

# 1 Investigation of subglacial weathering under the Greenland Ice Sheet using 2 silicon isotopes.

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38

## 39 Abstract

40 **Subglacial chemical weathering plays a key role in global silicate weathering budgets,**  
41 **contributing to the cycling of silicon (Si) in terrestrial and marine systems and the potential**  
42 **drawdown of carbon dioxide from the atmosphere. Here, we use data from two Greenland Ice**  
43 **Sheet (GrIS) catchments to demonstrate how Si isotopes from dissolved and amorphous**  
44 **particulate fractions ( $\delta^{30}\text{DSi}$  and  $\delta^{30}\text{ASi}$  respectively) can be used together with major ion data**  
45 **to assess the degree of secondary silicate weathering product formation and redissolution in**  
46 **subglacial environments. We compare a time-series of summer melt seasons from the two study**  
47 **sites, which differ in catchment size (~600 km<sup>2</sup> for Leverett Glacier (LG) and ~36 km<sup>2</sup> for**  
48 **Kiattuut Sermiat (KS)). Subglacial waters from LG have elevated Na<sup>+</sup> and K<sup>+</sup> ions in relation to**

49 **Ca<sup>2+</sup> and Mg<sup>2+</sup> ions, indicating a predominance of silicate weathering, whilst meltwaters from**  
50 **KS are characterised by carbonate weathering (hydrolysis and carbonation) throughout the**  
51 **melt season. Both catchments have mean  $\delta^{30}\text{DSi}$  values substantially lower than average riverine**  
52 **values (KS 0.41‰, LG -0.25‰, versus a global riverine mean of 1.25‰) and display a seasonal**  
53 **decline, which is more pronounced at LG. The  $\delta^{30}\text{ASi}$  values (discharge weighted mean values**  
54 **KS -0.44‰, LG -0.22‰) are lighter than the bedrock (mean values KS  $-0.18\pm 0.12\%$ , LG**  
55  **$0.00\pm 0.07\%$ ) in both catchments, indicating a secondary weathering product origin or leaching**  
56 **of lighter isotopes during initial weathering of crushed rock. When used in combination, the**  
57 **major ion and silicon isotope data reveal that the extent of silicate weathering and secondary**  
58 **phase redissolution are more pronounced at LG compared to KS. Contrasting weathering**  
59 **regimes and subglacial hydrology between catchments need to be considered when estimating**  
60 **the  $\delta^{30}\text{Si}$  composition of silica exported into polar oceans from the GrIS, with larger catchments**  
61 **likely to produce fluxes of lighter  $\delta^{30}\text{Si}$ . As larger catchments dominate freshwater export to the**  
62 **ocean, GrIS meltwater is likely to be very light in isotopic composition, and the flux is likely to**  
63 **increase with ice melt as the climate warms.**

64

## 65 **1. Introduction**

66 The weathering of silicate minerals plays an important role in biogeochemical cycles, producing  
67 dissolved silicate (DSi) and dissolvable amorphous silica (ASi) into terrestrial and marine systems,  
68 whilst consuming carbon dioxide (CO<sub>2</sub>) (Walker et al., 1981; Berner, 2003). DSi is generated through  
69 physical weathering and the subsequent chemical dissolution of silicate minerals, and then mobilised  
70 via fluvial hydrological systems. Primary weathering processes also lead to secondary mineral  
71 formation. Secondary weathering products, such as clays and ASi have varying solubility and may  
72 contribute DSi to the fluvial system through redissolution processes (Frings et al., 2016). The resultant  
73 DSi is converted to biogenic silica (BSi) by siliceous organisms, including diatoms, which are  
74 responsible for 35-70% of oceanic productivity (Nelson et al., 1995). Weathering processes,  
75 precipitation, and biogenic uptake result in fractionation of silicon isotopes, meaning isotopic analysis  
76 can provide insight into the weathering regimes and biological utilisation of Si within aquatic systems  
77 (De la Rocha et al., 1997; De La Rocha et al., 2000; Fontorbe et al., 2013; Frings et al., 2016).

78 Glaciation promotes physical and chemical weathering, with dynamic hydrological and microbial  
79 systems upon, within and beneath ice (Wadham et al., 2010; Tranter and Wadham, 2014). It is likely  
80 that weathering beneath large ice sheets is silicate mineral dominated (Stevenson et al., 2017), due to  
81 the enhanced residence times of subglacially stored waters, which result in exhaustion of carbonate  
82 minerals or saturation of meltwaters with respect to calcite (Wadham et al., 2010; Hindshaw et al.,  
83 2014; Michaud et al., 2016). This is evidenced by the ionic composition of meltwaters (high relative

84 proportions of Na<sup>+</sup> and K<sup>+</sup>) from the bed of the Greenland Ice Sheet (GrIS) (Graly et al., 2014) and the  
85 Antarctic Ice Sheet (AIS) (Michaud et al., 2016) compared to Alpine valley glaciers. Recently, there  
86 has been increased focus on the chemical composition of GrIS meltwaters in an attempt to improve  
87 the understanding of weathering processes in large ice sheet catchments and associated nutrient  
88 release (Graly et al., 2014; Hindshaw et al., 2014; Yde et al., 2014; Graly et al., 2017; Kohler et al.,  
89 2017). High fluxes of Si resulting from weathering processes have been reported from ice sheets. For  
90 example, Hawkings et al. (2017) estimated silica fluxes from GrIS to be approximately 50% of input  
91 from Arctic rivers (0.20 Tmol year<sup>-1</sup>). However, the mechanisms and control on silicate dissolution  
92 processes in ice sheet catchments are currently poorly understood, which is compounded by the  
93 difficulty of acquiring data over an entire melt season and across differing hydrological regimes in  
94 challenging environments. An improved understanding of the complex nature and extent of subglacial  
95 silicate weathering is necessary to fully understand and quantify silicon cycling on a global scale  
96 (Graly et al., 2014; Hindshaw et al., 2014; Yde et al., 2014).

97 Silicon isotopes (denoted by  $\delta^{30}\text{Si}$ ) are increasingly used as a tool for studying continental weathering  
98 (Ding et al., 2004; Georg et al., 2007; Opfergelt et al., 2013; Chemtob et al., 2015), especially in  
99 riverine systems (Ding et al., 2004; Cardinal et al., 2010; Ding et al., 2011; Fontorbe et al., 2013;  
100 Frings et al., 2016). Silicon has three stable isotopes; <sup>28</sup>Si, <sup>29</sup>Si and <sup>30</sup>Si with relative abundances of  
101 92.2%, 4.7% and 3.1% respectively (Ding et al., 2005). The isotopic composition is reported relative  
102 to international reference standard NBS-28, as a deviation from the ratio of <sup>28</sup>Si/<sup>30</sup>Si of the sample  
103 from the reference standard (Equation 1, Frings et al. 2016). The isotopic composition of the lower  
104 crust (-0.29±0.04‰), middle crust (-0.23±0.04‰) and bulk earth values (-0.29±0.08‰) are well  
105 characterised (Savage et al., 2010; Savage et al., 2013). Relatively large fractionation occurs during  
106 weathering processes as newly formed secondary weathering products are enriched with lighter  
107 isotopes (Andre et al., 2006; Frings et al., 2016). The range of DSi isotopic compositions in rivers  
108 draining from non-glaciated catchments is currently 0.4-3.4‰ (De La Rocha et al., 2000; Ding et al.,  
109 2004; Cardinal et al., 2010; Ding et al., 2011), with a global mean of 1.25±0.68‰ (Frings et al., 2016).  
110 Riverine DSi is therefore generally enriched in the <sup>30</sup>Si isotope compared with bulk silicate earth and  
111 suspended particulate matter, which is isotopically lighter on average (-0.18‰, Frings et al., 2016;  
112 Ding et al. 2004). Precipitated silica from the formation of secondary weathering or biological uptake  
113 is enriched in <sup>28</sup>Si, due to preferential uptake of the lighter isotope, resulting in an enrichment of <sup>30</sup>Si  
114 in the dissolved phase (De la Rocha et al., 1997; De La Rocha et al., 2000; Ziegler et al., 2002; Ding  
115 et al., 2004). However, whilst these generalisations about the fractionation of low temperature  
116 processes can be made, more definite fractionation factors for many environmental processes are still  
117 very uncertain (Frings et al., 2016).

