathering		
722	intensity falling roughly in the center of our simulation ranges.		
723			
724	DISCUSSION		
725			
726	The results from our series of CrunchFlow weathering simulations suggest a number of		
727	important controls on Li systematics and their relationship to weathering regimes. We first		
728	compare model output to global river observations and review the primary drivers of Li isotope		
729	and concentration signals including weathering congruence, mineral composition, mineral		
730	residence times in regolith, and fluid travel times. Next, we discuss the inability of fluid travel		
731	time controls to capture observed relationships between weathering regime, $\delta^7 \text{Li}_{\text{diss}}$ , and [Li <sup>+</sup> ].		
732	Instead, we show that the incorporation of soluble, Li-rich chlorite phases in our weathering		
733	experiments produces signals consistent with global observations. Additionally, we describe an		
734	alternative hypothesis outside the scope of CrunchFlow simulations that geothermal water		
735	contributions may also link weathering regimes and Li isotopes and concentrations in river		
736	systems. Lastly, we discuss model limitations along with need for future studies to evaluate these		
737	hypotheses in the field.		
738			
739	Insights from Reactive Transport Simulations		
740			

741 With a relatively simple representation of kinetic lithium isotopic fractionation during 742 clay precipitation, we are able to capture a number of key features of global riverine Li data. 743 First, as shown in figure 3, the simulated offset between  $\delta^7 \text{Li}_{\text{diss}}$  and  $\delta^7 \text{Li}_{\text{bedrock}}$  is a function of the 744 fraction of Li dissolved that remains in solution,  $f_{Li}$ . This relationship is linear, matching general 745 'batch reactor'-style behavior observed across the Amazon basin (Dellinger and others, 2015). 746 As our analysis is focused on hillslope processes, we do not simulate the scavenging of dissolved 747 Li by extant secondary phases as is theorized to occur in floodplains. The result is that we do not 748 capture significantly elevated  $\delta^7 \text{Li}_{\text{diss}}$  values (20-35‰), attributed to 'Rayleigh-style' floodplain 749 processes (fig. 3b), though transient overshoots in weathering incongruence during initial clay 750 precipitation capture similar style behavior.

751 Our relatively simple weathering mineralogy is also able to capture global ranges of riverine [Li<sup>+</sup>] from  $\sim 10^{0}$ - $10^{3}$  nM, though we note that simulated concentrations above 200 nM 752 753 require the inclusion of a soluble, Li-rich mineral such as chlorite or lithium-bearing alkali 754 amphibole. In particular, across simulations both with and without chlorite, the transition from 755 clay dissolution to primary mineral dissolution between high and medium weathering intensity environments captures the observed increase in [Li<sup>+</sup>] from  $\sim 10^{0}$  to  $\sim 10^{2}$  nM along with decreases 756 757 in  $\delta^7$ Lidiss values. Simulations with high chlorite-derived [Li<sup>+</sup>] are discussed in more detail in the 758 following section.

759 Our simulations also clarify the distinct controls of fluid travel time and mineral 760 residence time on Li systematics. Similar to previous reactive transport studies,  $\delta^7 \text{Li}_{\text{diss}}$  during 761 incongruent weathering increases with increasing fluid travel time (Wanner and others, 2014; 762 Bohlin and Bickle, 2019; Golla et al. 2021). We note, however, that our simulations reach 763 maximum  $\delta^7 \text{Li}_{\text{diss}}$  as incongruent weathering approaches a stoichiometric steady-state, while 764 solute concentrations continue to increase. This is occurs on much shorter timescales than those 765 required for the system to approach a meta-stable equilibrium between dissolving and 766 precipitating phases when net reaction rates approach zero and solute concentrations remain 767 constant. This travel time dependence broadly matches concentration-discharge behavior 768 observed in streams, in which high seasonal discharge periods associated with faster fluid travel 769 times exhibit lower [Li<sup>+</sup>] and  $\delta^7$ Li<sub>diss</sub> as compared to low discharge periods (Lemarchand and 770 others, 2010; Manaka and others, 2017). We demonstrate qualitative concentration-discharge

dynamics associated with our model output assuming a gamma-function travel time distributionin Appendix B.

773 In contrast, mineral residence times control the dominant weathering reactions, as 774 primary minerals are sequentially depleted from regolith with increasing mineral residence time. 775 This has previously been suggested to control the transition from primary mineral dissolution to 776 secondary mineral dissolution at high weathering intensities (Bouchez and others, 2013), and we 777 extend this framework to demonstrate potential transitions in dominant primary mineral reactions 778 and associated Li sources at low weathering intensities. We also note that mineral residence 779 times do not directly impact weathering congruence in our simulations except through these distinct transitions in weathering mineralogy. This is in contrast to previous studies in which 780 781 relative rates of primary mineral dissolution and secondary mineral precipitation are directly 782 dependent on mineral residence times (Chamberlain and others, 2005; Ferrier and Kirchner, 783 2008; Bouchez and others, 2013). While halloysite does not significantly accumulate under short 784 mineral residence times within our low WI simulations (fig. 5b), relative rates of halloysite 785 precipitation are still proportional to plagioclase dissolution integrated across the modeling 786 domain as enforced through the affinity-dependence terms in mineral precipitation/dissolution 787 rates (eq. 2). We offer a linkage between mineral residence time controls, fluid travel times and 788 associated weathering congruence by evaluating solute exports from the defined regolith zone 789 (gray shading in fig.'s 5 and 6), representing preferential flow through more porous regolith and 790 soil media relative to unweathered bedrock. This connection is implicit in previous studies, and 791 we discuss its incompatibility with observations of high [Li<sup>+</sup>] under low WI in the following 792 section.

793 CrunchFlow simulations also support the hypothesis that mineral residence times are 794 responsible for the broad trends between observed  $\delta^7 \text{Li}_{\text{sus}}$  and weathering intensity (Dellinger 795 and others, 2017; fig. 8). Specifically, at low weathering intensities when mineral residence 796 times are short,  $\delta^7 \text{Li}_{\text{sus}}$  largely represents contributions from primary minerals as interpreted by 797 Dellinger and others (2017). With increasing weathering intensity and the depletion of primary 798 minerals from the shallow, actively-eroding zone of regolith,  $\delta^7 Li_{sus}$  transitions to values 799 reflecting secondary minerals. We note that our simulations predict a much wider spread in 800  $\delta^7$ Li<sub>sus</sub> at high weathering intensities than observed. This primarily reflects the fact that low

(Li/Al)<sub>sec</sub>/(Li/Al)<sub>prim</sub> (i.e. *P<sub>Li</sub>*) are not observed at low weathering intensities (Dellinger and
others, 2017).

- 803
- 804

#### Weathering Intensity Controls on $\delta^7 \text{Li}_{\text{diss}}$

805

As described by both observations and theoretical considerations,  $\delta^7 \text{Li}_{\text{diss}}$  is related to WI 806 807 through a parabolic relationship (fig. 1, fig. 7e). Previous studies hypothesize that low  $\delta^7 Li_{diss}$ 808 values observed in low WI environments are caused by short fluid and mineral residence times 809 that inhibit secondary mineral precipitation relative to primary mineral dissolution, thereby 810 limiting the extent of Li fractionation (Bouchez and others, 2013; Dellinger and others, 2015). 811 Our CrunchFlow simulations demonstrate two potential issues with this hypothesis. First, the 812 treatment of weathering profiles within a multi-component RTM framework in which reaction 813 rates are based on solution chemistry suggests that there is no intrinsic connection between 814 mineral residence time and weathering congruence. Models that include this connection 815 implicitly assume that all weathering occurs within regolith, or mineral-depleted weathering 816 profiles, such that travel times are dependent on regolith thickness. However, this does not 817 account for infiltration of waters into relatively un-weathered bedrock, for example via fracture 818 flow. As shown in Appendix B, riverine  $\delta^7 Li_{diss}$  may reflect a lack of secondary mineral precipitation depending on the travel time distributions and mean travel times that dictate the 819 820 proportional contribution of these 'young', non-evolved waters (Kirchner, 2016; Jasechko and 821 others, 2016). However, for travel times to result in the observed  $\delta^7 \text{Li}_{\text{diss}}$ -WI relationships, young 822 water fractions would need to be highest in low WI environments. This connection between 823 travel times and erosional environments is an open question; however, we note that this is in 824 direct contrast to Jasechko and others (2016), who suggest relatively low proportions of 'young' 825 waters in high slope environments associated with low WI.

