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1 **The lithium isotope response to the variable weathering of soils in Iceland**

2

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16

17

18 Abstract

19 This study has analysed Li isotopes ratios from well-studied soil and pore water
20 profiles from Iceland that have the same parent material but have experienced different
21 degrees of chemical weathering. Thus, from least to most weathered, we have analysed
22 vitrosols (V), gleyic andosols (GA), brown andosols (BA), Histosols (H) and Histic Andosols
23 (HA). Although the most weathered H and HA soils have the highest content in clay-sized
24 material, they have the least fractionated $\delta^7\text{Li}_{\text{pore water}}$ values. In contrast, the least weathered
25 GA and BA pore waters are most fractionated for Li isotopes. Given that Li isotope ratios are

26 fractionated by clay mineral formation, this appears counter-intuitive. A single trend for all
27 samples of $\delta^7\text{Li}$ as a function of Li/Na ratios suggests that they are all controlled by a process
28 with a single fractionation factor, in this case likely the formation of poorly-crystalline
29 allophane, which dominates in the “least weathered” soils. This rapidly forming secondary
30 mineral dominates Li isotope fractionation over more slowly-forming crystalline clays. The
31 fractionation along a single path shows that the key process here in controlling the Li isotope
32 ratio of surface waters is the degree of Li uptake by secondary minerals. This does not
33 necessarily correspond to the amount of clay minerals present in the soil, but to the amount of
34 clay minerals that are being newly formed in a single passage of the pore water through the
35 soil, or are in equilibrium with soil solutions at the time of sampling.

36

37 1.0 Introduction

38 The chemical weathering of rocks at the Earth’s surface is the primary control on
39 atmospheric CO₂ levels on varying timescales. Weathering provides both alkalinity
40 (stemming from water-rock-atmospheric-carbon interactions) and cations for the formation of
41 marine carbonates, sequestering carbon (Berner et al., 1983; Walker et al., 1981). If the rocks
42 being weathered are silicates, containing virtually no carbon, then this carbon sequestration is
43 a net sink and a primary long-term climate controlling process. At the same time, continental
44 weathering provides a number of critical nutrients (e.g. iron, silicon, phosphorus) to the
45 coastal oceans, which stimulates primary productivity (Hawley et al., 2017; Lalonde et al.,
46 2012). This productivity temporarily also takes up atmospheric carbon, but most marine
47 organic matter is reoxidised before it is buried in marine sediments. However, continental
48 chemical weathering and physical erosion also provide clay particles into the coastal oceans.
49 These both increase nutrient fluxes to primary productivity (because microbes leach nutrients
50 from the rock particles (Grimm et al., 2019)), but, critically, also greatly increase organic

51 carbon burial, thus also sequestering carbon on the long term (Kennedy et al., 2014; Kennedy
52 and Wagner, 2011). Overall, this makes silicate weathering the primary atmospheric CO₂
53 controlling process on both long (via carbonate precipitation) and short, seasonal timescales
54 (via organic carbon burial).

55 Significant effort has focused on understanding and quantifying weathering processes
56 and fluxes, as well as on the processes that control weathering (Gislason et al., 2009; Maher,
57 2011; West et al., 2005). In particular, a feedback control, via a temperature-driven
58 “thermostat” is likely to be the overarching process that has mitigated climate over geological
59 time, although time periods also exist where an increase in supply may have overridden this
60 climate stabilisation process, such as in the Cenozoic (Raymo and Ruddiman, 1992).

61 The debates on these various processes have been at the forefront of biogeochemistry
62 for almost four decades, but finding reliable and unambiguous tracers of weathering
63 processes both in the present and geological past has proven tricky. One of the most
64 established of such tracers is lithium isotopes. Lithium is a moderately incompatible trace
65 element that is present in all silicate rocks, in orders of magnitude more than in carbonate
66 rocks (Penniston-Dorland et al., 2017; Pogge von Strandmann et al., 2020). Hence, in
67 weathering environments, Li arises from the dissolution of silicates, even in carbonate-
68 dominated catchments (Kisakürek et al., 2005; Millot et al., 2010). Lithium isotopes (⁶Li and
69 ⁷Li) are not fractionated by primary productivity (Pogge von Strandmann et al., 2016), and
70 while it remains poorly understood whether plants fractionate Li isotopes, the Li
71 concentration of plants is sufficiently low for surface waters to remain largely unaffected
72 (Clergue et al., 2015; Lemarchand et al., 2010; Li et al., 2020; Steinhoefel et al., 2021).
73 Instead, Li isotopes are fractionated by the silicate weathering process, in that when primary
74 silicate rocks (with a relatively low isotope ratio) are dissolved, they impart a low (rock-like)
75 $\delta^7\text{Li}$ to waters. Isotopically light lithium is preferentially taken up by formation of secondary

76 minerals (clays, oxides, zeolites), as well as by adsorption onto secondary minerals, imparting
77 a heavy isotopic signature to residual waters (Chan et al., 1992; Hindshaw et al., 2019; Huh et
78 al., 2001; Millot et al., 2010; Pistiner and Henderson, 2003; Pogge von Strandmann et al.,
79 2019a; Pogge von Strandmann et al., 2017b; Pogge von Strandmann and Henderson, 2015;
80 Teng et al., 2010; Vigier et al., 2008). Consequently, dissolved Li isotopes are controlled by
81 the weathering congruency: congruent dissolution of silicates imparts a low $\delta^7\text{Li}$, while
82 incongruent weathering (where secondary minerals form) yields a high $\delta^7\text{Li}$ signature relative
83 to the parent material (Misra and Froelich, 2012; Pogge von Strandmann and Henderson,
84 2015).

85 Li isotopes are now well established as a tracer of both present and past weathering
86 processes (Dellinger et al., 2015; Hathorne and James, 2006; Lechler et al., 2015; Misra and
87 Froelich, 2012; Pogge von Strandmann et al., 2021a; Pogge von Strandmann et al., 2017a;
88 Pogge von Strandmann et al., 2013), but important questions remain, especially concerning
89 precisely how Li isotopes respond to changes in weathering processes. For example, the
90 effect of water-rock interaction time (e.g. secondary mineral formation rates) on Li isotopes
91 remains poorly quantified, partly because the timing of adsorption vs. incorporation during
92 neoformation is poorly known, and because the effect of Li uptake by oxides/oxyhydroxides
93 vs. clay or zeolites is also poorly known (Hindshaw et al., 2019; Li and Liu, 2020). This
94 study aims to investigate the, as yet, little quantified effects arising from variable degrees of
95 weathering in soils – for example, high solution $\delta^7\text{Li}$ could be due to either a high fraction of
96 removal with a low fractionation factor, or lower removal with a high fractionation factor.
97 Here we examine soil and pore water profiles from different soil types from Iceland. These
98 different profiles represent weathering of bedrock with a similar composition, but reflect
99 differing degrees of weathering.

100

101 2.0 Samples

102 Five different Icelandic soil types were analysed that represent increasing degrees of
103 chemical weathering, collected in 2009 and 2010 (Opfergelt et al., 2014), and are shown on a
104 soil map of Iceland (Fig. 1). These particular samples have previously been analysed for the
105 isotopes of Mg, Si, Fe and Zn (Opfergelt et al., 2014; Opfergelt et al., 2017a; Opfergelt et al.,
106 2017b). These Icelandic soils have specific definitions (Arnalds, 2004): Vitrisols (V)
107 represent the least weathered soils from Iceland, and have <1% organic carbon or <1% Si_{ox}.
108 Their mineralogy is dominated by volcanic glass, and they have little (5–10%) plant cover.
109 Broadly, they represent a desert-like environment with little vegetation or soil formation, but
110 in a cold and wet environment.

111 Gleyic andosols (GA – where US spelling is Andisols) are slightly more weathered,
112 and tend to have organic carbon contents (<12%) in surface horizons. They contain allophane
113 contents of up to 20%.

114 Brown andosols (BA) are increasingly weathered, and are the classical freely drained
115 andosols found in Iceland. They tend to contain significant allophane (15–30%), as well as
116 some ferrihydrite (1–8%).

117 Histosols (H) and Histic Andosols (HA) are the most weathered soils in this study.
118 Histosols are dominated by organic materials (>20% organic C). Histic Andosols are found in
119 poorly drained areas, and have sufficient aeolian inputs to reduce their organic contents
120 below the 20% limit for Histosols. Allophane content is 2–10%, and they meet the (Al +
121 0.5Fe)_{ox} criteria for Andosols.

122 Overall, Iceland is covered by ~48% Andosols (of all types), ~40% Vitrisols and
123 Leptosols and ~1% Histosols (and 11% by glaciers and ice caps) (Arnalds, 2004). In general,
124 vegetated areas are dominated by Andosols, desert areas by Vitrisols and wetlands by
125 Histosols.

126 In this study, the vitrisol (V) is from SW Iceland, located south of the Langjökull and
127 west of the Myrdalsjökull ice caps. The gleyic andosol (GA) is from just north of Reykjavik.
128 The brown andosol (BA), the Histosol (H), and the Histic Andosol (HA) are from the
129 Borgarfjörður catchment in western Iceland. Peats were developed at the HA and H sites
130 (Opfergelt et al., 2014). Overall, GA and BA soils are well-drained, while the H and HA soils
131 are poorly drained, with inevitable consequences for the water-rock interaction time. All soils
132 developed from similar tholeiitic basalt, and their difference from each other is largely due to
133 the soil drainage conditions (Opfergelt et al., 2014).

134 Further to the soils, a series of local secondary minerals was acquired from the
135 Natural History Museum in Reykjavik. These minerals were not always formed in the same
136 areas as the soils (although many are from the same catchment), but are from similar
137 Icelandic weathering environments, and are examined here to provide some context of
138 secondary mineralogy. These include samples of different calcites, clays (unidentified) and
139 six different types of zeolite (heulandite, chabazite, mesolite, stilbite, epistilbite and
140 scolecite). Further, an Fe-oxyhydroxide was taken from the coating on a pebble from a side
141 stream on the nearby Norðura River (locality A6; (Pogge von Strandmann et al., 2006)). The
142 major element composition suggests that this coating is largely impure ferrihydrite,
143 containing Al, P and possibly Si admixtures (Si not measured), typical of soil and stream
144 environments, but the presence of other phases, such as Al-hydroxide, cannot be ruled out.

145

146 3.0 Methods

147 *3.1 Sampling*

148 Sampling is described in Opfergelt et al. (2014). Briefly, typical parent tholeiitic
149 basalt was sampled at the BA site. This represents the type of parent material for the HA, H,
150 BA and GA sites. For the Vitrisol, the C horizon was considered as unweathered parent

151 volcanic ash. Soil profiles were described following FAO guidelines and sampled by horizon.
152 Soil samples were air-dried and sieved at 2mm. The measurement of chemical and physical
153 properties of the soils (pH, cation exchange capacity (CEC), chemical composition) is
154 described by Opfergelt et al. (2014).

155 Three representative soils were sequentially leached to separate out different phases,
156 and analysed for Li concentrations. The leaching technique followed a modified version of
157 Tessier et al. (1979) detailed in Revels et al. (2021). Briefly, the exchangeable fraction was
158 leached using $Mg(NO_3)_2$ at pH5 for 1 hour; carbonates were leached using acetic acid
159 buffered with Na acetate to pH5 for 5 hours; oxides were leached using $NH_2OH-HCl$ in
160 acetic acid at $96^\circ C$; the silicate residue was dissolved using the method described for bulk
161 soils below.

162 Soil solutions were sampled over two different seasons (September 2009 and June
163 2010), using macro rhizon soil water samplers, which filter pore waters at $0.2 \mu m$. Elemental
164 concentrations were determined by ICP-MS and ion chromatography, as described by
165 Opfergelt et al. (2014). In their initial measurements, Li concentrations were not always
166 determined, because $[Li]$ was so low, and was hence below detection limits. For these
167 samples, Li concentration was re-determined by this study during isotope analysis, by drying
168 down at least 10ml of sample, and comparing the MC-ICP-MS beam intensity to that of a
169 known concentration standard.