118 Studies of Icelandic rivers have shown that catchments with greater glacial coverage have a lighter  
119 dissolved  $\delta^{30}\text{Si}$  composition ( $\delta^{30}\text{DSi}$ ) compared to non-glacial rivers (Georg et al., 2007). Opfergelt et

120 al. (2013) reported glacial rivers to have a mean composition of  $0.17 \pm 0.18\text{‰}$  compared to  $0.97 \pm$   
121  $0.31\text{‰}$  for non-glacial rivers. The dissolved phase exported from a large ice sheet catchment of GrIS  
122 has also been shown to have an isotopic composition lighter than the bulk bedrock ( $-0.25 \pm 0.12\text{‰}$   
123 compared to  $0.00 \pm 0.07\text{‰}$ , Hawkings et al., 2018). These studies indicate that subglacial processes  
124 could be influencing the isotopic composition of meltwaters. Si isotope systematics could potentially  
125 be used to provide insight into silicate dissolution and redissolution of secondary weathering products  
126 within the subglacial environment. However, we currently lack high resolution time series data to  
127 show whether these patterns are consistent across glacial catchments and to explain the mechanism  
128 driving the export of low  $\delta^{30}\text{Si}$ . We need to gain a better understanding of subglacial biogeochemical  
129 processes, and their impact on the  $\delta^{30}\text{Si}$  compositions of exported dissolved and particulate phases.  
130 This will allow us to quantify the impact of Si exported from GrIS on the wider silica cycle and make  
131 predictions about how export from large ice sheets may change over longer timescales.

132 Here we combine new  $\delta^{30}\text{Si}$  measurements of DSi and ASi phases with hydrogeochemical data from  
133 GrIS glacial catchments of contrasting scale over the summer melt season, in order to improve the  
134 current understanding of subglacial weathering processes.

## 135 **2. Methodology**

### 136 **2.1 Sampling locations**

137 We present hydrological and geochemical data from two polythermal- based, land terminating outlet  
138 glaciers from the GrIS, both of which have subglacial drainage systems that develop seasonally  
139 (Bartholomew et al., 2011). Data were collected from the proglacial river exiting Kiattuut Sermiat  
140 (KS;  $61.2^\circ\text{N}$ ,  $45.3^\circ\text{W}$ ; Fig. 1) from April to August 2013 and Leverett Glacier, (LG;  $67.06^\circ\text{N}$ ,  
141  $50.17^\circ\text{W}$ ; Fig. 1) from May to July 2015 to capture the composition of meltwaters exiting the glaciers  
142 during the summer ablation season. KS is a small coastal glacier, covering an area of  $36\text{km}^2$  and  
143 currently terminating in a proglacial lake of approximately  $0.5\text{km}^2$  (Hawkings et al., 2016; Dubnick et  
144 al., 2017). Calculations using discharge records find the turnover time of the proglacial lake to be  
145 relatively short once the glacial melt season begins (less than 24 hours from Day 157, Supplementary  
146 Fig. 1), indicating that the lake will likely not have a significant effect on the meltwater chemistry  
147 downstream. In comparison, LG is a much larger glacier, with a hydrologically active catchment of  
148 around  $600\text{km}^2$  (Cowton et al., 2012), which feeds into Watson River and then into the Davis Strait  
149 via the Søndre Strømfjord (Hawkings et al., 2016). The composition of bedrock beneath LG is  
150 dominated by Precambrian Shield crystalline gneiss and granite, which is representative of the  
151 majority of GrIS (Escher, 1976) and much of the bedrock that was covered by the Eurasian and North  
152 American Ice Sheets (Bouysse, 2014). The bedrock at KS is relatively similar to LG, but there are  
153 some potential differences. KS overlays the Gardar Province, containing basalts, trachytes and  
154 phonolites. This region may also include syenites from the intrusive rocks of the Julianhåb batholith

155 (Henriksen, 2009). However, previous work on  $\delta^{30}\text{Si}$  fractionation and elemental differentiation  
156 during high temperature processes in the mantle has shown limited isotope fractionation in  
157 comparison to weathering and biological processes (Savage et al., 2010).

158 We carried out hydrological monitoring of discharge, suspended sediment, pH and electrical  
159 conductivity using *in-situ* hydrochemical sensors (Fig. 1), as detailed by Kohler et al. (2017) and  
160 Hawkings et al. (2018). Water samples were collected at least once a day from just below the surface  
161 of the proglacial river using 1L Nalgene™ bottles (HDPE). Samples were filtered immediately using  
162 0.45 $\mu\text{m}$  cellulose nitrate membrane filters (Whatman®) mounted on a PS filtration stack (Nalgene™)  
163 and kept refrigerated in the dark until laboratory analysis. Samples for ASi concentrations and  
164 amorphous isotopic composition ( $\delta^{30}\text{ASi}$ ) were collected by filtering approximately 500ml of the  
165 same water sample collected for dissolved analysis through a 0.45 $\mu\text{m}$  cellulose nitrate membrane  
166 filters (Whatman®), retaining the sediment. Filters were kept refrigerated and in the dark until being  
167 gently air dried under a laminar flow hood prior to laboratory analysis.

168

## 169 **2.2 Major Ion Composition**

170 Water samples were analysed for major cation ( $\text{Na}^+$ ,  $\text{K}^+$ ,  $\text{Ca}^{2+}$  and  $\text{Mg}^{2+}$ ) and anion ( $\text{NO}_3^-$ ,  $\text{SO}_4^{2-}$  and  
171  $\text{Cl}^-$ ) concentrations by ion chromatography, with  $\text{HCO}_3^-$  estimated using the charge deficit (Tranter et  
172 al., 2002). Measurements were carried out using a Thermo Scientific™ Dionex™ capillary ICS-5000  
173 fitted with anion and cation columns (Hawkings et al., 2015; Dubnick et al., 2017).

174

## 175 **2.3 Silica concentrations**

176 DSi concentrations were determined spectrophotometrically using Flow Injection Analysis (FIA) on a  
177 LaChat 8500 series (QuikChem Method 31-114-27-1-D), as outlined by Hawkings et al. (2017). The  
178 limit of detection was 0.3 $\mu\text{M}$ . Precision and accuracy ( $\pm 0.54\%$  and  $\pm 1.9\%$ ) were determined by repeat  
179 measurements of a gravimetrically weighed 8.9 $\mu\text{M}$  standard (n=17). ASi was determined using an  
180 alkaline extraction method adapted from DeMaster (1981), as used by Hawkings et al. (2017). ASi  
181 was extracted from the suspended sediments collected on 0.45 $\mu\text{m}$  filters in the field. Briefly, 50ml  
182 0.1M  $\text{Na}_2\text{CO}_3$  was added to ~30mg (accurately weighed) of sediment at 85°C with 1ml aliquots taken  
183 at 2, 3 and 5 hours. Aliquots were neutralised immediately with 0.021M HCl and stored at 4°C until  
184 FIA analysis, using the same method as DSi concentrations. ASi (wt%) was calculated following  
185 DeMaster (1981).  $\text{Na}_2\text{CO}_3$  was preferred over NaOH extractions as it is the most commonly used  
186 alkaline extraction method of aquatic sediments, and is calibrated to clay minerals (Conley, 1998;  
187 Sauer et al., 2006; Hawkings et al., 2017). However, a 0.2M NaOH extraction method was required to  
188 extract the total reactive silica for silicon isotope analysis and both methods have been compared to  
189 ensure consistency between the ASi concentrations reported and the ASi extracted for silicon isotope  
190 analysis. The NaOH extraction produced similar or slightly lower concentrations compared to the

191 Na<sub>2</sub>CO<sub>3</sub> method (Hawkings et al., 2018). Saturation indices for ASi were calculated for each  
192 timepoint using hydrochemical data from KS and LG and Debye-Hückel ion interaction model in  
193 Geochemists Workbench® Student Edition.