826 Second, short fluid travel times that limit secondary mineral precipitation also limit 827 primary mineral dissolution and overall reaction progress. The result is that under short fluid 828 travel times, simulated [Li<sup>+</sup>] is extremely low, on the order of <10 nM. This is in direct conflict 829 with observations that show the highest [Li<sup>+</sup>] in low WI and rapidly eroding environments, on 830 the order of  $10^3$  nM (Dellinger and others, 2015; Pogge von Strandmann and Henderson, 2015). 831 Fluid travel times, are therefore unable to capture observed parabolic relationships between

832  $\delta^7 \text{Lid_{iss}}$  and [Li<sup>+</sup>] as shown in figure 7. Given that observed low WI [Li<sup>+</sup>] on the order of 10<sup>3</sup> is 833 difficult to account for with typical granitic Li concentrations and plagioclase mineral solubility, 834 we suggest that low WI observations instead represent Li release from soluble, Li-rich minerals 835 such as chlorite.

836 Under this hypothesis, only low WI and short mineral residence times within the 837 weathering zone are able to maintain chlorite in significant concentrations. Our simulated 838 chlorite weathering profiles broadly match a recent comparison of 'vermiculitization' profiles 839 under high, medium, and low erosion rates in which protolith chlorite concentrations are 840 maintained in shallow soils only under high erosion rates (1-3 mm/yr; Gu and others, 2020). Due 841 to the high solubility and relatively high Li concentrations of chlorite, low  $\delta^7$ Lidiss results from 842 high dissolved Li concentrations that overwhelm uptake by secondary mineral precipitation. In 843 our simulations, the transition from low to moderate weathering intensity then results in a change 844 in the dominant Li weathering reaction from chlorite dissolution to plagioclase dissolution, 845 which is an inherently more incongruent weathering reaction for Li. This mechanism, in contrast 846 to fluid travel times, predicts parabolic relationships between  $\delta^7 \text{Li}_{\text{diss}}$  and both WI and [Li<sup>+</sup>] 847 consistent with observations, as shown in figure 7. We note that while our simulations are 848 limited to idealized representations of chlorite leaching, this overall pattern is characteristic of a 849 general situation in which Li exists at relatively higher concentrations in a more soluble mineral 850 within the bedrock assemblage such as lithium-bearing alkali amphiboles.

We also suggest that while our simulations only involve steady surface erosion rates, these dynamics should theoretically apply to environments in which transient mass wasting events occur as well. Periodic mass wasting events would expose fresh bedrock and allow for highly weatherable minerals to be maintained close to the surface.

855 With respect to riverine observations, low WI primarily represents the weathering of 856 shale (Dellinger and others, 2015; Beni River) and metamorphically-altered (Pogge von 857 Strandmann and Henderson, 2015; New Zealand) bedrock, which typically contain Li-rich, ferro-858 magnesian minerals such as chlorite. As mentioned above, the inclusion of chlorite brings our 859 bedrock Li concentrations in line with typical shale values (Holland, 1984) and bedrock values 860 observed in the Beni River watershed (Dellinger and others, 2015). Further, a study of Amazon 861 tributary sediments found high stream chlorite and illite contents (~30%) in the Andean Madeira 862 River (of which the Beni is a tributary) suggesting rapid erosion of soluble minerals (Gibbs,

863	1967). Additionally, chlorite presence decreases significantly along the Andean Ucayali river		
864	and is significantly lower in Amazon shield rivers (Irion, 1983; Gibbs, 1967), also in line with		
865	our hypothesized mineralogic controls. While we demonstrate the potential for chlorite		
866	weathering to explain observed relationships between weathering intensity, $\delta^7 Li_{diss}$ , and [Li <sup>+</sup> ],		
867	only very limited inferences have been drawn regarding chlorite contributions to Li weathering		
868	in actively eroding field environments (Golla et al. 2021). Thus, targeted studies are needed to		
869	evaluate whether or not chlorite or other highly soluble, Li-bearing minerals are responsible for		
870	high [Li <sup>+</sup> ] and low $\delta^7 Li_{diss}$ in low WI environments. Other mechanisms such as geothermal water		
871	contributions may also be consistent with the parabolic relationships in figure 7 e-f, as we		
872	describe in the following section.		
873			
874	Alternative Hypothesis: Geothermal Waters		
875			
876	Our analysis confirms that changes in weathering zone mineralogy as modulated by		
877	weathering intensity are consistent with global patterns of riverine lithium concentrations and		
878	isotopes; however, other mechanisms may also contribute to observed patterns. One possibility is		
879	that geothermal waters play an increasingly important role in Li fluxes under low weathering		
880	intensity conditions. A number of studies have observed high [Li <sup>+</sup> ] (10 <sup>-1</sup> -10 <sup>4</sup> $\mu$ M) often		
881	coincident with bedrock-like $\delta^7$ Lidiss values in geothermal springs (Millot and Negrel, 2007;		
882	Millot and others, 2007, 2010b, 2012; Bernal and others, 2014; Godfrey and Álvarez-Amado,		
883	2020). This has been attributed to high reaction temperatures that alter dominant clay stability to		
884	phases that incorporate less Li and reduce fractionation factors (Verney-Carron and others, 2015;		
885	Pogge von Strandmann and others, 2016). If relative contributions of geothermal waters are		
886	highest in low weathering intensity regimes, for example due to active tectonic uplift		
887	(Chamberlain and others, 2002), this could result in the parabolic $\delta^7$ Lidiss patterns of figure 7e-f:		
888	(1) At high weathering intensity, clay dissolution results in low [Li <sup>+</sup> ] and low $\delta^7$ Li <sub>diss</sub> ; (2) at		
889	moderate weathering intensity, where geothermal waters do not contribute significantly to solute		
890	fluxes, primary mineral dissolution coupled with secondary mineral precipitation results in		
891	moderate [Li <sup>+</sup> ] and high $\delta^7$ Li <sub>diss</sub> ; and (3) at low weathering intensity, where geothermal waters		
892	contribute significantly to solute fluxes, the inhibition of secondary mineral precipitation during		
893	high temperature weathering results in high [Li <sup>+</sup> ] and low $\delta^7$ Li <sub>diss</sub> . Geothermal water		

894	contributions in rapidly uplifting collisional orogens have been shown, for example, to control	
895	radiogenic Sr budgets and overall solute fluxes in Himalayan rivers (Evans and others, 2001). As	
896	with our chlorite hypothesis, this mechanism is consistent with riverine observations, and it is	
897	supported by observations of geothermal waters; however, this has not been systematically	
898	evaluated across WI gradients. Future studies targeting the potential role of geothermal waters	
899	and evaluating these competing hypotheses may shed important light on the mechanistic	
900	relationships between weathering environments and Li systematics. In particular, future studies	
901	are needed that target low WI environments including the detailed characterization of weathering	
902	zone mineralogy (e.g. the presence of soluble Li-rich minerals, mineralogical $P_{Li}$ values for	
903	secondary phases, and fluid travel times).	
904		
905	Limitations of Approach	
906		
907	One primary limitation of our methodology is that the partition coefficient $P_{Li}$ is set via	
908	mineral stoichiometries rather than evolving as a function of fluid chemistry. Mechanisms for the	
909	latter may include: competing clay phases, recrystallization/Ostwald ripening, sorption-	
910	desorption, and concentration-dependent trace element incorporation. In modeling Li uptake	
911	during clay formation, we chose a simplistic representation of halloysite Li-uptake such that	
912	mineral solubility was only dependent on Si and Al concentrations rather than base cation	
913	concentrations. This allows us to systematically vary $P_{Li}$ across the full 0-1 range through	
914	specific trace Li concentration without inducing time- or depth-dependent changes in $P_{Li}$ within	
915	individual simulations. However, it also limits the ability to forward-model or predict $\delta^7$ Li	
916	behavior at a specific site without local constraints on bedrock and secondary phase lithium	
917	concentrations. In particular, it will require extensive field characterization in order to accurately	
918	simulate the effects of evolving clay stability on Li isotope dynamics. Similarly, we do not	
919	include any dependence of instantaneous $P_{Li}$ on [Li <sup>+</sup> ], as has been noted experimentally	
920	(Decarreau and others, 2012) and included in other reactive transport frameworks (Wanner and	
921	others, 2014; Bohlin and Bickle, 2019). The incorporation of these dynamics would not change	
922	the relationship between $f_{Li}$ and $\delta^7$ Lidiss at steady-state (fig. 3b, filled circles) but would change	
923	the approach to steady state (fig. 3b, colored lines). Additionally, $P_{Li}$ dependence on [Li <sup>+</sup> ] may	
924	extend the reaction timescales over which $\delta^7 Li_{diss}$ increases. While this may allow for more	

realistic fluid travel times to support  $\delta^7$ Lidiss observations in low WI environments, it would still result in low [Li<sup>+</sup>], in conflict with observations.