170

171 *3.2 Li isotope analyses*

172 Lithium isotope purification and analyses by our methods have been described in
173 elsewhere (Pogge von Strandmann et al., 2011; Pogge von Strandmann and Henderson,
174 2015). Briefly, soils were dissolved in $HF-HNO_3-HClO_4$, HNO_3 and HCl , and sufficient pore
175 water was dried down to attain 20ng Li. This was purified through a two-step cation

176 exchange method, and analysed relative to the LSVEC standard on a Nu Instruments HR
177 MC-ICP-MS at Oxford University. Splits taken before and after the Li collection bracket on
178 the columns show that >99.9% of Li analysed, which is essential for both precision and
179 accuracy. The total procedural blank for Li isotopes is effectively undetectable (<0.005 ng
180 Li). Long-term precision of this method and instrument is $\pm 0.6\%$ (2sd), based on repeated
181 seawater analyses of $31.2 \pm 0.6\%$ (2sd, n = 62), and BCR-2 basalt of $2.6 \pm 0.4\%$ (n = 6).

182

183 4.0 Results

184 Opfergelt et al. (2014) extensively characterised these soils, and results are given in
185 Tables 1 and 2. Briefly, the primary mineralogy of these soils (augite, Ca-plagioclase,
186 magnetite) reflects that of the primary tholeiitic basalt. Based on selective extraction,
187 secondary minerals are dominated by short-range ordered minerals, mainly allophane and
188 ferrihydrite, as also reported from other Icelandic soils (Arnalds, 2004; Sigfússon et al., 2006,
189 2008). No other secondary minerals were identified in the soils' clay-sized fraction (<2 μ m),
190 except in the HA soil, where XRD detected smectite and kaolinite (Opfergelt et al., 2014).
191 The allophane content is higher in the V-BA-GA soils than in the HA-H soils. This is because
192 allophane forms above pH 4.9 when Al is not complexing with humus and therefore the more
193 acidic HA-H soils have a lower allophane content (Mizota and van Reeuwijk, 1989). The
194 HA-H soils do contain some allophane, which is explained by aeolian deposition or by higher
195 pH conditions that may have existed earlier in the soil development cycle (Opfergelt et al.,
196 2014).

197 As described by Opfergelt et al. (2014), one of the primary chemical properties of the
198 bulk soils is the total reserve in bases (TRB = [Na] + [Mg] + [Ca] + [K]). The TRB is well-
199 defined as a relative index quantifying the leaching of these cations in the soils relative to the
200 parent material (e.g. Ameijeiras-Mariño et al., 2017; Delvaux et al., 1989; Henriot et al.,

201 2008), and hence provides an estimate of the content of the weatherable minerals in the soils.
202 Any fraction of cations that are included in present secondary, rather than primary minerals,
203 will still be included in the TRB. The TRB value in soils decreases with increased
204 weathering, and leads to the observation that the least weathered soil is the Vitrisol (average
205 TRB = 618 cmol_c/kg), compared to a value of 733 cmol_c/kg in the pristine basalt, followed by
206 the Gleyic Andosol (TRB = 401 cmol_c/kg), then the Brown Andosol (TRB = 358 cmol_c/kg),
207 then the Histosol (TRB = 194 cmol_c/kg), closely followed by the Histic Andosol (TRB = 158
208 cmol_c/kg), which is therefore the most weathered soil type. This relative degree of weathering
209 is supported by a direct correlation with the proportion of free Fe (Fe_d/Fe_t) (Opfergelt et al.,
210 2017b), which is also used as a weathering indicator. This is also further supported by the
211 total clay-sized content (<2μm), which is higher in the HA-H soils (53 and 43%,
212 respectively) compared to the BA-GA (both ~35%) and V (4.6%) soils (Table 1).

213

214 *4.1 Lithium in soils*

215 None of the soil profiles have a clear trend in Li concentrations with depth (Fig. 2a).
216 Overall, the GA soil has the highest Li concentrations (10.6 ± 2.8 (1sd) μg/g), followed by
217 the BA and V (8.3 ± 2.6 and 8.3 ± 3.0 μg/g, respectively), and then by the H and HA (5.2 ±
218 3.3 and 5.2 ± 2.8 μg/g, respectively). In comparison, the host basalt has a Li concentration of
219 5.89 μg/g, within the range for Icelandic basalts (Pogge von Strandmann et al., 2006; Pogge
220 von Strandmann et al., 2008; Pogge von Strandmann et al., 2012; Vigier et al., 2009).

221 The selective leaches of three soils (HA O1, H O6, BA BW2) show that between 0.2
222 and 3.4% of the total Li is in the exchangeable fraction, slightly less than observed during
223 experiments with Icelandic basalts (Pogge von Strandmann et al., 2019a), but similar to the
224 amount of Mg reported in this fraction (Opfergelt et al., 2014). Carbonates contain a similar
225 proportion of Li (1–4%). The oxide fractions (primarily Fe-oxides) contain significantly more

226 Li, ranging from 10% in the BA soil, to 25% in the HA soil. The silicate fraction contains
227 ~90% of the total Li in the BA soil, 70% in the HA soil, and 72% in the H soil (Table 1).
228 Although the recovered phases are slightly different compared to the BCR leaching of
229 Hawaiian basalt by Li et al. (2020) (e.g. that method combines exchangeable and carbonate
230 phases in a single leach), the results agree in that the silicate fraction dominates the Li budget,
231 followed by the oxide fraction. The least weathered leached soil (BA) has the least Li in any
232 of the secondary fractions.

233 Similarly, there are no trends with depth for soil $\delta^7\text{Li}$, except for the BA, where $\delta^7\text{Li}$
234 decreases linearly from 6.4 to 2.1‰ with increasing depth (Fig. 3). There is, however, a
235 difference in the average $\delta^7\text{Li}_{\text{soil}}$ between different soil types. From least to most weathered,
236 the V soil has an average of $1.9 \pm 1.9\text{‰}$, the GA of $5.2 \pm 1.3\text{‰}$, the BA of $2.7 \pm 3.0\text{‰}$, and H
237 of $-0.2 \pm 3.5\text{‰}$ and the HA of $0.5 \pm 1.4\text{‰}$. In other words, aside from the Vitrisol, the
238 average $\delta^7\text{Li}_{\text{soil}}$ decreases as the degree of weathering increases. The parent basalt has a $\delta^7\text{Li}$
239 of 3.6‰, within the range for Icelandic basalts and MORB. In particular, this parent value is
240 similar to that of 2010 Eyjafjallajökull ash (3.9‰; Pogge von Strandmann et al., 2012)), of
241 unaltered basalt at the Reykjanes Peninsular (3.7‰; Verney-Carron et al., 2015)), and
242 elsewhere in Iceland (3.1–4.0‰; Ryan and Kyle, 2004)), suggesting that there is little
243 variability across Iceland, and hence of the parent composition of our different soil profiles.

244

245 *4.2 Lithium in pore waters*

246 No pore waters exist from the very fast-draining Vitrisol. In the other soils, there is a
247 hint of increasing Li concentrations with depth in the HA and BA (Fig. 2b), but not in the
248 other soil profiles. On average, the Li concentrations increase with increasing degree of
249 weathering: the GA has an average pore water Li concentration of 0.09 ± 0.02 ng/ml,
250 compared to 0.15 ± 0.07 ng/ml in the BA, 1.78 ± 1.12 ng/ml in the HA and 3.27 ± 2.54 ng/ml

251 in the H. In comparison, Icelandic rivers range between 0.015 and 0.4 ng/ml, although up to
252 1.1 ng/ml in rivers with hydrothermal input (Pogge von Strandmann et al., 2006). A Histic
253 Andosol from elsewhere in Iceland has reported pore water Li concentrations of 0.67–1.94
254 ng/ml (Pogge von Strandmann et al., 2012).

255 There are sometimes significant differences between both the [Li] and the $\delta^7\text{Li}_{\text{PW}}$ (on
256 average $\sim 10\text{‰}$) from the same horizons, but from the two different years/seasons, collected
257 in September (late summer) 2009 and June (spring) 2010 (Fig. 3). Some variability between
258 shallow ($<70\text{cm}$) pore waters could be expected, given fluctuations in water-levels,
259 evaporation and redox variations, which could also explain the lack of consistent differences
260 between the different soils. Broad patterns appear to be maintained, however; for example, in
261 the HA, pore waters from both years trend towards lighter isotope ratios with depth (Fig. 3),
262 similarly to broadly flat profiles in the Gleyic Andosol.

263 Lithium isotope ratios show trends to lower values with depth in the HA and H
264 profiles, but not in the others. Overall, the pore water $\delta^7\text{Li}_{\text{PW}}$ is always higher than that of the
265 soils, except at the base of the HA soil, where pore waters become similar or even slightly
266 isotopically lighter than their corresponding soils. A precipitation correction (denoted by *) is
267 applied to the major element pore water data, based on the assumption that rainwater in
268 Iceland stems from the oceans (Pogge von Strandmann et al., 2006), and that all Cl^- in the
269 pore waters stems from marine aerosols, which is validated by the low Cl^- concentrations of
270 Icelandic basalt (Gíslason et al., 1996). We do not apply a precipitation correction to pore
271 water Li. This is because an attempted correction using seawater elemental ratios and isotope
272 compositions, and the measured [Cl], drives around a third of [Li] to negative values. We also
273 note that the pore water data do not lie on a mixing trend with seawater. This is a similar
274 approach to other basaltic Li studies (Liu et al., 2015; Pogge von Strandmann et al., 2006),
275 and we note that this may be due to analytical uncertainty in both [Cl] and [Li] values, as also

276 suggested by other studies (Pogge von Strandmann et al., 2010). We note that for samples
277 where the correction can be made, the contribution from sea spray is generally <10%, and
278 averages <5%.

279 Between different soil profiles there are large average $\delta^7\text{Li}_{\text{PW}}$ differences that
280 correlate with the degree of weathering. The least weathered GA has the highest $\delta^7\text{Li}_{\text{PW}}$ of
281 $26.7 \pm 3.4\text{‰}$ (1sd). The BA has an average $\delta^7\text{Li}_{\text{PW}}$ of $23.2 \pm 7.3\text{‰}$, the H of $14.3 \pm 3.3\text{‰}$ and
282 the HA of $6.6 \pm 6.9\text{‰}$. In comparison, Icelandic (non-hydrothermal) rivers have a $\delta^7\text{Li}$ of
283 $15.4\text{--}44\text{‰}$ (Pogge von Strandmann et al., 2006; Pogge von Strandmann et al., 2016; Vigier et
284 al., 2009), and another reported Icelandic Histic Andosol has pore waters that have a $\delta^7\text{Li}$
285 ranging between $7.4\text{--}21.4\text{‰}$ (Pogge von Strandmann et al., 2012).

286

287 *4.3 Lithium in secondary minerals*

288 While the fluids that correspond to the secondary minerals are not available (although
289 most were formed in the same catchment as most of the soils analysed here), analysis of the
290 solid secondary minerals, nevertheless, provides some insight into their effect on lithium
291 behaviour (Table 1). Li concentrations in the calcites are similar at $32\text{--}39\text{ ng/g}$, over two
292 orders of magnitude below Icelandic basalt. Of the silicate secondary minerals, the clays have
293 Li concentrations of $\sim 5\text{--}14\text{ }\mu\text{g/g}$, similar to that of the Fe oxyhydroxide ($14\text{ }\mu\text{g/g}$), but
294 significantly higher than the zeolites ($0.4\text{--}2.7\text{ }\mu\text{g/g}$).

295 Li isotope ratios in the calcites range between -1.0 and 5.4‰ . Inorganic calcites are
296 reported to have fractionation factors of $3\text{--}6\text{‰}$ from fluid (Marriott et al., 2004). The clays
297 exhibit a similarly wide variation in $\delta^7\text{Li}$, with values of -8.4 and 2.1‰ . The zeolites tend to
298 have fairly low $\delta^7\text{Li}$ values (-4.9 to -0.2‰), except for heulandite (7.9‰) and scolecite
299 (5.3‰). Given the differing locations and the variability in the pore waters detailed above, it
300 is not possible to determine fractionation factors from these samples. As such, the Li

301 concentrations potentially provide more information than the isotope ratios, supporting the
302 hypothesis that carbonate weathering does not significantly affect dissolved Li compared to
303 silicates (Kisakürek et al., 2005). In addition, the concentration data suggest that zeolites play
304 a more minor role in affecting Li behaviour than clays and Fe oxyhydroxides.