194

## 195 **2.4 Silicon Isotope Composition**

196 All silicon isotope analysis was completed in the Bristol Isotope Group laboratories (University of  
197 Bristol) using a Thermo Scientific™ Neptune Plus™ High Resolution MC-ICP-MS and a standard-  
198 sample-standard bracketing procedure with Mg doping (100µl 10ppm Mg) to correct for mass bias  
199 (Cardinal et al., 2003; Hendry and Robinson, 2012). International reference standard NBS-28 (NIST  
200 RM8546, purified quartz sand) was used as the bracketing standard and isotope compositions are  
201 reported in terms of δ<sup>30</sup>Si (Equation 1). Samples were doped with 50-100µL 0.1M sulphuric acid  
202 (Romil-UpA) to ensure matrix match between sample and standard, to reduce the mass bias effects of  
203 anion loading (Hughes et al., 2011). Instrument blanks were <1% of <sup>28</sup>Si signal and typical internal  
204 reproducibility was ±0.08‰ for δ<sup>30</sup>Si (2SD) and ±0.04‰ for δ<sup>29</sup>Si (2SD). Long term reproducibility  
205 of reference standards Diatomite and LMG08 (sponge) are reported as +1.22 ±0.15‰ and -3.45  
206 ±0.14‰ (2SD) respectively. Mass dependent fractionation is demonstrated by a three-isotope plot of  
207 all samples analysed during the study (Supplementary Fig. 2) with a gradient of 0.5118 (Reynolds et  
208 al., 2007).

209

210 Sediment samples for δ<sup>30</sup>ASi composition were prepared by adding 1ml 0.2M NaOH per 0.1mg ASi  
211 and heating at 100°C for 1 hour. Samples were diluted and acidified with 8N HNO<sub>3</sub> and filtered  
212 through 0.22µm PES syringe filters. Bulk bedrock samples were processed using alkaline fusion  
213 (Georg et al., 2006) and the full procedure is detailed in Hawkings et al. (2018). Briefly, coarse,  
214 unsorted debris were crushed and ground to form a fine powder using a planetary ball mill (Fritsch  
215 Planetary Mono Mill Pulverisette 6). Debris was collected in front of the portal at LG and an ice cave  
216 into the front of KS. Samples were then furnaceed at 730°C with a NaOH pellet (approximately  
217 200mg), diluted with MQ and acidified with 8N HNO<sub>3</sub> once cool. Water samples were prepared for  
218 δ<sup>30</sup>DSi by preconcentrating the samples via evaporation to approximately 2ml sample, ensuring 2ppm  
219 of Si for analysis. All samples were then added to pre-cleaned BioRad exchange resin (AG50W-X12)  
220 columns and eluted with MQ water (Georg et al., 2006).

221

## 222 **3. Results**

### 223 **3.1 Defining periods of the melt seasons at KS and LG**

224 We categorise the melt season into three parts at KS, following Dubnick et al. (2017); Early Season,  
225 Transition Period and Late Season. These categories are defined by distinct changes in the proglacial  
226 river hydrochemistry. The Early Season is defined by low discharge and a lack of relationship

227 between solutes, which is unlikely to reflect a subglacial signal. The Transition Period likely reflects  
228 the time when the subglacial system became hydrologically connected and the bulk meltwaters  
229 included a subglacially stored contribution (Hawkings et al., 2016; Dubnick et al., 2017). This period  
230 includes a “Spring Event”, which is a time of high glacier velocity, associated with increases in  
231 subglacial water pressure discharge, suspended sediment and the divalent to monovalent cation ratio  
232 (Dubnick et al., 2017, Mair et al., 2004; Fig. 2). The Late Season had a bulk melt water signature that  
233 was distinct from the other periods analysed and is more similar to Alpine glacier geochemistry  
234 (Tranter et al., 2002, Tranter and Wadham., 2014).

235 We also categorise the melt season at LG into Early Season and Late Season, defined by differences  
236 in the hydrochemical data, with the transition being the first outburst event of the melt season. After  
237 this outburst event, the hydrochemistry of the meltwaters shifts towards those indicative of silicate  
238 mineral dissolution, as longer residence time waters are flushed.

### 239 **3.2 Dissolved major ions, DSi and $\delta^{30}\text{DSi}$**

240 The ratio of  $(\text{Ca}^{2+} + \text{Mg}^{2+})$  to  $(\text{Na}^{+} + \text{K}^{+})$  (divalent to monovalent cations, reported hereafter as D:M) has  
241 traditionally been used to indicate the amount of silicate weathering in relation to carbonate  
242 weathering (Tranter et al., 2002; Wadham et al., 2010). Elevated monovalent ion concentrations in  
243 relation to divalent concentrations are associated with enhanced silicate mineral weathering. The  
244 dissolved ion composition of glacial runoff from the two catchments changed significantly with  
245 seasonal evolution of the subglacial drainage system (Fig. 3). The discharge-weighted mean (Qwt)  
246 D:M ( $\mu\text{eq L}^{-1} : \mu\text{eq L}^{-1}$ ) at LG was 1.36 compared to 6.91 at KS, with the D:M decreasing at LG as the  
247 melt season progressed, from 5.18 to 1.44, compared to an increase in the D:M of a smaller magnitude  
248 at KS (6.57 to 7.72, Fig. 3).  $\text{Ca}^{2+}$  was the dominant cation in KS over the entire melt season, consistent  
249 with previously documented meltwaters from Alpine and Polar glaciers. However, there was a shift  
250 from  $\text{Ca}^{2+}$  to  $\text{Na}^{+}$  as the dominant cation as the melt season progressed at LG (Ca/Na molar ratios  
251 started at 3.23 and decreased to 0.56 during outburst events, Fig. 4).

252 DSi concentrations in LG meltwaters decreased as the season progressed due to dilution from  
253 supraglacial meltwaters ( $56.9\mu\text{M}$  to  $9.21\mu\text{M}$ ). The discharge weighted mean DSi concentration was  
254  $20.8\mu\text{M}$ , similar to previously reported concentrations (Hindshaw et al., 2014; Hawkings et al., 2017).  
255 KS had a slightly higher discharge weighted mean DSi of  $22.2\mu\text{M}$ , with a range of  $14.8\mu\text{M}$  to  $41.8$   
256  $\mu\text{M}$ . Concentrations also decreased as the subglacial hydrological drainage system developed from  
257 inefficient to efficient drainage pathways (Supplementary Table 3 and 4).