927 Catchments are also complex 3-dimensional systems with variable distributions of fluid 928 travel times. As such, rivers may integrate a wide range of flow paths with different  $f_{Li}$  and resulting  $\delta^7$ Li<sub>diss</sub>, even in the absence of appreciable floodplain exchange. In addition, local 929 930 heterogeneity may impact the efficiency of reactions relative to that shown here for initially 931 homogeneous domains (Pandey and others, Jung and Navarre-Sitchler, 2018). In spite of the 932 large- and small-scale heterogeneities that exist in catchment and regolith profiles, respectively, 933 the general trends presented in the 1-D simulations here are still applicable (Maher, 2011; 934 Pandey and others, Jung and Navarre-Sitchler, 2018). Thus, although the simulations do not 935 capture full complexity of catchments and initial conditions, they do represent a broad spectrum 936 of flow rates, erosion rates and weathering rates and thus provide general insights into potential 937 behavior of Li isotopes.

- 938
- 939
- 940

#### CONCLUSIONS

941 In order to probe the relationships between weathering intensity and Li isotope 942 systematics, we incorporate Li isotope fractionation processes into CrunchFlow reactive 943 transport simulations of weathering granite. Using a parsimonious effective kinetic fractionation 944 parameterization, our simulations capture broad patterns in observed relationships between 945  $\delta^7 \text{Li}_{\text{diss}}$  and the fraction of dissolved Li remaining in solution following secondary mineral 946 precipitation. Upland weathering profiles are simulated across a range of weathering intensities 947 by varying water infiltration and bedrock uplift/erosion rates. Simulations produce a parabolic 948 relationship between  $\delta^7$ Lidiss and weathering intensity similar to global observations, controlled 949 by a combination of fluid travel times and mineral residence times in the weathering zone. 950 However, fluid travel and mineral residence time controls alone are unable to capture high 951 observed Li concentrations under low weathering intensity conditions: short fluid travel times 952 that limit clay precipitation also limit primary mineral dissolution and solute accumulation. 953 Instead, we demonstrate how mineral residence times may regulate the dominant Li weathering 954 reactions, where the weathering of soluble, Li-rich minerals may only be sustained with the rapid 955 supply of fresh bedrock. This mineralogic control on Li isotope systematics is able to capture

parabolic relationships between  $\delta^7$ Lidiss and both weathering intensity and Li concentrations. We suggest that as a viable hypothesis, this mineralogic control may be tested against other possible mechanistic ties between weathering regimes and Li systematics such as varying contributions from geothermal waters.

960 The refinement of a theoretical framework for Li isotope systematics may provide for 961 more accurate reconstructions of weathering dynamics in the past. In particular, direct links 962 between Li-specific congruence  $(P_{Li})$  and overall weathering congruence need to be established 963 in order for Li isotopes to serve as a direct proxy of weathering intensity. Additionally, based on 964 the parabolic relationships described above, tectonic uplift and attendant shifts in weathering may be result in either increasing or decreasing  $\delta^7 Li_{diss}$  values depending on the background 965 966 weathering intensity. Thus, understanding the mechanistic connections between uplift and  $\delta^7 Li_{diss}$ 967 may allow for more direct hypothesis evaluation, for example, through comparisons of proxy 968  $\delta^7$ Li with coeval records of hydrologic shifts or changes in depositional environment mineralogy 969 along active orogens that may indicate changes in fluid and mineral residence times.

970 971

972

980

982

#### ACKNOWLEDGEMENTS

We thank Daniel Ibarra, Jeremy Caves, and Friedhelm von Blanckenburg for useful comments
and conversations, along with the Associate Editor and two anonymous reviewers whose
comments and suggestions improved an earlier version of this manuscript. This work was
supported by the National Science Foundation (EAR-1254156) to K.M. The authors will make
updated CrunchFlow software source code and all input files used in this analysis available on
request.

981 Appendix A: Weathering Reaction Sensitivity Analysis.

To demonstrate the sensitivity of the presented simulations to assumptions of prescribed P<sub>Li</sub>
values in Figure 8 and to our idealized reaction network, we present analyses showing:

- 986 *A1*) How  $\delta^7$ Lidiss varies as a function of weathering intensity and [Li] for a range of  $P_{Li}$ 987 values, and
- 988A2) How the use of a 2:1 clay rather than 1:1 halloysite does not impact apparent989fractionations.
- 990
- 991 A.1 To show the sensitivity of  $\delta^7$ Lidiss to weathering intensity and [Li], we present an altered
- 992 version of Figure 7e,f in which simulations are color-coded by prescribed  $P_{Li}(0.08, 0.53, \text{ and}$
- 993 0.99). As  $P_{Li}$  primarily impacts the maximum extent of  $\delta^7 \text{Lid}_{\text{id}\text{iss}}$  increase during incongruent
- 994 plagioclase weathering, the effect of lowering  $P_{Li}$  is to flatten the parabolic curves of  $\delta^7$ Lidiss v.

995 WI and  $\delta^7 \text{Lid}_{iss}$  v. [Li]. In other words, smaller  $P_{Li}$  values result in lower ranges of  $\delta^7 \text{Lid}_{iss}$ 996 variability. Additionally,  $P_{Li}$  values impact the concentrations of Li during incongruent and clay 997 weathering along with  $\delta^7 \text{Li}_{clay}$  values. This results in variable  $\delta^7 \text{Lid}_{iss}$  at high weathering intensity 998 when clay dissolution dominates and  $\delta^7 \text{Lid}_{iss}$  matches  $\delta^7 \text{Lid}_{clay}$  (figure A1a). It also results in [Li<sup>+</sup>] 999 variability under both clay dissolution and incongruent weathering: as  $P_{Li}$  approaches 1, the

1000 variability under bour clay dissolution and incongruent weathering: as  $P_{Li}$  approaches 1, the difference in [Li<sup>+</sup>] under clay dissolution v. incongruent weathering decreases, causing a steeper

1001 left side of the parabolic curve in figure A1b.

1002

1003 A.2 - To demonstrate the insensitivity of our model simulations to alternative representations of 1004 clay mineralogy, we include batch reactor simulations similar to those presented in the main text 1005 where halloysite is replaced with a calcium beidellite (2:1 smectite group clay). Kinetic 1006 fractionation is implemented in the same manner, and we simulate a range of  $P_{Li}$  (0.08 – 0.93) 1007 values based on prescribed substitution of Li<sup>+</sup> for Ca<sup>2+</sup> as shown in table A1. We chose to 1008 simulate a calcium rather than iron or magnesium 2:1 clay based on the lack of magnesium and 1009 iron in our primary mineral dissolution reaction (Plagioclase An20). Selected results 1010 corresponding to main text figures 2a,b,d and 3a are shown in figure A2. In comparison to our

1011 halloysite reaction, the simulation of beidellite alters the patterns of primary mineral dissolution

and secondary mineral precipitation rates along with the steady-state weathering incongruence

value representing stoichiometric balance of plagioclase dissolution and beidellite precipitation.
 Changes in the balance of rates are largely related to changes in reaction affinity as kinetic rate

1015 constants are unchanged from halloysite simulations. Weathering incongruence reflects

1016 represents the aluminum-normalized stoichiometry. As shown in figure A2, the evolution of

1017  $\delta^7$ Lidiss is identical to halloysite both as a function of time and of prescribed  $P_{Li}$ . Even though the 1018 model is simulating a different mechanisms of Li substitution (i.e. for substituted cations rather

1019 than structural aluminum), the  $P_{Li}$  value and kinetic fractionation factor together set  $\delta^7 \text{Lid}_{\text{iss}}$  in the

same manner as with structural substation for Al in halloysite.