305

306 5.0 Discussion

307 *5.1 Elemental depletion in soils*

308 In Icelandic basalt weathering, Na is the most mobile major cation (Gíslason et al.,
309 1996), consequently it goes into solution preferentially over Mg and Ca, which are
310 approximately 10 times less mobile (Gíslason et al., 1996; Pogge von Strandmann et al.,
311 2006). Elemental depletion in soils can be assessed from τ calculations (Brimhall and
312 Dietrich, 1987; Chadwick et al., 1990), where the loss or gain of Li compared to the parent
313 material is compared to that of an immobile element. Positive τ values indicate a net
314 elemental gain, while negative ones indicate a loss. Here we use Zr as the immobile element,
315 although we note that τ values normalised to Nb yield values within $\pm 10\%$. Mobile major
316 cations such as Mg and Ca have consistently negative τ values (aside from one sample for
317 Mg), showing that they have been stripped from the soils. The average τ values for Mg and
318 Ca from each soil sequence decrease with increasing weathering grade (Fig. 4a,b), agreeing
319 with the other tracers of the degree of weathering, such as TRB. The overall τ values for
320 these elements are higher than, for example, in the highly weathered bauxite profiles
321 developed in the Columbia River Basalts (Liu et al., 2013), which is to be expected in the
322 more kinetically-limited weathering regimes of Iceland (Dellinger et al., 2015).

323 More immobile elements, such as Al, also tend to have negative τ values in most of
324 the samples (apart from at the base of the GA soil), albeit significantly higher numbers than
325 the mobile elements. The Gleyic Andosol has values close to zero, and these values then

326 decrease with increasing grade of weathering in the other profiles (Fig. 4c). In other words,
327 some Al has been stripped from the more weathered of these soils, but considerably less than
328 the mobile elements, and overall elemental loss is less than in more weathered Columbia
329 River Basalt soil profiles (where τ_{Mg} values are uniformly close to -1, and τ_{Al} values average
330 at ~ -0.55 (Liu et al., 2013)).

331

332 *5.2 Soil lithium isotopes*

333 Tau values were also calculated for Li, relative to Zr (Fig. 4d). Only the H and HA
334 soils show any Li loss, and it is the HA that shows the greatest Li loss (average $\tau_{Li,Zr} = -0.07 \pm$
335 0.6), while on average the Histosol has gained Li ($\tau_{Li,Zr} = 0.19 \pm 0.7$). The BA and GA soils
336 have gained even more Li ($\tau_{Li,Zr} = 0.27 \pm 0.15$ and 0.35 ± 0.28 , respectively). The vitrisol
337 has a tau value close to 0 (0.06), reflecting its relatively unaltered status. Positive τ values are
338 in contrast to soil profiles from Hawaii and the Columbia River Basalts, where soils are
339 almost always depleted in Li, and tend to exhibit much lower absolute τ_{Li} values in both
340 settings (Liu et al., 2013; Ryu et al., 2014). Relative Li enrichment at depth in Hawaii,
341 especially in older soils, is reported as due to Li retained in secondary minerals (Ryu et al.,
342 2014). In the Icelandic soils examined here, the average τ_{Li} for each profile decreases with
343 increasing weathering grade (based on TRB, and other tracers mentioned above), meaning
344 that the nominally least weathered profiles are enriched in Li relative to the parental rock,
345 while more weathered profiles are comparatively depleted in Li. The Gleyic Andosol in
346 particular is increasingly enriched towards the base of the soil, and this mimics enrichment in
347 Al (Fig. 4c), confirming that Li is being retained by secondary minerals – in this case largely
348 allophane.

349 There is no direct relationship between the δ^7Li_{soil} values of individual samples and
350 their τ_{Li} values. For example, both the Histic Andosol and especially the Histosol, show

351 layers that are relatively isotopically heavy compared to the rest of the soils (Fig. 3). In both
352 cases these correspond to ash layers. In the HA profile, the ash layer is mixed with a redox
353 layer of organic-rich material, while in the H profile it is pure ash.

354 Discounting those layers, however, there is a clear decrease in average $\delta^7\text{Li}_{\text{soil}}$ with
355 increasing level of weathering, except for the Vitrisol ($\delta^7\text{Li} = 1.9 \pm 1.9\text{‰}$), which overlaps
356 the composition of the parent basalt (3.6‰). Thus, $\delta^7\text{Li}$ values decrease from GA > BA > H ~
357 HA, which also broadly co-varies with the TRB values, although there is scatter within each
358 soil type. This also yields a positive co-variation between the average τ_{Li} and the $\Delta^7\text{Li}_{\text{soil-basalt}}$
359 (i.e. the difference in $\delta^7\text{Li}$ between the soil and the parent basalt) for each soil type (Fig. 5).
360 This trend is enhanced by highly depleted and isotopically light bauxites from the Columbia
361 River Basalts (Liu et al., 2013). Various Hawaiian soils also lie on this trend (Li et al., 2020;
362 Ryu et al., 2014), albeit with the humid profile from Li et al. (2020) having a large variability
363 (Fig. 5).

364 Isotopically lighter Li isotope ratios in soils could be taken to suggest a greater Li
365 accumulation in secondary minerals, relative to soils with heavier values (Clergue et al.,
366 2015; Pogge von Strandmann et al., 2012). However, in the case of these soils, those that are
367 enriched in Li relative to the parent basalt (positive τ_{Li}) are isotopically heavier compared to
368 those that are depleted in Li. There is also an overall positive co-variation between soil $\delta^7\text{Li}$
369 and the abundance of clays such as allophane (Fig. 6a), which is the dominant secondary Al-
370 silicate in these environments (Opfergelt et al., 2014; Stefansson and Gislason, 2001; Wada et
371 al., 1992). There is no co-variation between $\delta^7\text{Li}_{\text{soil}}$ and the abundance of ferrihydrite nor
372 clay-sized contents (Fig. 6b), showing that in these environments, Li sorption on iron
373 oxyhydroxides or incorporation into crystalline clays is not controlling isotope fractionation.
374 There is also is no correlation between soil $\delta^7\text{Li}$ and $\delta^{66}\text{Zn}$, where Zn is thought to be
375 controlled by the soil organic matter content (Opfergelt et al., 2017a), suggesting that Li

376 isotopes are not affected by organic matter, as also confirmed by other studies (Li et al.,
377 2020).

378 It therefore appears that the amount of total clay-sized material in the soils is not the
379 primary control on bulk soil $\delta^7\text{Li}$ values. The correlation between $\delta^7\text{Li}_{\text{soil}}$ and allophane
380 contents (and also $\delta^7\text{Li}_{\text{soil}}$ and τ_{Li}) is also hard to understand, because soil $\delta^7\text{Li}$ values extend
381 to values both higher and lower than those of unaltered basalt, or indeed Icelandic volcanic
382 ash (Pogge von Strandmann et al., 2012). Thus, it is unlikely that the isotopically light bulk
383 soils values of the H and HA are due to a low allophane content. Instead, as discussed below
384 (Section 5.3), it appears more likely that greater allophane content is associated with higher
385 $\delta^7\text{Li}$ values because the secondary minerals are precipitating from isotopically heavy soil
386 pore waters. In other words, greater clay formation is driving pore water $\delta^7\text{Li}$ higher, from
387 which secondary minerals are then precipitating. This can occur in a standard coupled
388 relationship of two reservoirs with a constant fractionation factor (Faure, 1986). It would,
389 however, also mean that the initial dissolution of basalt would at first yield secondary
390 minerals that are isotopically lighter than basalt. This initial isotopically lightest phase is not
391 being resolved here, however. Complete loss of this phase via erosion is relatively unlikely, if
392 this phase is an aluminosilicate, because allophane is the first of these phases to form
393 (Arnalds, 2004). Therefore, such a light phase would have to be of another composition, or,
394 more likely, is not resolvable in the analysis of bulk soils due to mass balance considerations.
395 This is discussed further below in the context of the pore waters.

396 An alternate hypothesis is that the least weathered soils (BA and GA) have a higher
397 $\delta^7\text{Li}$ because they have experienced a lower solid to solution partitioning of Li, i.e. a lower
398 weathering intensity. This can be tested by comparing the measured τ_{Li} value to those based
399 on a steady-state mass balance between the dissolved load, soil and bedrock. If modelled
400 (steady-state) τ_{Li} values are similar to measured values, it would imply that the system is in

401 steady-state, and weathering intensity (the Li in the water/solid ratio) is controlling the soil Li
402 gain or loss. In contrast, if modelled and measured values are not similar, then there is a
403 decoupling between the present-day solution and solids, and more rapidly forming secondary
404 minerals (i.e. allophane) are controlling solution compositions. We model τ_{Li} using equations
405 from Bouchez et al., 2013:

$$406 \quad w = \frac{\delta_{soil} - \delta_{rock}}{\delta_{soil} - \delta_{diss}} = -\tau^{steady-state}$$

407 Where δ is the δ^7Li values of the soil, rock and dissolved load. Overall, there is no
408 relationship between the measured and modelled steady-state τ_{Li} values ($r^2 = 0.15$). However,
409 there is a positive co-variation for the HA soils (Fig. 7), while there is none for the other soil
410 types. This strongly suggests that the isotopically heavy soils (BA and GA) are not at steady-
411 state between their solutions and soils, whereas the HA soil may be. In turn, this suggests that
412 our previous hypothesis is correct, and the BA and GA soils exhibit decoupling between their
413 solutions and soils, and therefore their high δ^7Li may be due to allophane taking up Li from
414 heavy pore waters.

415

416 *5.3 Allophane control on pore water profiles*

417 These soils also exhibit a positive co-variation between δ^7Li_{PW} and allophane contents
418 (significant $p < 0.01$ for Spearman Rank correlation; Fig. 8). Thus, while the total clay-sized
419 content co-varies negatively with δ^7Li_{PW} (greater clay-sized content in the poorly-drained
420 soils), the allophane content co-varies positively with δ^7Li_{PW} (greater allophane in the
421 rapidly-drained soils). This correlation is highly unlikely to be caused by redissolution of
422 isotopically light clay minerals, as observed in supply-limited regimes such as the tropics
423 (Dellinger et al., 2015), because weathering regimes in Iceland are dominantly kinetically-
424 limited (Dellinger et al., 2015; Gíslason et al., 1996). Instead, allophane is poorly crystalline

425 and can form rapidly in these soils, and together with ferrihydrite (which does not show a
426 useful correlation with $\delta^7\text{Li}$), tends to dominate both the secondary weathering products and
427 the exchangeable components in these volcanic soils (i.e., in many Icelandic soils it is the
428 dominant silicate secondary mineral), especially at $\text{pH} > 4.9$ (Opfergelt et al., 2014;
429 Sigfússon et al., 2008). As mentioned above, the similar trend in $\delta^7\text{Li}_{\text{soil}}$ with allophane
430 contents is then likely caused by allophane precipitating from isotopically heavy pore waters.
431 Formation from the congruent dissolution of volcanic ash is unlikely, because measured ash
432 has a $\delta^7\text{Li}$ lower than that of the heaviest soils (Pogge von Strandmann et al., 2012). This
433 requires that allophane precipitating from the very first step of dissolution (when solution
434 $\delta^7\text{Li}$ would presumably be close to primary rock $\delta^7\text{Li}$) is not resolved in this study – either
435 because it is not present (unlikely, given that allophane is the first silicate secondary mineral
436 to form in Iceland (Arnalds, 2004)), or more likely that it is not resolvable in bulk soils due to
437 mass balance. A mass balance, using the fractionation factor determined below and a basaltic
438 starting composition, suggests that the first 20% of Li transferred from solution to secondary
439 minerals would have a $\delta^7\text{Li} < 0\text{‰}$, and the total $\delta^7\text{Li}$ of the solid after taking up 60% of Li
440 from solution would be +1‰. The calculations below (Section 5.4) show that, on average, 80
441 $\pm 27\%$ of Li is removed from solution, which would yield a clay $\delta^7\text{Li}$ of 2.5‰, if starting
442 from congruent basaltic dissolution. In other words, the initial light precipitated solid is not
443 detectable in bulk measurements.