258 There was a substantial decline in  $\delta^{30}\text{DSi}$  at LG as the melt season progressed and the subglacial  
259 system became hydrologically connected, with drainage of more isolated regions of the bed further up  
260 catchment (Fig. 5d). The discharge weighted mean value of  $\delta^{30}\text{DSi}$  at LG ( $-0.25 \pm 0.12\text{‰}$ ) was lighter  
261 than the previous riverine average  $\delta^{30}\text{DSi}$  composition ( $+1.25\text{‰}$ , Frings et al., 2016), and studies of

262 glacially fed rivers in Iceland ( $+0.17 \pm 0.18\%$ , Opfergelt et al., 2013). The lightest values measured at  
263 LG are more comparable to long residence time groundwaters (up to  $-1.42\%$ , Georg et al., 2009). KS  
264 had a higher  $\delta^{30}\text{DSi}$ , with a discharge weighted mean of  $+0.41 \pm 0.10\%$ , although this value is also  
265 below the average riverine silicon isotope composition (Frings et al., 2016). The  $\delta^{30}\text{DSi}$  at KS also  
266 declined after the connection of the subglacial hydrological system (“Spring Event”, Hawkings et al.,  
267 2016, Mair et al. 2004) but the decline was not as substantial as that seen at LG (Fig. 5c).

### 268 **3.3 Saturation Indices**

269 Bulk meltwaters in both catchments were highly undersaturated with respect to ASi throughout the  
270 melt season (LG  $\text{SI}_{\text{ASi}} = -1.04$  to  $-1.85$  and KS  $\text{SI}_{\text{ASi}} = -1.12$  to  $-1.73$ ). The lightest  $\delta^{30}\text{DSi}$  composition  
271 occurred when the proglacial river was most undersaturated with respect to ASi at both catchments  
272 (Supplementary Fig. 3).

### 273 **3.4 Bedrock, SPM, ASi concentration and $\delta^{30}\text{ASi}$ .**

274 Suspended particulate matter (SPM) concentrations in bulk meltwaters generally increased over time  
275 in both catchments, as increasing amounts of sediment were entrained subglacially (Fig. 2). LG ASi  
276 and SPM concentrations were significantly higher than KS (Table 1), with peak SPM coinciding with  
277 subglacial outburst events (Hawkings et al. 2016, Fig. 2). Particles at LG also had a higher relative  
278 proportion of extractable ASi, contributing to the elevated ASi concentrations (LG Qwt mean =  
279  $0.73\text{wt}\%$ , compared to KS Qwt mean =  $0.23\text{wt}\%$ ). However, KS has a higher specific discharge than  
280 LG, so the yield of Si per  $\text{km}^2$  is higher at KS. We estimate a mean annual ASi yield of  $1.75 \times 10^4 \text{ kg}$   
281  $\text{km}^{-2}$  from KS (Day 128 - 221) compared to an estimated mean yield of  $1.24 \times 10^4 \text{ kg km}^{-2}$  from LG  
282 (Day 135 – 210).

283 The  $\delta^{30}\text{ASi}$  composition of SPM in bulk runoff was lighter at KS compared to LG, with little variation  
284 at either site over the monitoring period (discharge weighted mean of  $-0.47 \pm 0.06\%$  compared to -  
285  $0.22 \pm 0.06\%$ , Fig. 5). These values were lower than the local bedrock (KS  $-0.18 \pm 0.06\%$ , LG  $0.00 \pm$   
286  $0.07\%$ ) and bulk suspended sediment (KS  $-0.32 \pm 0.12\%$ , LG  $-0.09 \pm 0.07\%$ ).

## 287 **4. Discussion**

### 288 **4.1 Conceptual model of subglacial hydrology**

289 Differences in drainage system characteristics of glaciers at different spatial scale may result in  
290 contrasting chemical weathering environments and water export mechanisms (Wadham et al., 2010;  
291 Graly et al., 2014). Subglacial drainage systems of GrIS catchments likely include hydrological  
292 elements that exist for the majority of the year and those that develop over time and shut down during  
293 winter (Dubnick et al., 2017). The progression of the melt season is accompanied by an evolution  
294 from slow-inefficient distributed to efficient (and potentially channelised) drainage systems as more



295 supraglacial meltwater reaches the bed (Bartholomew et al., 2011). Previous studies have  
296 demonstrated that the seasonal evolution of subglacial hydrology impacts nutrient export dynamics  
297 via the connection of chemically distinct, solute sources at the glacier bed (Bartholomew et al., 2011;  
298 Bhatia et al., 2013; Hawkings et al., 2016; Dubnick et al., 2017; Kohler et al., 2017). LG has a  
299 hydrologically active catchment of  $\sim 600\text{km}^2$ , is 80km long and has inland ice of low topography,  
300 favouring the formation of large supraglacial lakes (often  $\sim$  kms in diameter, Hoffman et al., 2011).  
301 These supraglacial lakes can drain rapidly ( $<2$ hours) through moulins to the glacier bed as a result of  
302 hydrofracturing (Das et al., 2008; Bartholomew et al., 2011). Such drainage events can result in  
303 outburst events being recorded in the proglacial hydrochemical records (Bartholomew et al., 2011;  
304 Hawkings et al., 2015). Proglacial river discharge, suspended sediment and electrical conductivity  
305 rapidly rise (Bartholomew et al., 2011, Hawkings et al., 2016; Fig. 2), as subglacially stored waters  
306 and sediment become hydrologically connected and flushed downstream by incoming supraglacial  
307 water (Bartholomew et al., 2011; Cowton et al., 2012; Chandler et al., 2013). Subglacial hydrological  
308 evolution from inefficient to efficient drainage at the bed progresses from marginal to inland locations,  
309 with artificial tracer work showing efficient drainage existing up to 40km from the margin by late July  
310 (Chandler et al., 2013). This evolution is further supported by the seasonal progression of the  $^{14}\text{C}$  age  
311 of exported POC at LG (Kohler et al., 2017). The inland progression of the subglacial system could  
312 mean that increasingly isolated meltwaters drain from the bed, since the interval between basal  
313 flushing events via supraglacial lake drainage or moulins is longer in more inland locations  
314 (potentially 10 months) compared to marginal locations (Harper and Humphrey, 1995). Residence  
315 time differences between marginal and the most isolated inland waters are therefore likely to exist,  
316 especially if similar regions of the bed are flushed annually (Fig. 6), with implications for weathering  
317 and redissolution processes.

318 KS is only  $\sim 16\text{km}$  in length and there is comparatively little altitudinal difference from snout to the  
319 top of the ablation area. Supraglacial lakes do not form at KS, thus some more isolated parts of the  
320 bed may not be annually flushed by surface-to-bed water flow. The subglacial drainage system  
321 development at KS is driven entirely by the progression of the snowline and the consequent opening  
322 of new moulins and crevasses for surface to bed water flow, similar to smaller Arctic and Alpine  
323 glaciers. Thus, the subglacial drainage system develops from inefficient distributed drainage, where  
324 water follows a tortuous flow path, to an efficient, channelised system as the ablation season  
325 progresses and meltwater inputs increase (Fig. 6, Tranter et al., 2002, Nienow et al., 2014). The  
326 absence of supraglacial lake drainage and the smaller catchment size will shorten average residence  
327 times of water emerging as bulk runoff. Any seasonal variation in subglacial water residence times is  
328 likely on the order of weeks rather than months. We propose that the full length of KS could be  
329 compared with the evolution of the first 10-20km of subglacial hydrology at LG prior to the onset of  
330 outburst events, as demonstrated by the relationship between discharge and  $\delta^{30}\text{DSi}$  (Fig. 5b). As there

331 is a limited input of long residence time stored water with differing geochemical composition at KS,  
332 the chemical composition of the proglacial stream reflects a carbonate dominated weathering regime  
333 (Tranter et al., 2002).