1021

1022 Appendix B: Theoretical concentration-discharge dynamics.1023

In order to explore potential concentration-discharge dynamics encapsulated in our model simulations, we convolve idealized watershed travel time distributions with time-varying output from main text batch reactor simulations. Numerous studies have shown that times of high flow, either in response to transient infiltration events such as storms and snowmelt or sustained seasonal changes in precipitation, are associated with relatively lower mean travel times. To show how this may be manifested in stream solute data, we assume watershed travel time distributions may be represented by a gamma-type distribution as,

1031

1032 
$$h(\tau) = \frac{\tau^{\alpha-1}}{\left(\frac{\tau_m}{\alpha}\right)^{\alpha} \Gamma(\alpha)} e^{-\tau \alpha/\tau_m},$$

1033

1034 where *h* is a weight for a given travel time  $\tau$ ,  $\alpha$  represents a shape factor, and  $\tau_m$  is the mean 1035 travel time (e.g. Kirchner, 2016).

1036

1037 For the purposes of this exercise, we assume a shape factor of 1. We convolve travel time

1038 distributions with batch reactor experiment  $\delta^7 \text{Li}_{\text{diss}}$  for a range of mean travel time values by

1039 taking the weighted mean of Li concentration-weighted, time-dependent  $\delta^7$ Li<sub>diss</sub> values using

1040	h( $\tau$ )*[Li]( $\tau$ ) as the weights. Figure B1 shows how the convolved $\delta^7$ Lidiss varies in response to		
1041	mean travel times. As mean travel time decreases, the associated $\delta$ /Li <sub>diss</sub> decreases as well. This		
1042	relationship is muted relative to the time v. $\delta^7 Li_{diss}$ curves shown in Figure 3, due to the mixing		
1043	across a wide range of travel times as discussed for solutes (e.g. Maher, 2011) and stable isotope		
1044	ratios (Druhan & Maher, 2017) in previous papers. The largest changes are seen when mean		
1045	travel times are in the vicinity of 1-3 months corresponding roughly to timescales of		
1046	stoichiometric steady-state in the plagioclase dissolution – halloysite precipitation reaction		
1047	series. Under these idealized scenarios, an increase in flow resulting in a reduction in mean travel		
1048	time would cause a subsequent decrease in $\delta^7 L_{idiss}$ values consistent with field observations from		
1049	a number of studies. Importantly, while the magnitude of $\delta^7 L_{\text{disc}}$ change is dependent on the shift		
1050	and absolute value of mean travel time between low and high flow along with the shape factor		
1050	the direction of change is not. The precise relationship between mean travel time and discharge is		
1051	net well characterized; however, interfacing this stable isotone repetive transport model with		
1052	hot wen enalderenzed, nowever, interfacing uns stable isotope reactive transport model with		
1055	nilisiope or watersned hydrologic models may facilitate concentration-discharge relationship		
1054	predictions.		
1055			
1056	REFERENCES		
1057			
1058	Anghel, I., Turin, H. J., Reimus, P. W., 2002, Lithium sorption to Yucca Mountain tuffs: Applied		
1059	Geochemistry, v. 17, p. 819-824.		
1060			
1061	Arnorsson, S., Stefansson, A., 1999, Assessment of feldspar solubility constants in water		
1062	in the range 0 degrees to 350 degrees C at vapor saturation pressures: American Journal of		
1063	Science v. 299, p. 173–209.		
1064			
1065	Bagard, ML., West, J. A., Newman, K., Basu, A. R., 2015, Lithium isotope fractionation in the		
1066	Ganges-Brahmaputra floodplain and implications for groundwater impact on seawater isotopic		
1067	composition: Earth and Planetary Science Letters, v. 432, p. 404-414.		
1068			
1069	Bernal, N. F., Gleeson, S. A., Dean, A. S., Liu, X. M., Hoskin, P., 2014. The source of halogens		
1070	in geothermal fluids from the Taupo Volcanic Zone. North Island, New Zealand: Geochimica et		
1071	Cosmochimica Acta v 126 n 265-283		
1072	Cobino en intera, (*. 120, p. 200–200.		
1073	von Blanckenburg F. Hewawasam T. Kubik P. W. 2004 Cosmogenic nuclide evidence for		
1074	low weathering and denudation in the wet tropical highlands of Sri I anka: Journal of		
1074	Goophysical Poscarch: Earth Surface y 100 E02008 22 pp		
1075	Geophysical Research. Earth Surface, V. 109, 105008, 22 pp.		
1070	Deblin M.S. and Diable M.J. 2010. The reactive transment of Ling a manitor of weathering		
10//	Bonlin, M. S. and Bickle, M. J., 2019, The reactive transport of L1 as a monitor of weathering		
10/8	processes in kinetically limited weathering regimes: Earth and Planetary Science Letters, v. 511,		
10/9	p. 233-243.		
1080			
1081	Bouchez, J., von Blanckenburg, F., Schuessler, J. A., 2013, Modeling novel stable isotope ratios		
1082	in the weathering zone: American Journal of Science, v. 313, p. 267-308.		
1083			
1084	Caves-Rugenstein, J. K., Ibarra, D. E., von Blanckenburg, F., 2019, Neogene cooling driven by		
1085	land surface reactivity rather than increased weathering fluxes: Nature, v. 571, p. 99-102.		

1086 1087 Chamberlain, C. P., Koons, P. O., Meltzer, A. S., Park, S. K., Craw, D., Zeitler, P., and Poage, 1088 M. A., 2002, Overview of hydrothermal activity associated with active orogenesis and 1089 metamorphism: Nanga Parbat, Pakistan Himalava: American Journal of Science, v. 302(8), p. 1090 726-748. 1091 1092 Chamberlain, C. P., Waldbauer, J. R., Jacobson, A. D., 2005, Strontium, hydrothermal systems 1093 and steady-state chemical weathering in active mountain belts: Earth and Planetary Science 1094 Letters, v. 238, p. 351-366. 1095 1096 Chan, L. H., Edmond, J. M., Thompson, G., Gillis, K., 1992, Lithium isotopic composition of 1097 submarine basalts: implications for the lithium cycle in the oceans: Earth and Planetary Science 1098 Letters, v. 108, p. 151-160. 1099 1100 Decarreau, A., Vigier, N., Pálková, H., Petit, S., Vieillard, P., Fontaine, C., 2012, Partitioning of 1101 lithium between smectite and solution: An experimental approach: Geochimica et Cosmochimica 1102 Acta, v. 85, p. 314-325. 1103 1104 Dellinger, M., Gaillardet, J., Bouchez, J., Calmels, D., Galy, V., Hilton, R. G., Louvat, P., 1105 France-Lanord, C., 2014, Lithium isotopes in large rivers reveal the cannibalistic nature of 1106 modern continental weathering and erosion: Earth and Planetary Science Letters, v. 401, p. 359-1107 372. 1108 1109 Dellinger, M., Gaillardet, J., Bouchez, J., Calmels, D., Louvat, P., Dosseto, A., Gorge, C., 1110 Alanoca, L., Maurice, L., 2015, Riverine Li isotope fractionation in the Amazon River basin 1111 controlled by the weathering regimes: Geochimica et Cosmochimica Acta, v. 164, p. 71-93. 1112 1113 Dellinger, M., Bouchez, J., Gaillardet, J., Faure, L., Moureau, J., 2017, Tracing weathering 1114 regimes using the lithium isotope composition of detrital sediments: Geology, v. 45, p. 411-414. 1115 1116 Druhan, J. L., Steefel, C. I., Williams, K. H., DePaolo, D. J., 2013, Calcium isotope fractionation 1117 in groundwater: Molecular scale processes influencing field scale behavior: Geochimica et 1118 Cosmochimica Acta, v. 119, p. 93-116. 1119 1120 Druhan, J. L., & Maher, K., 2017, The influence of mixing on stable isotope ratios in porous 1121 media: A revised Rayleigh model: Water Resources Research, v. 53(2), p. 1101-1124. 1122 1123 Evans, M. J., Derry, L. A., Anderson, S. P., France-Lanord, C., 2001, Hydrothermal source of 1124 radiogenic Sr to Himalavan rivers: Geology, v. 29(9), p. 803-806. 1125 1126 Ferrier, K. L., Kirchner, J. W., 2008, Effects of physical erosion on chemical denudation rates: A 1127 numerical modeling study of soil-mantled hillslopes: Earth and Planetary Science Letters, v. 272, 1128 p. 591-599. 1129 1130 Flesch, G. D., Anderson Jr., A. R., Svec, H. J., 1973, A secondary isotopic standard for <sup>6</sup>Li/<sup>7</sup>Li