444 It therefore appears likely that the Li isotope composition of the pore waters is
445 largely being controlled by the formation of the more rapidly forming poorly crystalline
446 allophane, rather than the slower formation of crystalline clays such as smectite or kaolinite,
447 which are only detectable in the HA soil by XRD (Opfergelt et al., 2014). The clay fraction is
448 comprised of the $< 2\mu\text{m}$ fraction, so includes Fe-oxyhydroxides as well as clays. However,
449 there is no co-variation between pore water $\delta^7\text{Li}$ and Fe extracted by dithionite-citrate-

450 bicarbonate (DCB) (Opfergelt et al., 2017b), suggesting that neither crystalline clays nor Fe-
451 oxyhydroxides are controlling Li isotopes here.

452 The relatively low pH of the poorly-drained H and HA pore waters also results in a
453 co-variation between $\delta^7\text{Li}_{\text{PW}}$ and pH (Fig. 8d). Primary silicate dissolution will consume
454 protons, raising pH. In contrast, secondary mineral formation will tend to release protons,
455 thus lowering pH. Organic matter decomposition will also have the same effect (Gislason and
456 Eugster, 1987; Pogge von Strandmann et al., 2010), and in the peaty H and HA soils, it is
457 likely this latter process that is affecting pH. Thus, the soils with the highest clay-sized
458 contents have the lowest pore water pH, but this does not appear to be affecting the pore
459 water Li isotope ratio. Instead, the relationship is due to a positive co-variation between pH
460 and the saturation index of allophane. This agrees with the overall observation that allophane
461 only forms above pH 4.9 when Al stops forming a complex with humus (Mizota and van
462 Reeuwijk, 1989). This then provides further evidence that it is the formation of allophane
463 over other clays that is dominating Li isotope fractionation.

464

465 5.3.1 Using Li/Na ratios to constrain Li isotope fractionation factors

466 To further understand the processes controlling $\delta^7\text{Li}_{\text{PW}}$, the isotope ratios can be
467 compared to the Li/Na ratios (Liu et al., 2015). In these pore waters, there is a strong negative
468 correlation between $\delta^7\text{Li}_{\text{PW}}$ and Li/Na* (Fig. 9), as would be expected if secondary minerals
469 take up Li over Na, and the lighter Li isotope preferentially. Similar relationships have been
470 seen in basaltic rivers from Iceland and elsewhere (Liu et al., 2015; Pogge von Strandmann et
471 al., 2010; Pogge von Strandmann et al., 2006; Pogge von Strandmann et al., 2016; Pogge von
472 Strandmann et al., 2017b; Vigier et al., 2009). The interesting feature concerning the
473 relationship observed here, however, is that the pore waters from the soils that have
474 undergone the most weathering (HA and H, as determined from CRB depletion) are the least

475 fractionated from basalt or bulk soils for both Li/Na and $\delta^7\text{Li}_{\text{IPW}}$ (i.e. most congruent
476 dissolution), while those that have undergone the least weathering (BA and GA), and overall
477 have the lowest total clay-sized contents, are the most fractionated, implying that they also
478 have the most secondary mineral formation. In general, H and HA soils are more poorly
479 drained than BA or GA soils. In principle, that should lead to greater water-rock contact times
480 in the H and HA soils, which would promote secondary mineral formation and raise solution
481 $\delta^7\text{Li}$ (Dellinger et al., 2015; Pogge von Strandmann et al., 2017b; Pogge von Strandmann and
482 Henderson, 2015). Given that this is the opposite to what is observed, other processes must be
483 controlling the pore water $\delta^7\text{Li}$ composition.

484 Interestingly, both HA soil profiles (i.e., from this study and from Pogge von
485 Strandmann et al., 2012) plot along the same $\delta^7\text{Li}_{\text{IPW}}$ vs. Li/Na trend (Fig. 9), suggesting that
486 while their isotope ratios are different, the fractionation factors are the same in both soils. In
487 other words, a different amount of Li has been removed from the water by secondary mineral
488 formation, but with the same fractionation factor. Hence, the pore waters imply that the Histic
489 Andosol from this study is dissolving almost congruently at the base of its profile (Fig. 9).
490 These are the O layers of the soil, and it may be that organic acids are driving the pH low,
491 and causing more congruent dissolution of primary and/or secondary minerals.

492 If allophane is dominating Li removal from the remaining pore waters, then it is
493 possible to calculate an isotopic fractionation factor, based on the removal of Li relative to Na
494 from solution. The logarithmic relationship between Li/Na and $\delta^7\text{Li}$ for the trend shown by
495 the GA, BA and HA pore waters can be modelled using a Rayleigh fractionation relationship.
496 The starting composition can either be set to the host soils (as in Fig. 9b), or the pristine
497 basalt. The two starting compositions make little overall difference to the fractionation factor
498 α , which varies by 0.01 according to the initial material. The average Rayleigh α value for
499 these soils is 0.993 ± 0.03 (Fig. 9b), identical to the average α value of 0.992 for all basaltic

500 rivers (Pogge von Strandmann et al., 2017b), and to the α value of 0.991 determined in
501 inorganic basalt-water weathering experiments (Pogge von Strandmann et al., 2019a).
502 Equilibrium fractionation factors are also shown in Figure 9b, but the trends of the data do
503 not appear to follow linear trends, except potentially for the HA pore waters. In general,
504 therefore, the similarity in fractionation factors between soils, inorganic experiments and
505 rivers argues that the overall fractionation of Li isotopes during basalt weathering is similar
506 regardless of first order controls, such as runoff or climate. These factors may however be the
507 cause of the scatter observed in global basaltic rivers. In addition, the similarity of the soil and
508 river fractionation factors with inorganic experiments suggests that vegetation does not form
509 a significant control on the Li isotope composition of surface waters, or that vegetation
510 imposes an identical fractionation factor.

511 In general, if most of the pore waters can be modelled by a similar fractionation
512 factor, it would supply strong evidence that one secondary mineral is dominantly removing Li
513 from solution (Hindshaw et al., 2019). In this case, given the co-variation with $\delta^7\text{Li}_{\text{PW}}$, this
514 mineral would appear to be allophane, which also agrees with the similar values of the historic
515 andosol reported in Pogge von Strandmann et al. (2012), where allophane was also the
516 dominant silicate secondary mineral. The similar positive co-variation between bulk soil $\delta^7\text{Li}$
517 and allophane contents could therefore also imply that the allophane is forming through
518 uptake of Li from the pore waters, driving the bulk soil Li isotope ratio heavy as well. In
519 effect, this could explain why the ‘closed-system’ relationship of Rayleigh fractionation
520 appears to fit the fractionation behaviour better than equilibrium fractionation. This would be
521 because allophane formation (or at least its uptake of Li) is faster than the residence time of
522 the pore waters, meaning that they behave as a closed system. This applies even to the
523 rapidly-draining BA and GA soils, suggesting that Li uptake is very rapid, and indeed water-

524 basalt interaction experiments show that solution $\delta^7\text{Li}$ shows more than 12‰ fractionation
525 within 12 days, and more than 16‰ in a month (Pogge von Strandmann et al., 2019a).

526

527 *5.4 Element mobility and co-variation with Li isotopes*

528 The use of Li isotopes as a palaeo-weathering tracer (Hathorne and James, 2006;
529 Pogge von Strandmann et al., 2017a) opens the discussion as to whether Li isotopes only
530 inform on weathering regime changes (Dellinger et al., 2015), or whether they can inform on
531 CO_2 drawdown efficiency (Pogge von Strandmann and Henderson, 2015). The latter requires
532 that Li isotopes can be used to determine behaviour of the critical elements to the carbon
533 cycle, calcium and magnesium. This can be examined through the mobility of elements,
534 which is the tendency of elements to stay in solution (= mobile) compared to that of going
535 into secondary minerals (=immobile) (Gislason et al., 1996). This is determined by the ratio
536 of the elements in question (here, Li to Ca) of the dissolved load to that of a starting
537 composition (here, either parent basalt, or host soil). A relative mobility value of 1 implies
538 that the Li/Ca ratio of the water is the same as that of the starting material, while lower values
539 mean that Ca is more mobile than Li.

540 Icelandic rivers exhibit a very strong relationship between Li to Ca relative mobility
541 and their $\delta^7\text{Li}$ value ($r^2 = 0.7$; Fig. 10a), as well as a similar one for Li to Mg mobility ($r^2 =$
542 0.67 ; Fig. 10c) (Pogge von Strandmann et al., 2016; Pogge von Strandmann et al., 2020). In
543 turn, this means that Li isotopes ratios could inform on the sequestration efficiency of CO_2 .
544 The soil pore waters exhibit a similar, albeit more scattered, relationship. The GA waters
545 have the lowest Li mobility (compared to Ca or Mg), and, given that the decrease in Li
546 mobility is due to uptake by secondary minerals, the highest $\delta^7\text{Li}_{\text{PW}}$. The HA and H waters
547 show contrasting behaviour (Fig. 10a), with the HA waters having low $\delta^7\text{Li}_{\text{PW}}$ but still higher

548 Ca or Mg than Li mobility, while the H waters have intermediate $\delta^7\text{Li}_{\text{PW}}$, but a greater Li
549 mobility than Ca.

550 When comparing the mobility vs. $\delta^7\text{Li}_{\text{PW}}$ trend of these pore waters to basaltic rivers
551 from Iceland (Pogge von Strandmann et al., 2006) and the Azores (Pogge von Strandmann et
552 al., 2010), as well as the other Icelandic Histic Andosol (Pogge von Strandmann et al., 2012),
553 the HA soils from this study lie off the overall trend. If they are removed from the overall
554 pore water trend, then that trend and its r^2 becomes similar to that for the rivers ($r^2 = 0.7$ for
555 Ca, Fig. 10a; $r^2 = 0.55$ for Mg, Fig. 10c).

556 The behaviour of the pore waters can be taken further: a simple Rayleigh relationship,
557 using the best-fit alpha value from Section 5.3.1, can be used to determine the fraction of Li
558 taken into solution (Pogge von Strandmann et al., 2012; Vigier et al., 2009)(Fig. 10b). Thus,
559 on average the GA pore waters have the lowest fraction of Li in solution (average $7 \pm 4\%$,
560 where the uncertainty stems from the uncertainty in α), which increases to averages of $9 \pm$
561 6% in the BA pore waters, $14 \pm 7\%$ in the H and $50 \pm 6\%$ in the HA (up to 97% in some HA
562 waters). Again, as above, excepting the HA pore waters, there is a significant ($p < 0.01$,
563 Spearman Rank) relationship between the Li to Ca relative mobility (and Li to Mg relative
564 mobility) and $\delta^7\text{Li}_{\text{PW}}$, or the fraction of Li in solution that is derived from the $\delta^7\text{Li}_{\text{PW}}$ value
565 (Fig. 10b and d). In effect, when $\sim 10\%$ total Li is in solution, then Li and Ca or Mg are
566 similarly mobile (relative mobility $\sim 1_{-0.5}^{+1.5}$). In contrast, where $\sim 2\%$ of total Li is in solution,
567 Li is approximately a factor of 50 times less mobile than Ca (relative mobility $\sim 0.02_{-0.015}^{+0.06}$),
568 due to uptake into secondary minerals. This type of relationship has been reported in
569 experimental basalt-water interactions (Pogge von Strandmann et al., 2019a), as well as in
570 basaltic, andesitic and gneissic rivers (Clergue et al., 2015; Pogge von Strandmann et al.,
571 2010; Pogge von Strandmann et al., 2006; Pogge von Strandmann et al., 2020; Wimpenny et
572 al., 2010), and suggests that the $\delta^7\text{Li}$ value can be used to both determine the fraction of Li in

573 solution (i.e. the weathering regime) and from that the behaviour of Ca in solution. This
574 would be interesting especially from a palaeo-weathering perspective (Misra and Froelich,
575 2012; Pogge von Strandmann et al., 2017a; Pogge von Strandmann et al., 2013), because it
576 implies that the behaviour of an element directly involved in sequestering CO₂ could be
577 quantified. However, the HA pore waters here exhibit a trend with a significantly different
578 gradient from the other samples (Fig. 10b and d), and this highlights that more characterising
579 work is necessary. In this case, as mentioned above, the HA pore waters appear to follow a
580 linear rather than log trend in $\delta^7\text{Li}$ vs. Li/Na space (Fig. 9), potentially suggesting equilibrium
581 fractionation, and hence a subtly different fractionation factor. This may be because there are
582 hints that other clays are forming in these soils, compared to the other profiles (Opfergelt et
583 al., 2014; Opfergelt et al., 2017a; Opfergelt et al., 2017b).