#### 334 **4.2 Differences in weathering regimes**

335 Glacier size is hypothesised to have a major impact on chemical weathering via its influence upon  
336 hydrological flow path length, and hence water residence times (Wadham et al., 2010). Our data show  
337 clear differences in the major ion composition of bulk meltwaters draining from the study glacial  
338 catchments (Figs. 3 and 7), indicating contrasting subglacial chemical weathering regimes. In larger  
339 catchments silicate mineral dissolution is enhanced, as subglacially stored meltwaters reach saturation  
340 with respect to calcite, due to the long residence times and subglacial isolation (Wadham et al., 2010;  
341 Hawkings et al., 2016). Carbonate hydrolysis was previously believed to dominate over carbonation  
342 in subglacial environments, due to the limited atmospheric connectivity, resulting in enhanced  
343 concentration of divalent ions in meltwaters ( $\text{Ca}^{2+} + \text{Mg}^{2+}$ ; Equation 2, Tranter et al., 2002). By  
344 comparison, silicate hydrolysis results in elevated concentrations of monovalent ions ( $\text{K}^+ + \text{Na}^+$ ;  
345 Equation 3, Tranter and Wadham, 2014).

#### 346 Leverett Glacier

347 At LG, the progressive evolution in the D:M ratio is consistent with a shift towards silicate dominated  
348 mineral weathering as the melt season progresses (Fig. 3). This trend in D:M is also consistent with  
349 the Na-normalised molar ratios, which can be used to compare the hydrochemical signature to silicate  
350 and carbonate endmembers in mixing diagrams (Gaillardet et al., 1999). Fig. 4 shows the evolution of  
351 the LG towards the silicate endmember as the melt season progresses, with the outburst events lying  
352 closest to this endmember. This suggests that isolated meltwaters characterised by long residence  
353 times become hydrologically connected to a fast, efficient drainage systems and are exported to the  
354 ice margin.

355 Subglacial meltwaters in inland regions may only be flushed by surface melt (e.g. via lake drainage  
356 and moulins) after ~10 months when the snowline retreats this far from the ice margin and inland  
357 hydrological systems become connected. During the intervening period, these subglacial meltwaters  
358 are inferred to undergo enhanced silicate dissolution (Wadham et al., 2010; Chandler et al., 2013;  
359 Graly et al., 2014). Hydrolysis of silicates also increases the pH within subglacial waters (Equations 2  
360 and 3, Fig. 2), and these higher pH conditions further enhance the dissolution of aluminosilicate  
361 minerals (Georg et al., 2009; Tranter and Wadham, 2014). Evolution towards low D:M ratios, and  
362 elevated pH is particularly pronounced from Day 170 onwards in 2015 at LG after the first subglacial  
363 outburst event ( $Q_{wt}$  pH after first outburst event 8.71, Fig. 2).

364 Coupled carbonate dissolution and sulphide oxidation is a more important process, during the Early  
365 Season at LG. This importance is illustrated by gradients of  $\text{Ca}^{2+}+\text{Mg}^{2+}$  versus  $\text{SO}_4^{2-}$  (Fig. 7a) and of  
366  $\text{Ca}^{2+}+\text{Mg}^{2+}$  versus  $\text{HCO}_3^-$  (Fig. 7b) close to 2 (1.93 and 1.50 respectively, Tranter et al., 2002), which  
367 closely resemble molar equivalent ratios of 2:1 for  $\text{Ca}^{2+}+\text{Mg}^{2+}$  and  $\text{SO}_4^{2-}$ , and  $\text{Ca}^{2+}+\text{Mg}^{2+}$  and  $\text{HCO}_3^-$ ,  
368 according to Equation 5. These findings are similar to those found by Graly et al. (2017), but are not  
369 seen in a study which focused upon the middle to late part of the melt season at LG (Hindshaw et al.,  
370 2014). We find higher concentrations of  $\text{SO}_4^{2-}$  in 2015 during the Early Season when discharge was  
371 lower (Supplementary Fig. 4), which we can attribute to sulphide oxidation, highlighting the  
372 importance of categorising the development of the melt season due to potentially large seasonal  
373 differences. Furthermore, during the Early Season, the sulphate mass fraction (SMF, Equation 6) is  
374 higher than later in the season (Supplementary Fig. 5), indicating a larger proportion of protons result  
375 from sulphide oxidation. However, a SMF of  $<0.5$  throughout the melt season indicates carbonation is  
376 overall more important than sulphide oxidation as a proton supplier.

377 Following the first outburst event (Late Season), the gradients of  $\text{Ca}^{2+}+\text{Mg}^{2+}$  v  $\text{SO}_4^{2-}$  and  $\text{Ca}^{2+}+\text{Mg}^{2+}$   
378 v  $\text{HCO}_3^-$  became shallower ( $\text{Ca}^{2+}+\text{Mg}^{2+}:\text{SO}_4^{2-} = 1.50:1$ ,  $\text{Ca}^{2+}+\text{Mg}^{2+}:\text{HCO}_3^- = 0.79:1$ ). The SMF also  
379 decreases ( $\sim 0.45$  to  $\sim 0.25$ ), indicating protons are largely provided from carbonation reactions. It is  
380 likely that silicate dissolution via hydrolysis and carbonation is occurring in the Late Season at LG  
381 (Wadham et al., 2010), especially when considering the concomitant increase in the gradient of the  
382 lines of best fit for associations between  $\text{HCO}_3^-$  and  $\text{SO}_4^{2-}$  (Fig. 7c and embedded Table) from Early to  
383 Late season at LG (0.96 to 1.17). Carbonation of silicates and carbonates consumes  $\text{CO}_2$  in solution  
384 and result in formation of  $\text{HCO}_3^-$ , which, combined with silicate hydrolysis, also helps to explain the  
385 increasing pH as the melt season progresses. We expect that this increase in pH is the result of  
386 increased hydrolysis and carbonation of silicates, due to the decline of  $\text{Ca}^{2+}+\text{Mg}^{2+}:\text{HCO}_3^-$  ratios as the  
387 season progresses, indicating that the addition of  $\text{HCO}_3^-$  is occurring in the absence of  $\text{Ca}^{2+}+\text{Mg}^{2+}$ .

388 The meltwater geochemistry therefore suggests that LG begins as a system with the imprint of  
389 sulphide oxidation coupled to carbonate dissolution (SMF =  $\sim 0.45$ , Equation 5). As the season  
390 progresses and more isolated waters are exported, there is an increased importance of silicate  
391 weathering linked to carbonation reactions (SMF =  $\sim 0.2$ ).

### 392 Kiattuut Sermiat

393 Our conceptual model of the hydrology at KS leads to predictions of a relatively consistent subglacial  
394 weathering regime once an efficient drainage system has developed. The source of solute in the  
395 meltwaters at KS appears fairly consistent over the melt season. D:M ratio and mixing diagrams at KS  
396 indicates that the system is dominated by carbonate weathering (Figs. 3, 4; also see Dubnick et al.  
397 2017). We can use the ionic ratios to interrogate these relationships further and assess differences  
398 between Early Season, Transition Period and Late Season.

399 During the Early Season at KS there is no significant relationships between  $\text{Ca}^{2+}+\text{Mg}^{2+}$  and  $\text{HCO}_3^-$   
400 versus  $\text{SO}_4^{2-}$ . We would expect the higher  $\text{SO}_4^{2-}$  concentrations to be a result of sulphide oxidation.  
401 Sulphide oxidation would usually be coupled with carbonate dissolution in glacial systems, which  
402 should be reflected in molar equivalent ratios (Equation 5). However, the discharge is low at this time,  
403 so it is likely that the proglacial river reflects groundwaters diluted with some snowmelt and/or  
404 precipitation. Groundwater flow may be torturous, impacting upon the ionic ratios, and it is also likely  
405 that these waters are more affected by atmospheric deposition than later in the season when discharge  
406 is much greater (Bhatia et al., 2013). In the Early Season there are also longer turnover times of lake  
407 water due to the lower discharge entering the lake, which results in slower flushing of system  
408 (Supplementary Fig. 1).