1131 determinations: International Journal of Mass Spectrometry Ion Physics, v. 12(3), p. 2655-272.

1132 1133 Gibbs, R. J., 1967, The geochemistry of the Amazon River system: Part I. The factors that 1134 control the salinity and the composition and concentration of the suspended solids: Geological 1135 Society of America Bulletin, v. 78, p. 1203-1232. 1136 1137 Godfrey, L., Álvarez-Amado, F., 2020, Volcanic and saline lithium inputs to the Salar de 1138 Atacam: Minerals, v. 10, pp. 17. 1139 1140 Golla, J. K., Kuessner, M. L., Henehan, M. J., Bouchez, J., Rempe, D. M., & Druhan, J. L., 2021, 1141 The evolution of lithium isotope signatures in fluids draining actively weathering 1142 hillslopes: Earth and Planetary Science Letters, v. 567, p. 116988. 1143 1144 Gu, X., Rempe, D. M., Dietrich, W. E., West, A. J., Lin, T.-C., Jin, L., Brantley, S. L., 2020, 1145 Chemical reactions, porosity, and microfracturing in shale during weathering: The effect of 1146 erosion rate: Geochimica et Cosmochimica Acta, v. 269, p. 63-100. 1147 Heidari, P., Li, L., Jin, L., Williams, J. Z., Brantley, S. L., 2017, A reactive transport model for 1148 1149 Marcellus shale weathering: Geochimica et Cosmochimica Acta, v. 217, p. 421-440. 1150 1151 Hindshaw, R. S., Tosca, R., Goût, T. L., Farnan, I., Tosca, N. J., Tipper, E. T., 2019, 1152 Experimental constraints on Li isotope fractionation during clay formation. Geochimica et 1153 Cosmochimica Acta, v. 250, p. 219-237. 1154 1155 Holland H., 1984, The chemical evolution of the atmosphere and oceans: Princeton University 1156 Press. 1157 Huh, Y., Chan, L.-H., Edmond, J. M., 2001, Lithium isotopes as a probe of weathering 1158 1159 processes: Orinoco River: Earth and Planetary Science Letters, v. 194, p. 189-199. 1160 1161 Irion, G., 1983, Clay mineralogy of the suspended load of the Amazon and of rivers in the 1162 Papua-New Guinea mainland: Mitteilungen Geologie und Palaontolgie Institut Unversitat 1163 Hamburg, v. 55, p. 483-504. 1164 1165 Jasechko, S., Kirchner, J. W., Welker, J. M., McDonnell, J. J., 2016, Substantial proportion of 1166 global streamflow less than three months old: Nature Geoscience, v. 9, p. 126-129. 1167 1168 Jin, L., Ravella, R., Ketchum, B., Bierman, P. R., Heaney, P., White, T., Brantley, S. L., 2010, 1169 Mineral weathering and elemental transport during hillslope evolution at the Susquehanna/Shale 1170 Hills Critical Zone Observatory: Geochimica et Cosmochimica Acta, v. 74(13), p. 3669-3691. 1171 1172 Jung, H. and Navarre-Sitchler, A., 2018, Physical heterogeneity control on effective mineral 1173 dissolution rates: Geochimica et Cosmochimica Acta, v. 227, p. 246-263. 1174 1175 Kirchner, J.W., 2016, Aggregation in environmental systems - Part 1: Seasonal tracer cycles quantify young water fractions, but not mean transit times, in spatially heterogeneous 1176

1177 catchments: Hydrology and Earth System Sciences, v. 20, p. 279-297.

1178

- 1179 Lara, M.C., Buss, H., Henehan, M.J., Schuessler, J.A., McDowell, W.H., 2022, Secondary
- minerals drive extreme lithium isotope fractionation during tropical weathering: JGR Earth
  Surface, v. 127, e2021JF006366.
- 1182
- Lasaga, A.C., 1998, Kinetic theory in the earth sciences: v. 811. Princeton, NJ: Princetonuniversity press, 1998.
- 1185
- Lebedeva, M. I., Fletcher, R. C., Brantley, S. L., 2010, A mathematical model for steady-state
  regolith production at constant erosion rate: Earth Surface Processes and Landforms, v. 35, p.
  508-524.
- 1189

1190 Lebedeva, M. I. and Brantley, S. L., 2013, Exploring geochemical controls on weathering and 1191 erosion of convex hillslopes: beyond the empirical regolith production function: Earth Surface

1192 Processes and Landforms, v. 38(15), p. 1793-1807.

- 1193
  1194 Lechler, M., Pogge von Strandmann, P. A. E., Jenkyns, H. C., Prosser, G., Parente, M., 2015,
  1195 Lithium-isotope evidence for enhanced silicate weathering during OAE 1a ,Early Aptian Selli
  1196 event: Earth and Planetary Science Letters, v. 432, p. 210-222.
- event: Earth and Planetary Science Letters, v. 432, p. 210-222.
- Lemarchand, E., Chabaux, R., Vigier, N., Millot, R., Pierret, M.-C., 2010, Lithium isotope
  systematics in a forested granitic catchment, Strengbach, Vosges Mountains, France:
  Geochimica et Cosmochimica Acta, v. 74(16), p. 4612-4628.
- Li, G. and West, J. A., 2014, Evolution of Cenozoic seawater lithium isotopes: Coupling of
  global denudation regime and shifting seawater sinks: Earth and Planetary Science Letters, v.
  401, p. 284-293.
- Lowson, R. T., Comarmond, M.-C. J., Rajaratnam, G., Brown, P. L., 2005, The kinetics of the
  dissolution of chlorite as a function of pH and at 25°C: Geochimica et Cosmochimica Acta, v.
  69(7), p. 1687-1699.
- 1210 Maffre, P., Godderis, Y., Vigier, N., Moquet, J.-S., Carretier, S., 2020, Modelling the riverine 1211  $\delta^7$ Li variability throughout the Amazon Basin: Chemical Geology, v. 532, 119336.
- 1212
- Maher, K., Steefel, C. I., DePaolo, D. J., & Viani, B. E., 2006, The mineral dissolution rate conundrum: Insights from reactive transport modeling of U isotopes and pore fluid chemistry in marine sediments: Geochimica et Cosmochimica Acta, v. 70(2), p. 337-363.
- 1215
- Maher, K., 2010, The dependence of chemical weathering rates on fluid residence time: Earthand Planetary Science Letters, v. 294, p. 101-110.
- 1219
- 1220 Maher, K., 2011, The role of fluid residence time and topographic scales in determining
- 1221 chemical fluxes from landscapes: Earth and Planetary Sciences, v. 312, p. 48-58.
- 1222

Maher, K., Chamberlain, C. P., 2014, Hydrologic regulation of chemical weathering and the