584

585 *5.5 Li vs. Mg isotopes*

586 Magnesium is a critical element in the carbon cycle, because of the formation of Mg-
587 carbonates that sequester CO₂, and the exchange of Mg for Ca at mid-ocean ridges
588 (Humphris and Thompson, 1978; Walker et al., 1981). Mg isotopes have been suggested as a
589 tracer of Mg behaviour, but the processes that control Mg isotopes are complex, and include
590 lithology, secondary minerals and plants (Hindshaw et al., 2020; Tipper et al., 2006a; Tipper
591 et al., 2006b). It may be possible to constrain weathering processes using a combination of
592 isotopic systems, including those of Li and Mg. In particular, because Li substitutes for Mg in
593 both primary and secondary silicates (Hindshaw et al., 2019). Magnesium isotopes from these
594 particular soils and pore waters have been interpreted as representing a combination of Mg
595 adsorption by the soil exchangeable fraction (of heavy Mg isotopes preferentially) and of Mg
596 uptake by vegetation (which also takes up heavy Mg isotopes preferentially) (Opfergelt et al.,
597 2014). A strong control on Mg isotopes by adsorption onto allophane has also been reported

598 in other Icelandic soils (Pogge von Strandmann et al., 2012). When plotted against Li
599 isotopes (Fig. 11a), the pore waters are always isotopically lighter than the host basalt and
600 soils for Mg isotopes, but (as mentioned above), some pore waters have similar isotope ratios
601 to their host soils for Li isotopes.

602 Excluding the largely unfractionated HA from this study (bearing in mind that the
603 Histic Andosol studied by Pogge von Strandmann et al. (2012) behaves slightly differently –
604 Fig. 11a), there is a general negative relationship between Li and Mg isotopes, largely due to
605 the preferential uptake and sorption of light Li and heavy Mg by secondary minerals. Hence,
606 although there is scatter in any potential relationship, it is interesting that the pore waters
607 from soils that are nominally the least weathered are the most fractionated for both Li and Mg
608 isotopes. This supports the hypothesis that the secondary mineral that is affecting the sampled
609 pore waters (or at least their Li and Mg behaviour) is allophane, rather than other clays that
610 take longer to form, because it is the most fractionated (least weathered) soils that yield the
611 most favourable chemical conditions for allophane formation.

612 However, it is also clear that there is significant scatter in the relationship for these
613 pore waters between Li and Mg isotopes (Fig. 11a). In the absence of Mg-rich secondary
614 minerals in these soils (Opfergelt et al., 2014; Pogge von Strandmann et al., 2012; Sigfússon
615 et al., 2006, 2008), both Li and Mg are controlled by incorporation by substitution and
616 adsorption. However, the substitution and especially the sorption kinetics are different for the
617 two elements (experimentally by a factor of ~ 2.6 (Pogge von Strandmann et al., 2021b)),
618 meaning that similar behaviour will decouple the two elements and their isotopes. Scatter is
619 also caused by Mg uptake by plants, which causes isotope fractionation, and in these soils is
620 thought to be an important controlling process for Mg isotope ratios (Opfergelt et al., 2014).
621 Thus, even in the relatively simple weathering environment of Iceland, elements and isotopic

622 systems that are coupled by the same overall process (silicate weathering) and decoupled by
623 the specifics of the process for each element.

624

625 *5.6 Li vs. Si isotopes*

626 Silicon is another critical element in the carbon cycle, largely because it is the
627 weathering of silicates that ultimately removes atmospheric CO₂, and also because Si can be
628 a critical nutrient in certain ocean environments (Frings et al., 2016; Hatton et al., 2019). Si
629 isotopes are often interpreted similarly to Li isotopes, because secondary minerals
630 preferentially take up the light Si isotopes, driving waters isotopically heavy, and because
631 both trace silicate weathering processes (De la Rocha et al., 2000; Georg et al., 2007).
632 Because of this, both isotope systems exhibit a general positive covariation when comparing
633 all global river and pore water data (Pogge von Strandmann et al., 2017b). However, some
634 studies, including from Iceland, have shown a significant degree of scatter between Li and Si
635 isotopes, taken to be either influenced by plants (which preferentially take up light Si
636 isotopes, but do not fractionate Li isotopes), or due to kinetic differences if Si is dominated
637 by neoformation, compared to faster incorporation or adsorption for Li isotopes (Georg et al.,
638 2007; Georg et al., 2009; Opfergelt et al., 2013; Pogge von Strandmann et al., 2006; Pogge
639 von Strandmann et al., 2019b; Pogge von Strandmann et al., 2012; Vigier et al., 2009).

640 In our data, all the soils are isotopically lighter than the host basalt for Si isotopes
641 (Fig. 11b), due to the formation of secondary minerals (Opfergelt et al., 2017b). In contrast,
642 soil $\delta^7\text{Li}$ of the present study is only occasionally lower than the value for basalt, and there is
643 no correlation between bulk soil $\delta^7\text{Li}$ and $\delta^{30}\text{Si}$ for any of the soil profiles. Given that
644 secondary minerals clearly do take up the light isotopes of both elements (because pore
645 waters are isotopically heavy), this behaviour may be a function of the overall mass balance.
646 Because Si makes up a significant portion of clays (for example, Si makes up 20 molar% of

647 forsterite, 28% of Ca-montmorillonite and 14% of allophane), but Li does not (at most 0.01%
648 of clays), the bulk soil $\delta^{30}\text{Si}$ will be measurably perturbed by clay formation, while the bulk
649 soil $\delta^7\text{Li}$ will largely not be.

650 In contrast, the pore waters are clearly affected by secondary mineral formation in
651 both systems, driving an overall positive co-variation between $\delta^7\text{Li}$ and $\delta^{30}\text{Si}$ (Fig. 11b).
652 However, this co-variation does not exist for individual soil profiles. Further, the pore waters
653 that are most fractionated for Li isotopes (GA and BA) are not for Si, where the heaviest
654 values are from the Histosol (and also the HA of Pogge von Strandmann et al., 2012). From
655 the relationship between $\delta^7\text{Li}$ and Li/Na discussed above (Fig. 11a), it is clear that the Li
656 isotopes in the pore waters are only being fractionated by a single process (or processes with
657 a uniform fractionation factor), which then suggests that more variable processes are
658 affecting the Si isotope ratios of the pore waters. These likely include uptake by plants, and
659 the varying (and generally slower) kinetics that can affect the uptake of a major element into
660 secondary minerals (Opfergelt et al., 2017b; Pogge von Strandmann et al., 2012).

661 The relationships between $\delta^7\text{Li}$ and Mg and Si isotopes indicate the differing and
662 multiple processes that control Mg and Si over those that control Li (given the virtually linear
663 relationship between $\delta^7\text{Li}$ and Li/Na). This therefore demonstrates the potential
664 complications that can affect major elements over trace elements with relatively simple
665 behaviour during weathering such as Li (Pogge von Strandmann et al., 2019b), and also
666 confirms the potential of Li isotopes as a weathering tracer.

667

668 6.0 Conclusions

669 This study has analysed Li isotope ratios from well-studied soil and pore water
670 profiles from Iceland that have experienced different amounts of chemical weathering. Pore
671 waters from the soils that have experienced least weathering, based on the total reserve in

672 bases (Gleyic Andosol and Brown Andosol) have the highest $\delta^7\text{Li}_{\text{PW}}$ values (14–34‰), and
673 these soils are also enriched in Li compared to their parent basalts. In contrast, those that are
674 more weathered (HA and H) have the lowest $\delta^7\text{Li}_{\text{PW}}$ values (-1.4–20‰), and the bulk soils
675 are depleted in Li.

676 The pore waters form a well-defined negative trend for $\delta^7\text{Li}$ as a function of Li/Na
677 ratios, suggesting that they are controlled by a process with a single fractionation factor,
678 likely the formation of allophane. Hence, a large range of solution $\delta^7\text{Li}$ (~36‰) can be
679 generated by weathering with a single fractionation factor, due to variable proportion of Li
680 uptake. The correlation between $\delta^7\text{Li}$ and uptake presents the possibility of combining
681 isotope ratio and flux into a single solution when interpreting palaeo—records. The isotopic
682 fractionation does not correspond to the amount of clay-sized material present, but to the
683 amount of clay undergoing neoformation. In other words, it is secondary minerals that are
684 currently forming, rather those that have previously formed, which control the $\delta^7\text{Li}$ of surface
685 waters. Hence, profiles that have experienced only little overall chemical weathering, but
686 have formed a relatively large amount of amorphous, rapidly-formed, secondary minerals
687 (allophane), can exhibit substantial Li isotope fractionation.

688 Most of the pore waters also plot along the same trend between $\delta^7\text{Li}$ and the relative
689 mobility of Li to Ca or Li to Mg as other Icelandic surface waters. This suggests that
690 dissolved Li isotope ratios can be used not only to inform on Li behaviour and the silicate
691 weathering regime, but also Ca behaviour. Effectively, this can then be used to determine the
692 fraction of Ca in solution vs. secondary minerals in palaeo-records, and thus CO₂ drawdown
693 efficiency. In other words, with data from more weathering lithologies, Li isotopes may be
694 useful for quantitatively determining past CO₂ drawdown behaviour.