409 It is during the Transition Period that the gradients for all the major ion relationships most closely  
410 match those during the Early Season at LG (Fig. 7). There was an increase of  $\text{HCO}_3^-$  and  $\text{Ca}^{2+}+\text{Mg}^{2+}$   
411 in relation to  $\text{SO}_4^{2-}$  (Fig. 7a and 7c), which could be evidence of the connection of the subglacial  
412 system to the proglacial hydrological system. As the dissolution of freshly comminuted glacial  
413 sediments preferentially release  $\text{HCO}_3^-$  and  $(\text{Ca}^{2+}+\text{Mg}^{2+})$  (via hydrolysis of trace carbonates), it would  
414 be expected that the  $\text{HCO}_3^-$  and  $\text{Ca}^{2+}+\text{Mg}^{2+}$  ratios with respect to  $\text{SO}_4^{2-}$  would increase. The elevated  
415  $\text{HCO}_3^-$  could also result from the microbial oxidation of organic carbon (Wadham et al., 2010). The  
416 higher intercepts of  $\text{HCO}_3^-$  v  $\text{SO}_4^{2-}$ , and  $(\text{Ca}^{2+}+\text{Mg}^{2+})$  v  $\text{SO}_4^{2-}$  compared to LG, suggest rapid  
417 acquisition of  $(\text{Ca}^{2+}, \text{Mg}^{2+})$  and  $\text{HCO}_3^-$  in relation to  $\text{SO}_4^{2-}$ , potentially due to carbonate hydrolysis.  
418 Also, during the Transition Period, the  $(\text{Ca}^{2+}+\text{Mg}^{2+})$ :  $\text{HCO}_3^-$  ratio increased to 1.50 (Fig. 7b), and the  
419 SMF increased to ~0.18 (Supplementary Fig. 5). Collectively, these observations indicate some  
420 influence from carbonate dissolution coupled to sulphide oxidation (Equation 5), similar to the Early  
421 Season meltwaters at LG.

422 We see a prevalence of carbonate over silicate weathering during the Late Season at KS, providing  
423 evidence of continuous availability of carbonate minerals within the subglacial system and a lack of  
424 calcite saturation. Relationships close to 1:1 for  $(\text{Ca}^{2+}+\text{Mg}^{2+})$ :  $\text{HCO}_3^-$  (Fig. 7b) show that carbonation  
425 of carbonates and carbonate hydrolysis are the dominate weathering reactions throughout the melt  
426 season, as a 1:1 ratio is expected from Equation 2.

427 The values measured at KS are similar to bulk meltwaters of Alpine glaciers studied, although the  
428  $\text{SO}_4^{2-}$  concentrations in the outflow from KS are lower (Tranter et al., 2002). The overall major ion  
429 relationships indicate a shift from carbonate hydrolysis (Equation 2) and coupled carbonate  
430 dissolution-sulphide oxidation (Equation 5) to carbonation of carbonates as the melt season progresses,  
431 with little evidence of enhanced silicate weathering.

#### 432 **4.3 Catchment hydrology as a driver of $\delta^{30}\text{DSi}$ seasonal variability**

433 By combining our conceptual model of subglacial hydrology at LG and KS with the observed major  
434 ion ratio chemistry, we can begin to understand the geochemical drivers behind variations in  $\delta^{30}\text{Si}$   
435 composition of the meltwaters in each catchment. An increased predominance of silicate dissolution  
436 as the melt season progresses at LG should theoretically result in higher  $\delta^{30}\text{DSi}$ , as silicate weathering  
437 and the formation of secondary weathering products preferentially incorporate the lighter isotopes into  
438 the newly formed solid (De La Rocha et al., 2000; Frings et al., 2016). While this pattern is observed  
439 in non-glacial regimes, the lowest  $\delta^{30}\text{DSi}$  values measured at LG coincided with the most pronounced  
440 silicate weathering signals, and likely, by association, the longest residence time waters. We  
441 hypothesise that the low  $\delta^{30}\text{DSi}$  values measured at LG after Day 170 reflect the dissolution of  
442 isotopically light ASi and secondary weathering products in subglacially stored waters and/or the  
443 dissolution of isotopically light fresh mineral surface layers formed by enhanced physical weathering  
444 (Hawkings et al., 2018). As the melt season progressed, the hydrologically active part of the  
445 catchment retreated further inland and more isolated subglacial waters became hydrologically  
446 connected (Hawkings et al., 2015), resulting in decreasing  $\delta^{30}\text{DSi}$  (Fig. 5b) and D:M ratios as  
447 discharge rose (Fig. 5a). The declining D:M ratios indicate a move to increasingly dominant silicate  
448 weathering and redissolution of secondary weathering products from the more isolated subglacial  
449 system or longer residence time waters, resulting in more dissolution of these finely ground  
450 weathering crusts, which results in the lowering of the  $\delta^{30}\text{DSi}$  composition.

451 Some decline in  $\delta^{30}\text{DSi}$  also occurred during the Transition Period at KS (Fig. 5c), which was likely  
452 linked to the opening of subglacial hydrological pathways and flushing of a formerly distributed  
453 drainage system. We would expect redissolution of secondary weathering products to occur at KS due  
454 to the high pH and the undersaturation of ASi (Crompton et al., 2015, Supplementary Fig. 3).  
455 However,  $\delta^{30}\text{DSi}$  composition at KS is consistently higher than  $\delta^{30}\text{ASi}$  and bedrock values (Fig. 4d),  
456 indicating that the  $\delta^{30}\text{DSi}$  exported is a result of fractionation due to net secondary weathering product  
457 formation (Crompton et al., 2015). We hypothesise that the shorter residence times beneath KS  
458 compared to LG reduce the potential for redissolution of secondary weathering products to occur. It  
459 is more likely that there are fewer inland regions that remain isolated at KS, and the supraglacial  
460 waters are routed more efficiently through the subglacial system.

461 KS and LG have broadly similar bedrock compositions (Hawkings et al., 2016), with any differences  
462 unlikely to have a major impact on the overall  $\delta^{30}\text{Si}$  composition of the measured bulk bedrock  
463 (Savage et al., 2010). The intrusive rocks of the Julianhåb batholith and the Gardar Province at KS  
464 would be expected to have a limited range in  $\delta^{30}\text{Si}$  values due to minimal isotope fractionation during  
465 high temperature mantle processes (Savage et al., 2010; Savage et al., 2014). Therefore, our  
466 measurements of crushed proglacial rock debris for both catchments are in good agreement with the  
467 published range of values measured in West Greenland (Andre et al., 2006). The measured mean  
468 bedrock values of  $\delta^{30}\text{Si}$  at KS are lighter compared to LG ( $-0.18 \pm 0.06\%$  compared to  $0.00 \pm 0.07\%$ ),

469 making them more comparable to values for bulk silicates on the Earth's surface ( $-0.29 \pm 0.08\%$ ,  
470 Savage et al. 2010). The heavier  $\delta^{30}\text{Si}$  composition measured at LG could be an artefact of sampling  
471 bedrock with a weathering crust, rather than pristine samples. However, basalts have a lower isotopic  
472 composition ( $-0.3\%$  to  $-0.2\%$ , Georg et al., 2007; Chemtob et al., 2015), so it is possible KS bedrock  
473 is isotopically lighter, as a result of basaltic intrusions. Nevertheless, we attribute variations in silicon  
474 isotope composition of SPM ASi to be largely a result of weathering processes at the glacier bed. The  
475 isotopic fractionations caused by weathering conditions has been shown to be much greater than any  
476 variations in crustal samples, despite often significant differences in compositions (Ziegler et al., 2005;  
477 Savage et al., 2010; Savage et al., 2013).