1223

1224 geologic carbon cycle: Science, v. 343, p. 1502-1504. 1225 1226 Maher, K., von Blanckenburg, F., 2016, Surface ages and weathering rates from <sup>10</sup>Be ,meteoric, 1227 and <sup>10</sup>Be/<sup>9</sup>Be: Insights from differential mass balance and reactive transport modeling: Chemical 1228 Geology, v. 446, p. 70-86. 1229 1230 Manaka, T., Araoka, D., Yoshimura, T., Hossain, H. M. Z., Nishio, Y., Suzuki, A., Kawahata, 1231 H., 2017, Downstream and seasonal changes of lithium isotope ratios in the Ganges-1232 Brahmaputra river system: Geochemistry, Geophysics, Geosystems, v. 18(8), p. 3003-3015. 1233 1234 Maxwell, R. M., Condon, L. E., Kollet, S. J., Maher, K., Haggerty, R., Forrester, M. M., 2015, 1235 The imprint of climate and geology on the residence times of groundwater: Geophysical 1236 Research Letters, v. 43(2), p. 701-708. 1237 1238 Millot, R., Girard, J. P., 2007, Lithium isotope fractionation during adsorption onto mineral 1239 surfaces: Clay in Natural and Engineered Barriers for Radioactive Waste Confinement, 3rd 1240 International Meeting – Lille. 1241 Millot, R., Negrel, P., 2007, Multi-isotopic tracing  $\delta^7 \text{Li}$ ,  $\delta^{11}\text{B}$ ,  ${}^{87}\text{Sr}/{}^{86}\text{Sr}$ , and chemical 1242 1243 geothermometry: evidence from hydro-geothermal systems in France: Chemical Geology, v. 1244 244, p. 664-678. 1245 1246 Millot, R., Negrel, P., Petelet-Giraud, E., 2007, Multi-isotopic Li, B, Sr, Nd, approach for 1247 geothermal reservoir characterization in the Limagne Basin, Massif Central, France: Applied 1248 Geochemistry, v. 22, p. 2307-2335. 1249 1250 Millot, R., Vigier, N., Gaillardet, J., 2010a, Behavior of lithium and its isotopes during 1251 weathering in the Mackenzie Basin, Canada: Geochimica et Cosmochimica Acta, v. 74, p. 3897-1252 3912. 1253 1254 Millot, R., Scaillet, B., Sanjuan, B., 2010b, Lithium isotopes in arc geothermal systems: 1255 Guadeloupe, Martinique, French West Indies, and experimental approach: Geochimica et 1256 Cosmochimica Acta, v. 74, p. 1852-1871. 1257 1258 Millot, R., Hegan, A., Negrel P., 2012, Geothermal waters from the Taupo Volcanic Zone, New 1259 Zealand: Li, B, and Sr isotopes characterization: Applications in Geochemistry, v. 27, p. 677-1260 688. 1261 1262 Misra, S., Froelich, P. N., 2012, Lithium isotope history of Cenozoic seawater: changes in 1263 silicate weathering and reverse weathering: Science, v. 335, p. 818-823. 1264 1265 Murphy, S. F., Brantley, S. L., Blum, A. E., White, A. F., Dong, H., 1998, Chemical weathering 1266 in a tropical watershed, Luquillo Mountains, Puerto Rico; II. Rate and mechanism of biotite 1267 weathering: Geochimica et Cosmochimica Acta, v. 62(2), p. 227-243. 1268

1269 1270 1271	Palandri, J. L. and Kharaka, Y. K., 2004, A compilation of rate parameters of water-mineral interaction kinetics for application to geochemical modeling Rep. 2004–1068: US Geological Survey Open File Report.
1272	
1272 1273 1274 1275 1276	Pandey, S. and Rajaram, H., 2016, Modeling the influence of preferential flow on the spatial variability and time-dependence of mineral weathering rates: Water Resources Research, v. 52(12), p. 9344-9366.
1270	
1277	Penniston-Dorland, S., Liu, XM., Rudnick, R. L., 2017, Lithium Isotope Geochemistry:
1278 1279	Reviews in Mineralogy and Geochemistry, v. 82, p. 165-217.
1280	Pistingr I.S. and Handarson G.M. 2003 Lithium isotona fractionation during continental
1200	ristilici, J. S., and Hendelson, O. M., 2005, Entituti-isotope fractionation during continental
1281	weathering process: Earth and Planetary Science Letters, v. 214, n. 1–2, p. 327–339.
1282	
1283	Pogge von Strandmann P A E Henderson G M 2015 The Li isotone response to mountain
1200	unlift: Goology y 12 n 67 70
1204	upint. Ocology, v. 45, p. 07-70.
1285	
1286	Pogge von Strandmann, P. A. E., Jenkyns, H. C., Woodfine, R. G., 2013, Lithium isotope
1287	evidence for enhanced weathering during Oceanic Anoxic Event 2: Nature Geoscience, v. 6, p.
1288	668-672.
1289	
1290	Pogge von Strandmann P A F Burton K W Onfergelt S Firiksdottir F S Murnhy M I
1201	Financeson A Cisloson S P 2016 The effect of hydrothermal spring weathering processes and
1291	minimum and distinguity on lithium isotonogy Lake Mayzeta Lasland, Chamical Coology v. 445 a.
1292	primary productivity on itinium isotopes. Lake Myvain, iceland: Chemical Geology, v. 445, p. 4-
1293	13.
1294	
1295	Pogge von Strandmann, P. A. E., Desrochers, A., Murphy, M. J., Finlay, A. J., Selby, D., Lenton,
1296	T. M., 2017, Global climate stabilisation by chemical weathering during the Hirnantian
1297	glaciation: Geochemical Perspectives Letters, v. 3, p. 230-237.
1298	$\mathcal{S}$
1290	Pogge von Strandmann P & F Kasemann S & Wimnenny I B 2020 Lithium and lithium
1200	isotonos in Earth's surface surface Elements y 16(4) n 252 259
1201	isotopes in Earth's surface cycles. Elements, v. 10(4), p. 255-258.
1301	
1302	Proust, D., Eymery, JP., Beaufort, D., 1986, Supergene vermiculitization of a magnesian
1303	chlorite: Iron and magnesium removal processes: Clays and Clay Minerals, v. 34, p. 572-580.
1304	
1305	Steefel C I C A J Appelo B Arora D Jacques T Kalbacher O Kolditz V Lagneau P C
1306	Lichtner K II Mayer I C I Meeussen S Molins D Moulton H Shao I Simunek N
1207	Snuchar S. P. Vahusaki and G. T. Vah. 2015. Pasativa transport and a for subsurface
1200	Spycher, S. B. Labusaki and O. T. Tell, 2015, Reactive transport codes for substitute $\frac{1}{10}$
1308	environmental simulation. Computational Geosciences, V. 19(3), p. 445-478.
1309	
1310	Sullivan, P. L., Hynek, S. A., Gu, X., Singha, K., White, T., West, N., Kim, H., Clarke, B.,
1311	Kirby, E., Duffy, C., Brantley, S. L., 2016, Oxidative dissolution under the channel leads
1312	geomorphological evolution at the Shale Hills catchment: American Journal of Science. v. 316.
1313	p 981-1026
1314	r
1.7 1 1	