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696

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Bulk soils				C	CEC	TRB	Ferrihydrite	Allophane	Total clay	Na	Al	Fe	Ca	Mg	Zr	Li	$\delta^{7}\text{Li}$	2sd
Soil name	ID	Horizon	Depth (cm)	%	cmolc.kg-1	cmolc.kg-1	%	%	%	g/kg	g/kg	g/kg	g/kg	g/kg	µg/g	µg/g	‰	
Histic Andosol	HA	A1	0-15	18.5	40.1	148	7.1	4.9	56.1	4.3	50.5	165.7	14.6	6.4	167	4.70	0.9	0.3
		A2	15-26	16.9	55.3	177	5.4	9.5	54.3	5.5	72	98.9	17.7	7.3	237	5.80	-0.3	0.2
		Bw1	26-40	14.5	61.8	138	11.9	8.3	55.2	4.4	60.4	175.1	13.8	5.6	201	4.60	1.4	0.3
		Bw2	40-57	16.9	53.8	156	7.1	6.9	45.5	6.5	63	164.4	16.1	5.0	263	4.53	-0.7	0.4
		O1	57-67	28.7	59.4	103	4.4	6.6	53.8	3.3	44	55.7	11	3.9	275	2.36	-0.8	0.3
		redox	67-83	12.8	28.3	332	1.4	6.9		18.7	78.6	57.6	30.2	9.9	431	11.0	3.0	0.2
		O2	83+	42.7	71.3	52.4	3.2	1.4	55.3	2.1	22.3	42.9	5.4	1.7	280	3.21	0.3	0.5
Histosol	H	O1	0-13	21.0	49.5	266	6.8	2.5	52.3	6.4	39.5	120.3	28.5	11.1	112	7.86	-1.1	0.2
		O2	13-26	22.9	39.5	192	3.8	1.8	8.4	5.6	37	83.8	19.2	8.1	131	6.30	-4.3	0.3
		O3	26-40	18.6	52.6	188	5.9	6.7	38.2	7.5	56.4	141.7	18.6	6.8	252	4.32	0.3	0.3
		T	40-52	12.0	43.2	60.9	43.5	9.8		1.2	20.9	341.5	7.6	2.1	74	0.83	6.9	0.5
		O4	52-63	10.3	30.0	283	3.3	4.6	44.4	15.1	63.4	101.6	26.4	8.6	339	10.3	-0.7	0.4
		O5	63-72	29.3	65.6	115	4.7	2.3	70.9	2.8	34.3	140.7	12.6	4.6	132	2.04	0.0	0.3
		O6	72+	30.4	85.3	118	3.9	3.5	42.7	3.6	35.5	86.6	12.7	4.4	212	4.54	-2.2	0.2
Brown Andosol	BA	A1	0-21	7.29	36.8	403	7.8	10.7	33.2	10	83.2	111.9	39.1	19.2	223	9.05	6.4	0.5
		A2	21-40	7.92	35.2	370	8.6	11.0	29.7	9.6	86.8	117.2	36.1	17.2	241	8.97	5.0	0.4
		Bw1	40-52	6.47	36.8	377	7.3	9.2	37.2	10.1	91.3	122.7	35.7	17.8	217	9.66	3.9	0.2
		Bw2	52-96	7.25	50.9	317	9.7	13.0	41.0	8.5	97	126.1	31.1	14.5	262	9.08	3.7	0.2
		Bw3/C	96+	5.71	42.7	324	8.5	12.6		9.8	94.4	127.9	31.5	14.1	246	11.8	2.1	0.4
Gleyic Andosol	GA	A1	0-12	9.51	47.0	333	7.4	9.8	37.1	7.3	84.1	104.5	30.7	17.3	191	8.94	5.6	0.3
		A2	12-29	6.20	46.5	377	8.5	12.2	41.2	7.7	85.1	119.2	35	19.7	218	10.3	5.1	0.3
		Bw1	29-43	6.86	46.7	275	6.4	14.7	38.9	7.3	91	116.7	27.5	12.2	250	10.3	6.6	0.4
		Bw2	43-56	6.76	49.9	293	4.6	15.4	36.3	7.6	93.6	105.5	29.7	12.8	262	11.0	5.0	0.3
		C	56-64	2.70	23.5	649	4.0	7.4		10.3	78.3	119.2	38	38	144	8.92	6.3	0.3
		2Bw1	64+	3.26	40.2	480	1.8	21.1	22.9	8.5	112.5	59.2	22.5	22.5	196	16.6	5.3	0.3
Vitrisol	V	A	0-9	0.32	13.2	667	3.9	8.9	4.6			61.5	32.0	235	10.8	2.1	0.5	
		B/C	9-33	0.26	32.7	568	7.1	14.2	4.6			53	27.9	219	9.08	3.6	0.3	
		C	33+		8.87										137	4.98	-0.2	0.1
Parental basalt										20.6	72.6	103.8	74	32.1	164	5.89	3.6	0.3
Secondary minerals																		
Calcite	Iceland spar									0.38	0.02	0.009	336	0.01				
Calcite	11551									0.30	0.03	0.003	346	3.32		0.04	0.4	0.6
Calcite	11547									0.29	0.05	0.028	358	3.77		0.03	-1.0	0.5
Calcite	7869									0.33	0.39	0.133	363	0.19		0.04	5.4	0.5
Clay	11571									0.40	83.6	292	6.10	55.7		13.4	-8.4	0.2
Clay	21637									3.38	55.2	109	11.8	16.7		5.35	2.1	0.2
Heulandite	11607									9.52	68.1	0.010	29.5	0.05		0.83	7.9	0.3
Chabazite	15227									6.07	89.7	0.12	49.5	0.07		1.00	-0.2	0.2
Mesolite	21633									29.0	139.5	1.17	72.4	0.75		2.29	-4.9	0.2
Stilbite	11933									11.9	72.1	7.60	36.8	2.88		2.09	-1.8	0.1
Epistilbite	11613									11.9	97.9	0.057	57.2	0.03		2.57	-1.0	0.3
Scolecite	4221									16.8	102	0.060	43.1	0.02		2.33	5.3	0.6
Fe-oxyhydroxide	A6									11.0	110	283	55.6	27.7		14.1		
Leaches (% Li of total)			Exchangeable	Carbonate	Oxide	Residue												
HA 01			1.4	3.2	25.6	69.9												
H O6			3.4	4.0	20.6	72.0												
BA BW2			0.2	0.9	9.8	89.1												

940 Table 1. Bulk soil and secondary mineral data. All soil data aside from Li isotope ratios are
941 from Opfergelt et al., 2014. The final section shows the results for Li from sequential
942 leaching of three soils.

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Pore waters	Depth cm	pH	T °C	Li ng/ml	Na µg/ml	Mg µg/ml	Al ng/ml	K µg/ml	Ca µg/ml	Si µg/ml	Fe ng/ml	Cl µg/ml	$\delta^7\text{Li}$ ‰	2sd
2009														
HA H1	7.5	6.18	11.6	0.61	16.5	5.19	185	5.64	5.86	8.37	40.4	34.0	15.5	0.3
HA O1	62	3.64	11	3.02	7.87	3.60	10200	0.116	30.1	31.1	115	11	2.1	0.1
HA O2	83	3.71	10.9	3.02	8.95	3.56	9670	0.127	24.4	30.8	122	12	-1.4	0.2
HO1	6.5	4.34	11.3	4.17	9.17	4.16	316	0.390	9.88	17.40	13.8	9.0	14.6	0.5
HO5	67.5	4.78	11.7	3.02	16.1	3.33	15.8	0.318	4.29	13.90	519	22	14.3	0.3
HO6	72	5.48	11.6	5.94	19.6	10.9	90.8	0.746	11.2	18.3	25000	21	10.7	0.4
BA BW1	46	5.83	11.9	0.04	7.26	0.73	90.2	0.036	1.64	0.74	65.8	14.2	34.2	0.1
GA A1	6	5.64	13.1	0.12	7.83	1.46	106	0.326	3.78	4.89	38.6	25.9	28.5	0.3
GA BW1	36	5.90	13	0.08	7.53	1.20	9.6	0.041	3.54	7.95		13.6	34.2	0.2
GA 2BW1	64	5.88	12.9	0.08	6.93	2.00	11.6	0.011	5.94	13.80		12.1	29.5	0.4
2010														
HA H1	7.5	5.05	19.6	0.61	12.6	8.64	320	3.32	11.3	9.60	26.5	7.9	12.3	0.4
HA BW1	33	4.67	20.3	0.71	8.29	3.67	475	0.089	16.2	9.60	22.8	12.1	15.1	0.3
HA O1	62	3.99	18.9	1.80	6.70	4.13	6510	0.088	27.8	24.2	32.3	10.0	7.4	0.1
HA redox	75	3.98	18.5	2.97	8.54	3.60	7530	0.128	23.2	26.5	36.2	11.3	0.8	0.2
HA O2	83	3.95	18.3	2.80	9.52	4.34	7640	0.078	24.5	30.3	36.5	13.0	0.8	0.4
HO1	6.5	4.88	22.3	2.86	9.91	4.68	54.3	3.13	7.20	10.5	282	4	19.7	0.3
HO2	19.5	5.47	20.3	0.79	12.0	6.38	23.8	0.439	7.26	5.30	58.5	28.7	18.4	0.4
HO3	33	5.15	19.2	1.95	14.7	4.53	205	0.315	8.94	17.4	193	23	11.7	0.3
HO4	57.5			3.27	21.0	6.12	101	0.788	7.14	19.2	15300		14.1	0.6
HO6	72			7.48	25.4	17.1	45.7	1.72	15.2	18.8	20000		11.3	0.4
BA A1	10.5			0.06	6.38	1.51	4.93	1.23	2.13	5.90	12.1		28.4	0.6
BA A2	30.5	5.84	19.1	0.14	5.18	0.74	3.54	0.063	1.11	3.50	9.86	9.60	18.6	0.3
BA BW1	46	5.88	19	0.25	6.89	0.93	50.8	0.043	1.55	3.60	34.5	13.5	13.7	0.3
BA BW2	74	5.96	19.1	0.14	7.22	0.74	4.61	0.040	1.52	4.40	8.03	15.50	18.4	0.4
BA BW3	96	5.46	20	0.14	8.05	0.92	1.57	0.101	1.80	5.00	3.51	14.20	20.5	0.2
GA A1	6			0.05	6.87	1.79	17.3	2.318	3.75	5.70	13.6		31.6	0.4
GA A2	20.5	5.77	17.8	0.12	6.69	0.68	28.2	0.995	1.55	7.50	45.4	6.6	24.3	0.2
GA BW2	36			0.09	6.65	0.68	5.69	0.173	1.76	8.10	14.7		27.0	0.5
GA BW2	49.5	5.6	17.4	0.10	8.28	1.27	40.0	0.280	3.75	12.0	11.3	8.5	22.1	0.2
GA 2BW1	64	5.46	17.2	0.08	11.3	2.48	10.4	0.368	7.14	12.4	4.96	9.30	25.6	0.2