478 Neither catchment showed a seasonal trend in SPM  $\delta^{30}\text{ASi}$ , which exhibited a constant offset towards  
479 lower values compared to our bulk bedrock measurements ( $\sim 0.1$  to  $0.2\%$ ; Fig. 5c, d). The offset could  
480 be the result of fractionation induced precipitation reactions, weathering of silicate rocks,  
481 comminution of particles, or a combination of all three (Andre et al., 2006; Chemtob et al., 2015). The  
482 precipitation of ASi at low temperatures results in the preferential uptake of  $^{28}\text{Si}$  into the solid phase  
483 (e.g. Geiler et al. 2014). High resolution transmission electron microscope photomicrographs of ASi  
484 in SPM show it to be associated with edges of particles and with elements such as Al and Fe,  
485 suggesting it potentially forms as a result of aluminosilicate mineral weathering (Hellmann et al.,  
486 2012; Hawkings et al., 2017). The presence of ASi with elevated Al/Si ratios indicates that it would  
487 also be enriched in  $^{28}\text{Si}$ , based on prior low temperature laboratory experiments (Oelze et al., 2015).  
488 However, the offset in SPM  $\delta^{30}\text{ASi}$  from the bedrock measurements could also be linked to physical  
489 grinding, which has been demonstrated to result in the formation of reactive amorphous surface layers  
490 (Lin and Somasundaran, 1972; Hawkings et al., 2017). ASi formed in this way is likely to be  
491 characterised by isotopically light compositions as it is derived from the alteration of a freshly crushed  
492 outer mineral layer, enriched in  $^{28}\text{Si}$ , due to kinetic fractionation (Zielger et al., 2005). All three  
493 processes result in ASi enriched in  $^{28}\text{Si}$ , so our current data is unable to infer which of these is most  
494 important as ASi from both catchments are lighter than bedrock values across the season.

#### 495 **4.4 Understanding the Isotopic Mass Imbalance**

496 The interpretations presented above highlight a potential mass imbalance that arises because the  
497 subglacial waters export both DSi and ASi that is isotopically lighter than bedrock values at LG. In  
498 addition to this, when the  $\delta^{30}\text{DSi}$  and  $\delta^{30}\text{ASi}$  compositions at LG are summed, considering the relative  
499 contributions of both, the total  $\delta^{30}\text{Si}$  is consistently lighter than the measured bedrock across the melt  
500 season. Whilst we have sampled the majority of the melt season at LG ( $>60\%$ ), we did not continue  
501 sampling until the shutdown of the subglacial system. Therefore, we have carried out a simple mass  
502 balance calculation to ensure the mass imbalance seen at LG is not simply an artefact of the  
503 unmonitored part of the season (Supplementary Table 5). Whilst we do not have geochemical data

504 past Day 210, we have a continuous discharge (Q) record until much later in the season. From this  
505 record we calculated the proportion of the measured discharge compared to the total discharge. We  
506 assumed that DSi concentrations were similar to the discharge weighted mean from the measured  
507 period. As the  $\delta^{30}\text{ASi}$  composition was relatively constant across the measured period, this trend  
508 would likely continue into the latter stages of the season. We estimate the  $\delta^{30}\text{Si}$  composition of the  
509 unmeasured DSi required to ensure the total Si exported had a  $\delta^{30}\text{Si}$  composition that matched the  
510 bulk bedrock composition. The overall  $\delta^{30}\text{Si}$  composition after the sampling period would need to be  
511 +0.44‰, with a  $\delta^{30}\text{DSi}$  composition of +2.22‰, in order to balance the  $\delta^{30}\text{Si}$  composition over the  
512 rest of the melt season. The aim of this simple calculation was to demonstrate whether the mass  
513 imbalance could be realistically resolved by only considering the latter stages of the melt season,  
514 which we were unable to sample. A  $\delta^{30}\text{DSi}$  composition of +2.22‰ is likely unrealistic, considering  
515 the range of values measured over the rest of the season are significantly lighter. It is therefore likely  
516 the subglacial processes are driving the mass imbalance seen at LG.

517

518 One hypothesis to explain this mass imbalance is that the continuing light  $\delta^{30}\text{DSi}$  values of bulk  
519 meltwaters reflect physical erosion processes. High physical erosion of bedrock under the GrIS results  
520 in the formation of fresh finely ground rock flour, with very high surface areas (Cowton et al., 2012;  
521 Telling et al., 2015; Hawkings et al., 2016; Nienow et al., 2017). Published dissolution experiments  
522 have shown there is a preferential dissolution of  $^{28}\text{Si}$  from the fresh mineral surface, as a result of  
523 kinetic fractionation (Ziegler et al., 2005). Therefore, we expect that the freshly crushed subglacial  
524 minerals to result in delivery of dissolved silicon enriched in  $^{28}\text{Si}$ . The high pH and under saturation of  
525 waters with respect to ASi subsequently promotes the outer amorphous mineral layers to undergo  
526 further dissolution, resulting in the export of light  $\delta^{30}\text{DSi}$ . The higher  $\delta^{30}\text{DSi}$  composition at the  
527 beginning of the season reflects the weathering environment at marginal areas of the ice sheet. These  
528 areas are accessed more regularly by surface melt, with efficient hydrological drainage for longer  
529 parts of the season (Chandler et al., 2013), and with potentially less active grinding of bedrock (as  
530 demonstrated by lower suspended sediment concentrations during the Early Season). As the melt  
531 season progresses, more isolated inland regions of the bed are accessed, where dissolution has  
532 occurred over longer time periods and where physical erosion is enhanced, potentially evidenced by  
533 the linear relationship between increasing SPM concentrations and decreasing  $\delta^{30}\text{DSi}$  exported from  
534 LG (Supplementary Fig. 6). Hence, the  $\delta^{30}\text{DSi}$  of meltwaters exported is lower. By comparison, SPM  
535 concentrations exported from KS are consistently lower. This could be a result of some settling in the  
536 proglacial lake, but we believe it is more likely to result from lower rates of physical erosion in this  
537 smaller catchment. We therefore hypothesise that higher physical erosion rates and longer residence  
538 times in larger catchments (Wadham et al., 2010, Hawkings et al., 2016), help to explain the  
539 differences in the  $\delta^{30}\text{Si}$  composition of meltwaters from LG and KS.

540 To quantitatively assess if this hypothesis is realistic, we have modelled the proportion of DSi that  
541 would need to result from the dissolution of ASi associated with SPM, assuming complete  
542 dissolution, when considering the  $\delta^{30}\text{Si}$  composition of the measured DSi and corresponding  
543 endmembers (Equation 7). Our model was based on a range of fractionation factors for the initial  
544 formation of ASi ( $\epsilon$ ) from the alteration of bedrock, as low temperature fractionation processes are  
545 still poorly understood (Geilert et al., 2014; Frings et al., 2016). An open system was chosen, rather  
546 than using Rayleigh fractionation, as we do not expect a finite pool of Si within the subglacial system  
547 in the timescales we are considering.