- 1315 Sun, H., Xiao, Y., Gao, Y., Zhang, G., Casey, J. F., Shen, Y., 2018, Rapid enhancement of
- 1316 chemical weathering recorded by extremely light seawater lithium isotopes at the Permian-
- 1317 Triassic boundary: Proceedings of the National Academy of Sciences, v. 115(15), p. 3782-3787.
- 1318
- Tardy, Y., Krempp, G., Trauth, N., 1972, Le lithium dans les minéraux argileux des sédiments et
  des sols: Geochmica et Cosmochimica Acta, v. 36, p. 397-412.
- 13211322 Templeton, A. S., Chamberlain, C. P., Koons, P. O., and Craw, D., 1998, Stable isotopic
- 1323 evidence for mixing between metamorphic fluids and surface-derived waters during recent uplift
- 1324 of the Southern Alps, New Zealand: Earth and Planetary Science Letters, v. 154, p. 73-92.
- 1325
- Teng, F. Z., McDonough, W. F., Rudnick, R. L., Dalpe, C., Tomascak, P. B., Chappell, B. W.,
  Gao, S., 2004, Lithium isotopic composition and concentration of the upper continental crust:
- 1328 Geochimica et Cosmochimica Acta, v. 68, p. 4167-4178.
- 1329
- Tomascak, P. B., 2004, Developments in the understanding and application of lithium isotopes in
  the earth and planetary sciences in geochemistry of non-traditional stable isotopes: Reviews in
  Mineralogy and Geochemistry, v. 55, p. 153-195.
- 1333
- 1334 Verney-Carron, A., Vigier, N., Millot, R., Hardarson, B. S., 2015, Lithium isotopes in
- hydrothermally altered basalts from Hengill, SW Iceland: Earth and Planetary Science Letters, v.411, p. 62-71.
- 1337
- 1338 Vigier, N., Decarreau, A., Millot, R., Carignan, J., Petit, S., France-Lanord, C., 2008,
- Quantifying Li isotope fractionation during smectite formation and implications for the Li cycle:
  Geochimica et Cosmochimica Acta, v. 72, p. 780-792.
- 1341
- Vigier, N., Gislason, S., Burton, K., Millot, R., Mokadem, F., 2009, The relationship between
  riverine lithium isotope composition and silicate weathering rates in Iceland: Earth and Planetary
  Science Letters, v. 287, p. 434-441.
- 1345
- Wang, Y., Xu, H., 2001, Prediction of trace metal partitioning between minerals and aqueous
  solutions: a linear free energy correlation approach: Geochimica et Cosmochimica Acta,
  v. 65, p. 1529-1543.
- 1348 1349
- 1350Wanner, C., Sonnenthal, E. L., Liu, X.-M., 2014, Seawater  $\delta^7$ Li: A direct proxy for global CO21351consumption by continental silicate weathering?: Chemical Geology, v. 381, p. 154-167.
- 1352
- 1353 Williams, L. B., Hervig, R. L., 2005, Lithium and boron isotopes in illite-smectite: the
- 1354 importance of crystal size: Geochimica et Cosmochimica Acta, v. 24, p. 5705-5716.
- 1355
- 1356 Wimpenny J., Gisalason, S. R., James, R. H., Gannoun, A., Pogge von Strandmann, P. A. E.,
- 1357 Burton, K. W., 2010, The behavior of Li and Mg isotopes during primary phase dissolution and
- 1358 secondary mineral formation in basalt: Geochimica et Cosmochimica Acta, v. 74, 5259-5279.
- 1359

1360 Winnick, M. J., Maher, K., 2018, Relationships between CO<sub>2</sub>, thermodynamic limits on silicate 1361 weathering, and the strength of the silicate weathering feedback: Earth and Planetary Science 1362 Letters, v. 485, p. 111-120. 1363 1364 Wolery, T. J., Jackson, K. J., Bourcier, W. L., Bruton, C. J., Viani, B. E., Knauss, K. G., Delany, 1365 J. M., 1990, Current status of the EQ3/6 software package for geochemical modeling: ACS 1366 Symposium Series, v. 416, p. 104–116. 1367 1368 Zhang, L., Chan, L. H., Gieskes, J. M., 1998, Lithium isotope geochemistry of pore waters from 1369 Ocean Drilling Program Sites 918 and 919, Irminger Basin: Geochimica et Cosmochimica Acta 1370 v. 62, p. 2437-2450. 1371 1372 Zhang, X., Saldi, G. D., Schott, J., Bouchez, J., Kuessner, M. Montouillout, V., Henehan, M., 1373 Gailardet, J., 2021, Experimental constraints on Li isotope fractionation during the interaction 1374 between kaolinite and seawater. Geochimica et Cosmochimica Acta v. 292, p. 333-347. 1375 1376 Zhu, C., 2005, In situ feldspar dissolution rates in an aquifer: Geochimica et Cosmochimica 1377 Acta, v. 69(6), p. 1435-1453. 1378 1379 1380 FIGURE CAPTIONS 1381 1382 Table 1. CrunchFlow Reaction Network. 1383 1384 Table 2. Weathering simulation scenarios. 1385 1386 Table 3. Variables calculated from model output. 1387 1388 Fig. 1. Conceptual diagram of lithium isotope systematics based on observational and theoretical 1389 studies (Bouchez and others, 2013; Dellinger and others, 2015). The top graph shows general 1390 patterns of  $\delta^7 Li_{diss}$  as a function of weathering intensity (WI), soil residence time, and dissolved 1391 Li concentrations. Points A, B, and C represent medium, high, and low WI, respectively, and are 1392 described in the Introduction. The bottom panel shows relative rates of regolith production, 1393 physical erosion (E), and chemical weathering (W), with increasing WI (left to right), along with 1394 depictions of idealized regolith profiles showing the weathering of primary minerals (gray) and 1395 accumulation of secondary minerals (brown). 1396 1397 Fig. 2. Batch reactor (Scenario 1) results for granitic weathering (no chlorite) showing the 1398 temporal evolution of (a) plagioclase and halloysite dissolution/precipitation rates; (b) 1399 weathering incongruence calculated as the ratio of hallovsite precipitation to plagioclase dissolution rates; (c) dissolved  $Ca^{2+}$  concentration representing fully congruent weathering; and 1400 1401 (d) dissolved Li<sup>+</sup> concentrations across a range of partitioning coefficients (colored lines). 1402 1403 Fig. 3. Lithium isotope behavior in batch reactor experiments. (A) Temporal evolution of  $\delta^7 Li_{diss}$ for a range of partition coefficients (as in Fig. 2D); (B)  $\delta^7$ Lidiss as a function of fraction Li left in 1404

solution. Amazon basin data from Dellinger and others (2015) shown in gray, and samples

interpreted to reflect floodplain uptake processes denoted.

1405

1407				
1408	Fig. 4. No Erosion (Scenario 2) CrunchFlow simulations of weathering profiles showing (a)			
1409	plagioclase solid phase (vol. %); (b) halloysite solid phase (vol. %); (c) weathering incongruence			
1410	(ratio of instantaneous halloysite precipitation to plagioclase dissolution rates); (d) dissolved			
1411	$[Ca^{2+}]$ , (e) dissolved $[Li^+]$ , and (d) $\delta^7 Li_{diss}$ . Simulation timesteps are denoted by line color.			
1412	Simulations depicted have a $P_{Li}$ value of 0.9.			
1413				
1414	Fig. 5. Crunchflow simulations of granitic weathering (Scenario 3) including uplift and erosion.			
1415	Weathering profile results for low (a-e), medium (f-j), and high (k-o) weathering intensity			
1416	conditions. Columns from left to right show solid phase plagioclase, solid phase halloysite,			
1417	weathering incongruence, dissolved [Li <sup>+</sup> ], and $\delta^7 Li_{diss}$ . Gray shading denotes un-weathered			
1418	bedrock defined as model cells with <101% initial porosity. Colored lines denote model			
1419	timestep. Red bars in d, i, and n denote approximate observational ranges of [Li <sup>+</sup> ] from Dellinger			
1420	and others (2015) for low, medium, and high weathering intensity, respectively. Simulations			
1421	depicted have a $P_{Li}$ value of 0.9.			
1422				
1423	Fig. 6. Crunchflow simulations including chlorite (Scenario 4) and vermiculite minerals from			
1424	Table 1. Weathering profile results for low (a-d), medium (e-h), and high (i-l) weathering			
1425	intensity conditions. Columns from left to right show solid phase plagioclase, solid phase			
1426	chlorite, dissolved [Li <sup>+</sup> ], and $\delta^7$ Lidiss. Gray shading denotes un-weathered bedrock defined as			
1427	model cells with <101% initial porosity. Colored lines denote model timestep. Red bars in c, g,			
1428	and k denote approximate observational ranges of [Li <sup>+</sup> ] from Dellinger and others (2015) for			
1429	low, medium, and high weathering intensity, respectively. Simulations depicted have a $P_{Li}$ value			
1430	of 0.9.			
1431				
1432	Fig. 7. Relationships between $\delta^7$ Li <sub>diss</sub> and Weathering Intensity (left) and dissolved Li			
1433	concentrations (right). Results show model output (a-b) without chlorite; (c-d) with chlorite; and			
1434	(e-f) global riverine observations from Dellinger and others (2015) and references therein.			
1435	Simulations depicted have a $P_{Li}$ value of 0.9.			
1436				
1437	Fig. 8. $\delta^7$ Li of fine-grained suspended material as a function of weathering intensity for			
1438	CrunchFlow simulations (black dots; no chlorite simulations) and observational data from the			
1439	Amazon basin (Dellinger and others, 2017). Model output includes the full range of lithium			
1440	partition coefficients shown in Figure 3, and CrunchFlow output range is denoted by gray			
1441	shading.			
1442				
1443	APPENDIX FIGURE CAPTIONS			
1444				
1445	Table A1. CrunchFlow reaction network for plagioclase dissolution – beidellite precipitation			
1446	batch reactor sensitivity experiments.			
1447	_			
1448	Fig. A1. Relationships between $\delta'$ Lidiss and Weathering Intensity (left) and dissolved Li			
1449	concentrations (right) for erosion/uplift simulations with chlorite across a range of $P_{Li}$ values.			
1450				