947 Table 2. Pore water data. All data aside from Li isotope ratios are from Opfergelt et al., 2014.

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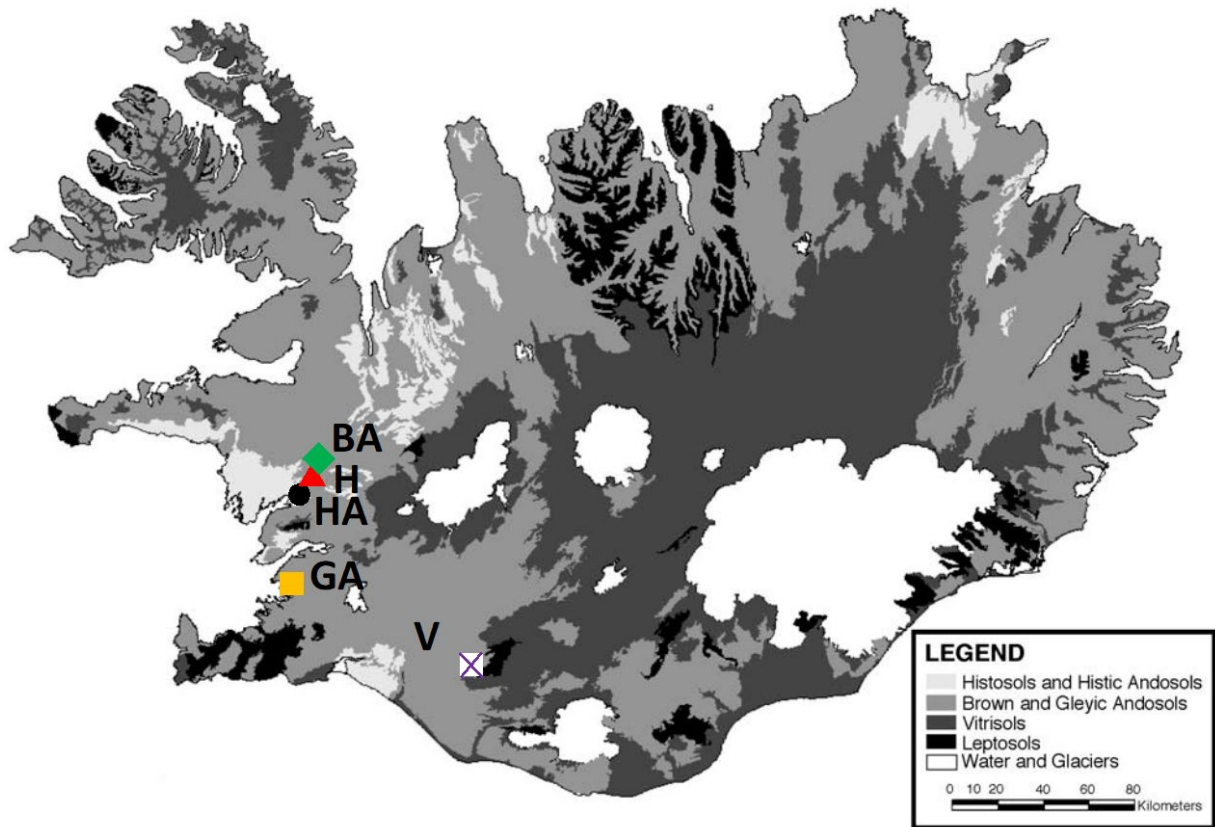
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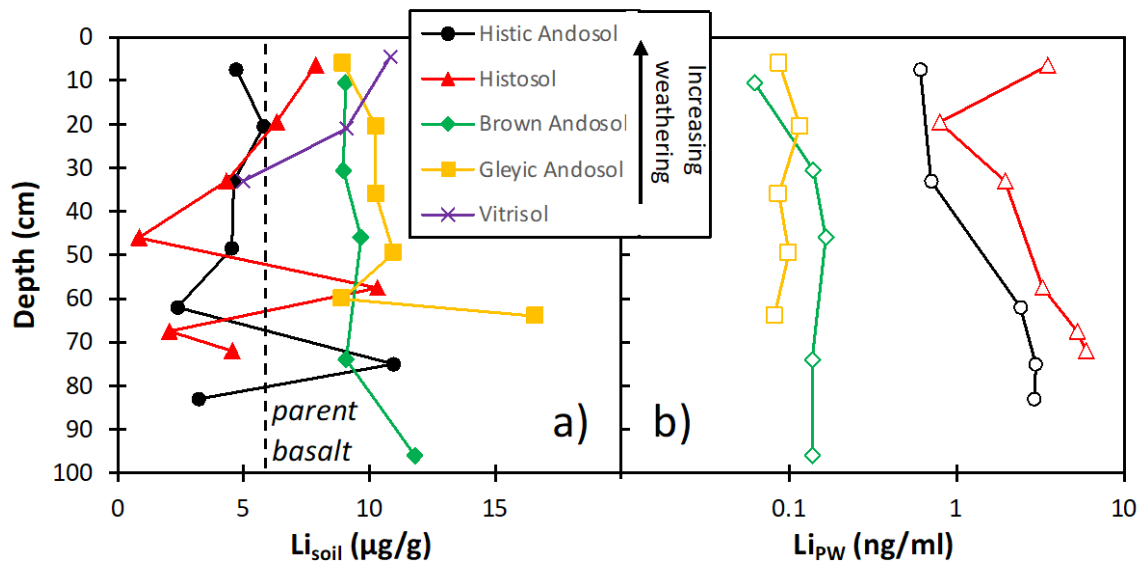
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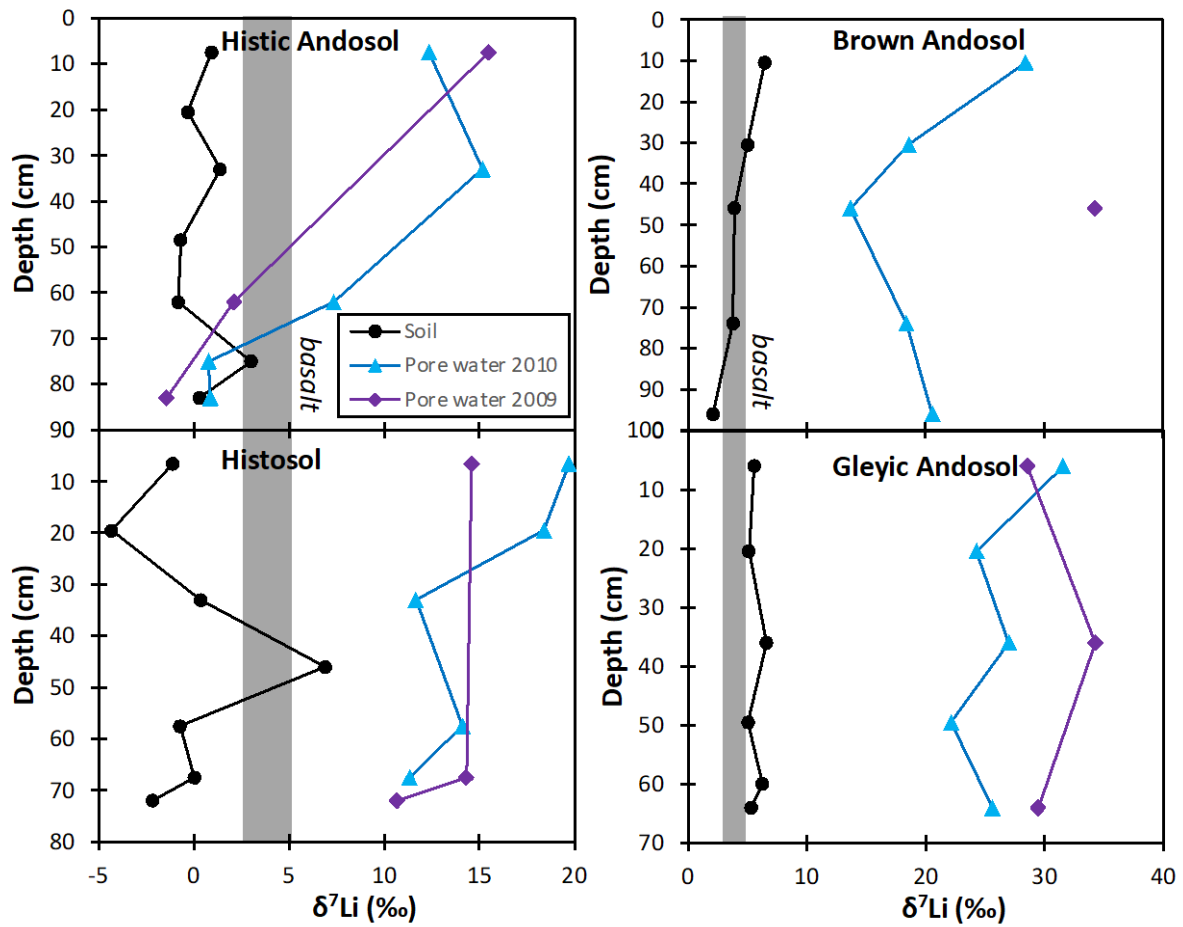
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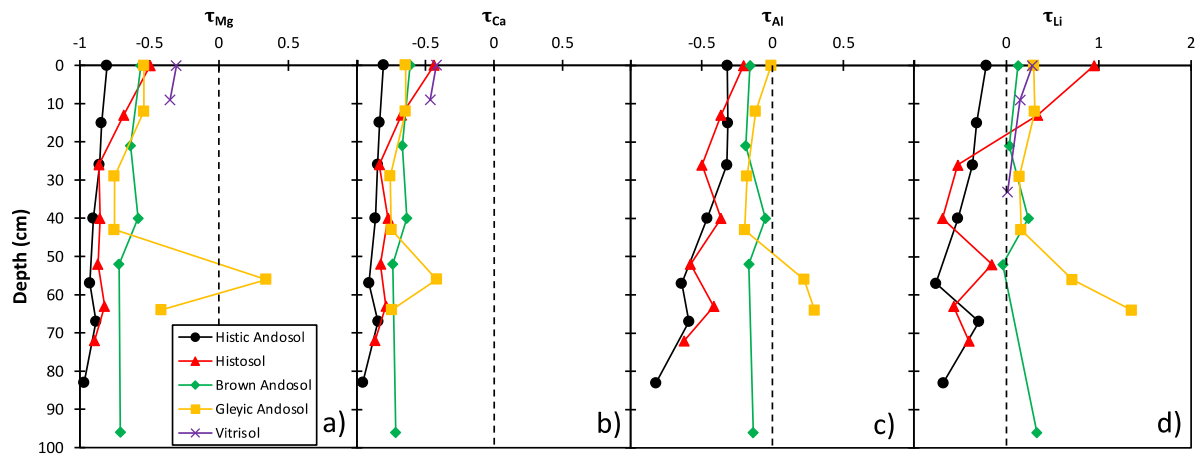
955 Figure 1. Sample locations on a soil map of Iceland. Soil symbols correspond to the key of
 956 Figure 2. Adapted from Opfergelt et al., 2014.



958 Figure 2. Depth profiles for Li concentrations for bulk soils and parent basalt depicted by the
 959 vertical dashed line (a), and pore waters (b). The pore water concentrations have been
 960 averaged from the two different collection years for this diagram (Table 2).
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965 Figure 3. Li isotope depth profiles for bulk soils, and the two different seasons of pore waters.
 966 In 2009, samples were taken in September (late summer), while in June (late spring) in 2010.
 967 The range of Icelandic basalt is depicted by the shaded area. Note the difference in horizontal
 968 axis scales.
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972 Figure 4. Tau values for the soils for the elements Mg, Ca, Al and Li. The dashed vertical
 973 lines are at tau = 0.

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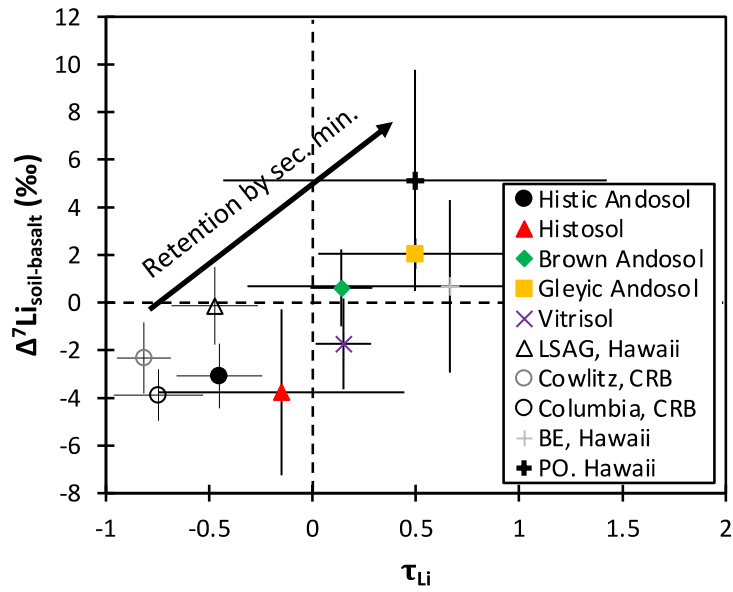
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984 Figure 5. Average τ_{Li} and $\Delta^7Li_{soil-basalt}$ values for each soil profile. The error bars represent the
 985 1sd spread in the data. Also shown are data from the Columbia River Basalts (Liu et al.,
 986 2013) and multiple sections from Hawaii (Li et al., 2020; Ryu et al., 2014). Due to high
 987 aeolian input, the topsoil from Li et al. (2020) has been removed from the averages.

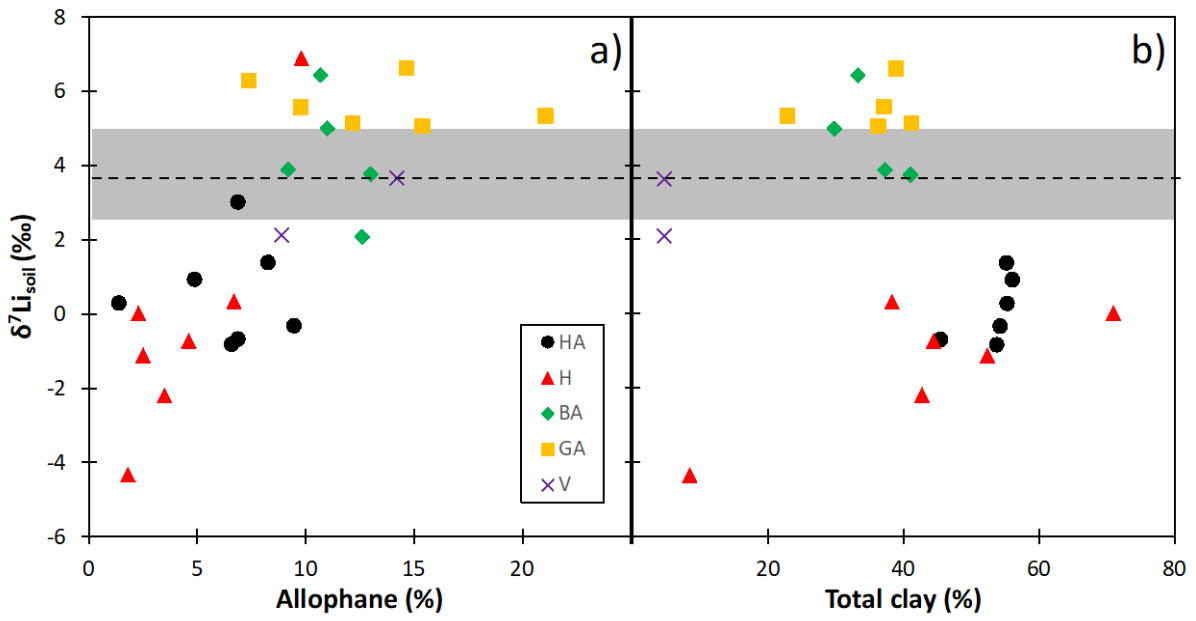
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994 Figure 6. Bulk soil Li isotopes as a function of allophane content (a) and total clay-sized
 995 content (b).

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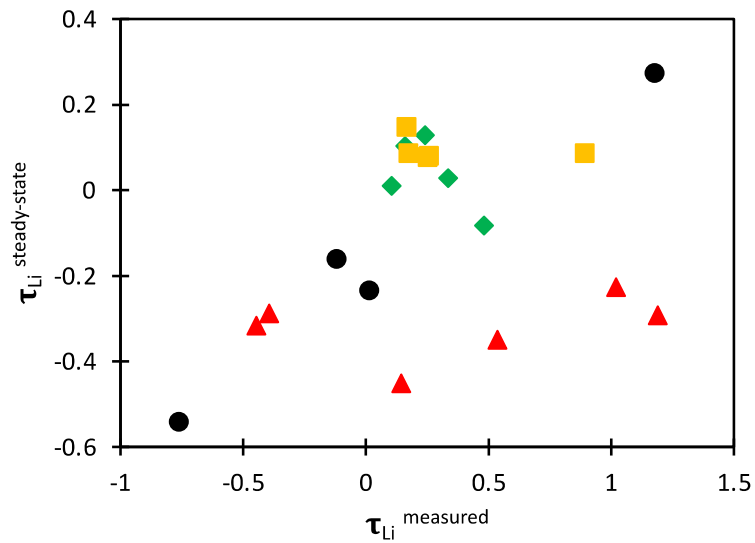
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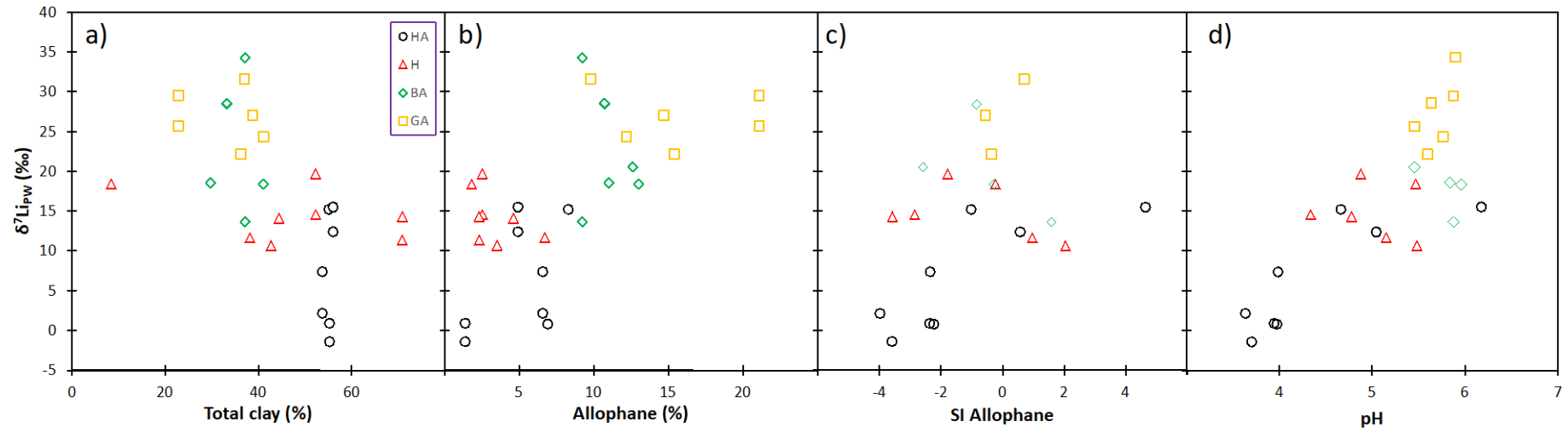
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1012 Figure 7. Comparison of measured τ_{Li} values, compared to steady-state τ_{Li} values calculated
 1013 from Li isotope ratios (Bouchez et al., 2013). Co-variation (as for HA soils) would suggest
 1014 that depletion or enrichment may be due to weathering intensity, while no correlation (as for
 1015 the other soils) suggest decoupling between processes controlling soil and present-day water
 1016 composition. See text for details. Symbols are the same as figure 6.

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1019 Figure 8. Pore water Li isotope ratios as a function of soil total clay content (a), soil allophane content (b), and the PHREEQC-derived allophane
1020 saturation index (c), and pH (d).

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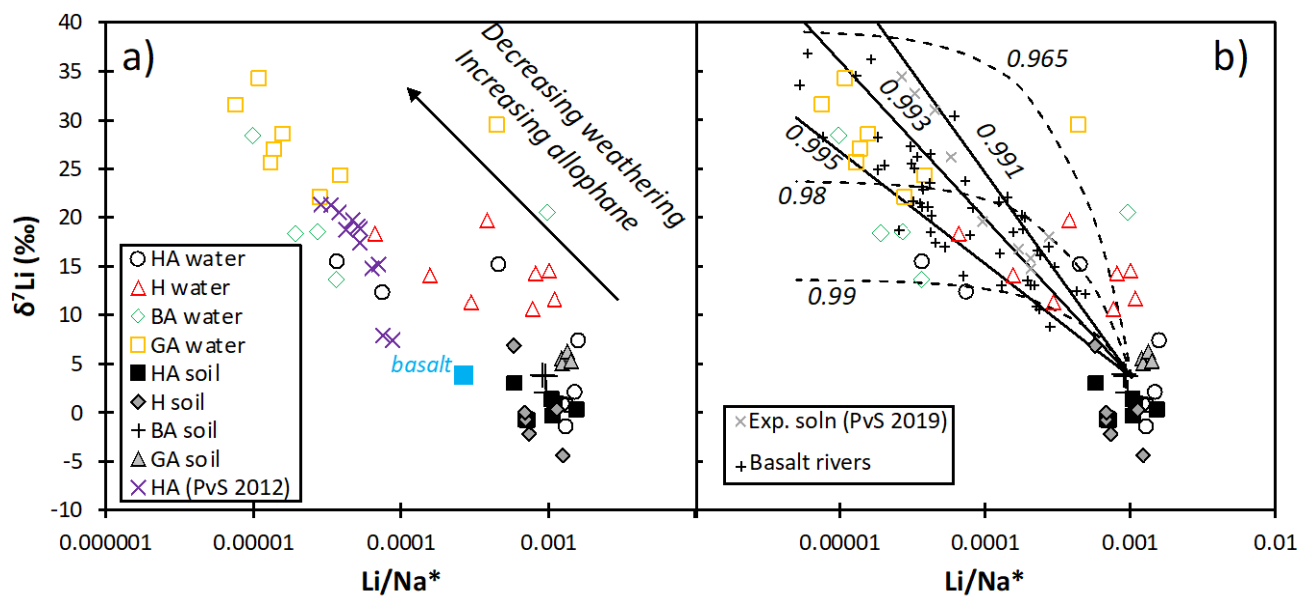
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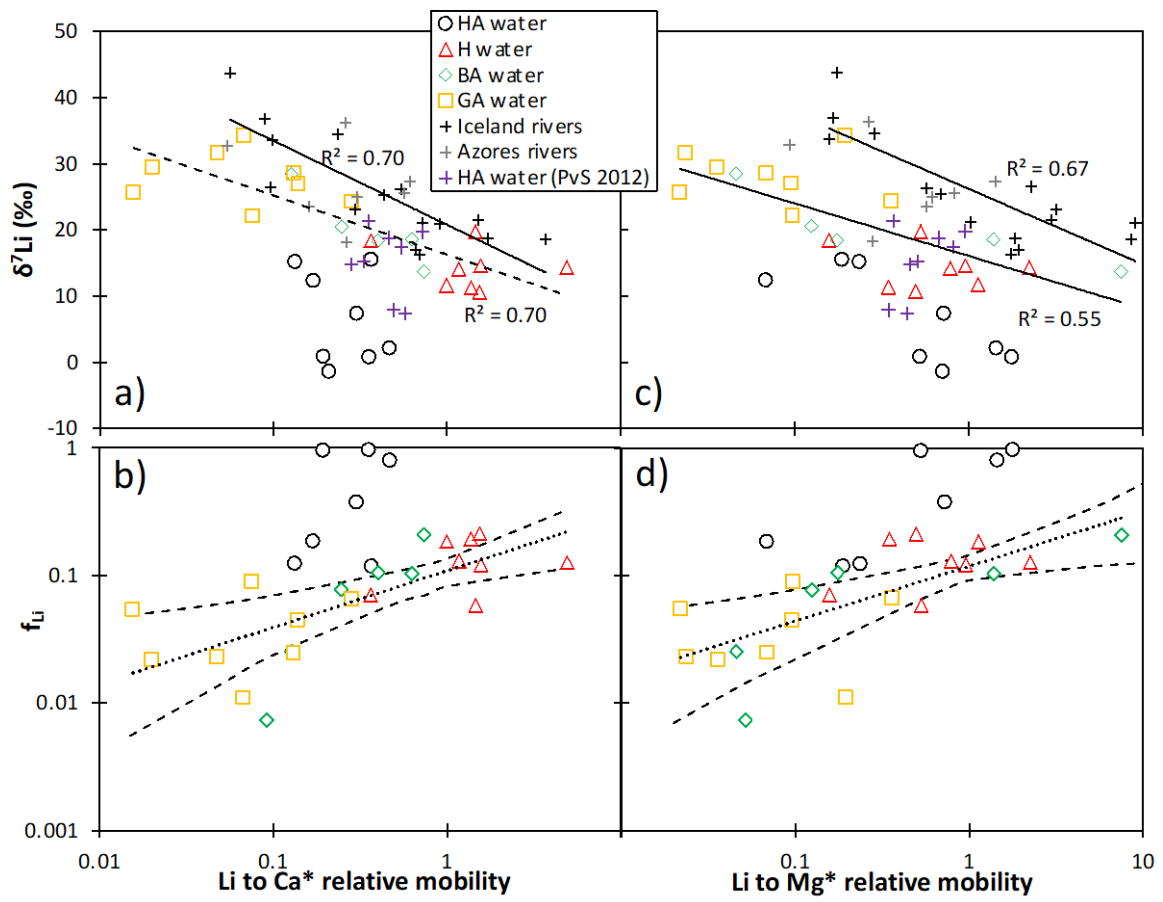
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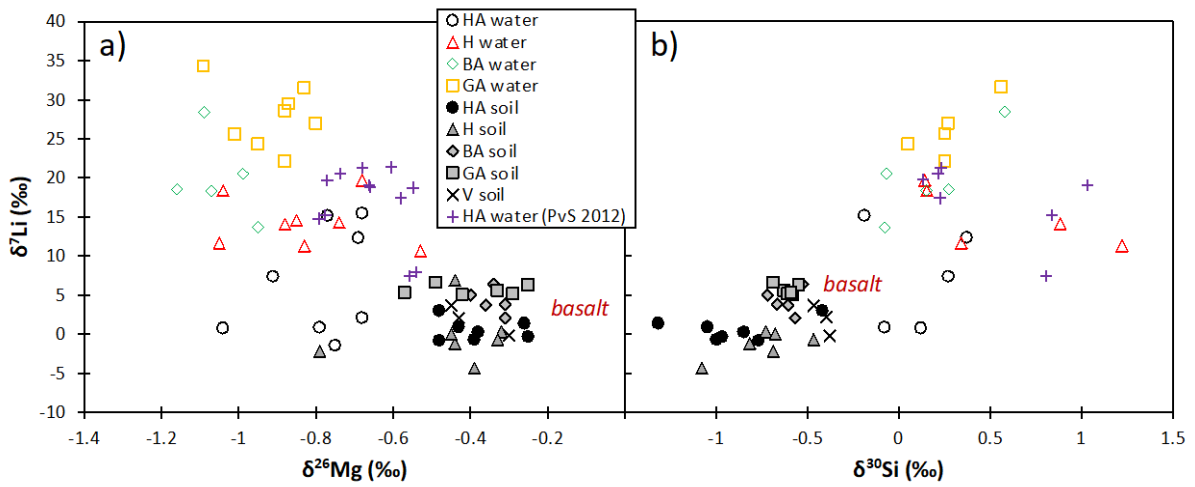


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1058 Figure 11. (a) Li isotope ratios as a function of Mg isotope ratios (Opfergelt et al., 2014;

1059 Pogge von Strandmann et al., 2012); (b) Li isotopes as a function of Si isotope ratios

1060 (Opfergelt et al., 2017b; Pogge von Strandmann et al., 2012).

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