- 1451 Fig. A2. Batch reactor output for plagioclase dissolution and beidellite precipitation showing
- 1452 (left) mineral precipitation/dissolution rates; (center) weathering congruence; and (right)  $\delta^7 Li_{diss}$ 1453 as a function of time for a range of  $P_{Li}$  values.
- 1454
- 1455 Fig. B1. Changes in Li concentrations and  $\delta^7$ Li<sub>diss</sub> with mean fluid transit time assuming a
- 1456 gamma-function transit time distribution with alpha value of 1. The discharge arrow shows
- 1457 qualitative relationships between discharge and mean transit times and demonstrating negative
- 1458 C-Q relationships for Li concentrations and  $\delta^7 Li_{diss}$  in simulations.
- 1459
- 1460

#### Table 1. CrunchFlow reaction network.

Mineral	Reaction	log(K <sub>eq</sub> ) @ 298 K	log(k)
Plagioclase (An <sub>20</sub> )	$Ca_{0.2}Na_{0.79739}{}^{7}Li_{0.002409}{}^{6}Li_{0.0001948}Al_{1.2}Si_{2.8}O_{8} \leftrightarrow 0.2Ca^{2+} + 0.79739Na^{+} + 0.002409{}^{7}Li^{+} + 0.0001948{}^{6}Li^{+} + 1.2AlO_{2}{}^{-} + 2.8SiO_{2}(aq)$	-20.076	-14.84
Quartz	$SiO_2(qz) \leftrightarrow SiO_2(aq)$	-3.9993	-55
Halloysite_7Li	$Al_{2-0.33x}^{7}Li_{x}Si_{2}O_{5}(OH)_{4} + 6H^{+} \leftrightarrow (2-0.33x)AlO_{2}^{-} + x^{7}Li^{+} + 2SiO_{2}(aq) + H_{2}O^{*}$	-36.4537	-17.14
Halloysite_ <sup>6</sup> Li	$Al_{2-0.33x}{}^{6}Li_{x}Si_{2}O_{5}(OH)_{4} + 6H^{+} \leftrightarrow (2-0.33x)AlO_{2}^{-} + x^{6}Li^{+} + 2SiO_{2}(aq) + H_{2}O^{*}$	-36.4537	Precip: -17.13225 Diss: -17.14
Chlorite <sup>a</sup>	$\begin{split} Mg_{1.03969}{}^{7}Li_{0.0005769}{}^{6}Li_{0.00004683}Fe_{0.2}K_{1.28}Al_{2.24}Si_{3}H_{13.52}O_{18} + 10.48H^{+} \\ & \qquad \qquad$	14.996	-13
Vermiculite <sup>a</sup>	$\begin{split} Mg_{0.1997}{}^{7}Li_{0.000576923}{}^{6}Li_{0.0004683}Fe_{0.441}K_{1.3}Al_{4.4}Si_{1.6}H_{8.52}O_{15.2} + 15.5H^{+} &\longleftrightarrow 0.2Mg^{2+} + 0.010962^{7}Li^{+} + \\ &0.0008897^{6}Li^{+} + 0.3941Fe^{2+} + 1.3K^{+} + 4.4Al^{3+} + 1.6SiO_{2}(aq) + 12H_{2}O \end{split}$	-5.39	-13
* x varies from 0.00035 - 0.0043		0035 - 0.0043	
		a	

<sup>a</sup> Based on Heidari et al. (2017)



#### Table 3. Variables calculated from model output

Model Output/Metric	Formula	Description
Erosion Rate (Mass) (t/km <sup>2</sup> /yr)	E = RP*BD	Prescribed volumentic regolith production (mm/yr) multiplied by bulk density (BD; kg/m3) in the top model cell
Chemical Denudation Rate (t/km <sup>2</sup> /yr)	$W = Q^{*}([Ca^{2+}] + [Mg^{2+}] + [Na^{+}] + [K^{+}] + [SiO2_{(aq)}])^{*}1000$	Prescribed infiltration rate (Q; m/yr) multiplied by dissolved solute load (g/L), evaluated at model domain base and weathering zone base
Weathering Intensity	WI = W/(W + E)	Ratio of chemical denudation rate to total denudation rate
Suspended Load δ <sup>7</sup> Li (‰)	$\begin{split} \delta^{7}Li_{sus} &= (M_{plag} * C_{plag} * \delta^{7}Li_{plag} + M_{chlor} * C_{chlor} * \delta^{7}Li_{chlor} + \\ M_{halloy} * C_{halloy} * \delta^{7}Li_{halloy})/(M_{plag} * C_{plag} + M_{chlor} * C_{chlor} + M_{halloy} * C_{halloy}) \end{split}$	Lithium mass-weighted $\delta^7$ Li composition of non-quartz minerals in the eroding top model cell, where <i>M</i> represents mineral mass and <i>C</i> represents mineral lithium concentration
Weathering Incongruence	$I = -R_{halloysite}/R_{plagioclase}$	Instaneous ratio of halloysite precipitation to plagioclase dissolution rates



**Regolith Production = Erosion + Weathering** 





# Figure 3



# Figure 5









# Figure 8



#### Table A1. CrunchFlow reaction network with 2:1 clay phase.

Mineral	Reaction	log(K <sub>eq</sub> ) @ 298 K	log(k)
Plagioclase (An <sub>20</sub> )	$\begin{split} & Ca_{0.2}Na_{0.79739}{}^{7}Li_{0.002409}{}^{6}Li_{0.0001948}Al_{1.2}Si_{2.8}O_{8} \\ \leftrightarrow 0.2Ca^{2+} + 0.79739Na^{+} + 0.002409{}^{7}Li^{+} + 0.0001948{}^{6}Li^{+} \\ & + 1.2AlO_{2}{}^{-} + 2.8SiO_{2}(aq) \end{split}$	-20.076	-14.84
Quartz	$SiO_2(qz) \leftrightarrow SiO_2(aq)$	-3.9993	-55
Ca-Beidellite_ <sup>7</sup> Li	$Ca_{0.165-0.5x}{}^{7}Li_{x}Al_{2.33}Si_{3.67}O_{10}(OH)_{2} + 7.32H^{+} \leftrightarrow (0.165-0.5x)Ca^{2+} + 2.33Al^{3+} + x^{7}Li^{+} + 3.67SiO_{2}(aq) + 4.66H_{2}O^{*}$	5.59	-17.14
Ca-Beidellite_ <sup>6</sup> Li	$Ca_{0.165-0.5x}{}^{6}Li_{x}Al_{2.33}Si_{3.67}O_{10}(OH)_{2} + 7.32H^{+} \leftrightarrow (0.165-0.5x)Ca^{2+} + 2.33Al^{3+} + x^{6}Li^{+} + 3.67SiO_{2}(aq) + 4.66H_{2}O^{*}$	5.59	Precip: -17.13225 Diss: -17.14

\* x varies from 0.000402 - 0.004945



Figure A2