548

549 The subglacial environment is complex; therefore, so we have simplified our model by assuming  
550 overall fractionation during bedrock alteration to form ASi is similar to that observed during low  
551 temperature ASi precipitation from solution with  $\epsilon$  ranging from -2 ‰ to -5 ‰. Extrapolation from  
552 experiments by Geilert et al. (2014) produces a fractionation factor of -2.34‰ at 0°C. However their  
553 conclusions indicate that the fractionation factor at low temperatures is system dependent, due to  
554 differences in fractionation based on external factors such as saturation state and surface area. Oelze  
555 et al. (2015) found a fractionation factor of -5‰ for initial stages of experiments with high Al/Si  
556 ratios, which could simulate subglacial conditions considering the potential formation of ASi through  
557 aluminosilicate weathering (Hawkings et al., 2017). We use the first  $\delta^{30}\text{DSi}$  value measured in the  
558 Early Season at KS and a value measured at the subglacial portal in the Early Season at LG as the  
559 initial  $\delta^{30}\text{DSi}$  endmember in the model. The  $\delta^{30}\text{ASi}$  endmember value is calculated by subtracting the  
560 chosen fractionation factor from the  $\delta^{30}\text{Si}$  composition of the bulk rock for each catchment.

561

562 In our modelled scenarios for LG we show that during outburst periods (and thus when  $\delta^{30}\text{DSi}$   
563 composition is lightest), a maximum of 56% of the measured DSi results from ASi dissolution (Fig. 8).  
564 This proportion equates to  $\sim 12\mu\text{M}$  of ASi compared to an overall ASi concentration of  $\sim 290\mu\text{M}$   
565 measured at the same timepoint, suggesting that even the maximum modelled values can be  
566 considered reasonable due to the large ASi reservoir. Experimental data also suggest the most  
567 negative  $\epsilon$  values relate to solids formed rapidly and with unidirectional kinetic fractionation effects  
568 (Oelze et al., 2015), which could also relate to these subglacial weathering crusts which are highly  
569 reactive and would presumably undergo rapid dissolution. While all the modelled  $f_a$  values can be  
570 considered realistic when converted to molar concentrations, we consider the lower values as more  
571 representative of subglacial systems, considering the experimental data from Oelze et al. (2015).

572 Despite the lower ASi concentrations at KS, the modelled values can also be considered realistic for  
573 this system. As we expect the ASi dissolution to be less important at KS compared to at LG, it follows  
574 that we calculate lower proportions of ASi required to undergo dissolution in order to produce the  
575 measured  $\delta^{30}\text{DSi}$ . We would expect the DSi at KS to be a result of silicate dissolution, from



576 hydrolysis reactions, with the formation of clay minerals as a secondary weathering product  
577 (Crompton, 2015), as evidenced by increasing pH and the heavier  $\delta^{30}\text{DSi}$  when compared to LG.

578 An alternative hypothesis to explain the mass imbalance at LG requires a temporal offset between  
579 weathering product formation and re-dissolution. For example, the isotopically light secondary  
580 weathering products may be a pre-glacial feature formed in a past, warmer climatic regime,  
581 sequestered by an advancing ice sheet under a cooler climate and were stored subglacially (Hawkings  
582 et al., 2018). The presence of palaeosols (>2.7million years, Bierman et al. 2014 ) beneath the ice  
583 sheet, which are exported in runoff, supports the notion that there are pre-glacial features present  
584 (Lawson et al., 2014; Kohler et al., 2017). The high pH of LG meltwaters (up to 9.63),  
585 undersaturation of waters with respect to ASi, and high ASi concentrations (Qwt mean 209 $\mu\text{M}$  after  
586 Day 170), could result in the redissolution of these isotopically light amorphous secondary weathering  
587 products when the isolated parts of the subglacial system become hydrologically connected  
588 (Hawkings et al., 2018). Whilst this hypothesis provides explanation for the potential mass imbalance  
589 and palaeosols are evidenced beneath ice sheets, it may be expected that this isotopically light source  
590 of Si would get depleted over time and any enhanced weathering currently occurring *in-situ* would  
591 result in heavier  $\delta^{30}\text{DSi}$  exported in the meltwaters. Also, ASi measurements in the current proglacial  
592 plain display very low concentrations (average 0.01wt%, Supplementary Table 6), suggesting these  
593 amorphous phases are not currently being formed in the current proglacial environment, or have been  
594 stripped away or aged. Therefore, whilst this hypothesis has potential to provide explanation for light  
595  $\delta^{30}\text{DSi}$  compositions in glacial meltwaters, our data suggests our first hypothesis focusing on physical  
596 erosion is currently the most likely scenario.

#### 597 **4.5 Implications and Conclusions**

598 The two Greenland Ice Sheet (GrIS) catchments have different subglacial weathering regimes  
599 primarily driven by subglacial hydrology. These weathering regimes appear related to water residence  
600 time and therefore catchment size. Kiattuut Sermiat (KS) was dominated by carbonate hydrolysis,  
601 whereas a predominance of silicate mineral weathering occurred as the melt season progressed at  
602 Leverett Glacier (LG). Subglacial silicate weathering results in the production of isotopically light  
603 amorphous secondary weathering solid phases in both catchments, with heavier isotopes recorded in  
604 the dissolved fraction during Early Season meltwater discharge. As the melt season progressed, long  
605 residence time subglacial waters became connected to an efficient drainage system. The larger of the  
606 two catchments in this study, LG, exhibited a marked decrease in  $\delta^{30}\text{DSi}$  from 0.87 to -0.55‰, as  
607 meltwater discharge rose, and the subglacial drainage system expanded inland, tapping increasingly  
608 remote subglacial water pockets. We interpret this isotopic shift as evidence of increasing contribution  
609 from silicate weathering products, including the dissolution of ASi and other secondary weathering  
610 products, and leaching of freshly crushed rock surfaces. The smaller catchment, KS, discharged

611 meltwaters with a  $\delta^{30}\text{DSi}$  similar to smaller valley glaciers, with carbonate weathering the  
612 predominant solute acquisition pathway.

613 Our results show that  $\delta^{30}\text{DSi}$  and  $\delta^{30}\text{ASi}$  can be used alongside major ion data to assess the degree of  
614 silicate weathering, redissolution of weathering products and hydrological drainage characteristics in  
615 glacial meltwaters. We postulate that access to meltwater present in subglacial drainage systems with  
616 limited connection to an efficient drainage system will increase in the future, as snowlines retreat  
617 further inland and melt increases under climatic warming (Hawkings et al., 2015). It is therefore likely  
618 that the intensity of subglacial geochemical weathering and the composition of glacial waters entering  
619 the ocean will be impacted. These findings are critical when attempting to estimate the fluxes of  
620 nutrients from rapidly melting glaciated regions and their impact on elemental cycles, in the past (e.g.  
621 during deglaciation events, Hawkings et al., 2018), present and future. The residence time of  
622 subglacial waters will have an influence on the weathering reactions occurring, and therefore the  
623 fluxes of key nutrients (Fe, P and Si) exported from glacial systems (Stevenson et al., 2017). More  
624 robust estimation of Si fluxes and the  $\delta^{30}\text{Si}$  composition of exported waters from the GrIS requires full  
625 consideration and further study of catchment size, hydrological development, weathering regime, and  
626 other processes within the complex subglacial system. The degree of silicate weathering in glaciated  
627 regions may be much higher than previously thought, considering that it is the larger catchments that  
628 discharge the majority of meltwater into the ocean from the GrIS and AIS. The quantity of meltwater  
629 delivered Si from ice sheets and its isotopic composition should therefore be considered when  
630 calculating global silicon budgets.

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816  
817 **Table 1: Hydrological comparison of two studied catchments.** Qwt = discharge weighted mean,  
818 D:M = divalent to monovalent ion ratio, DSi = Dissolved silicate, ASi = Amorphous silica.  
819

	Kiattuut Sermia (2013)	Leverett Glacier (2015)
Mean Q ( $\text{m}^3\text{S}^{-1}$ )	22.80 <sup>+</sup>	85.35
Total Q ( $\text{km}^3$ )	0.22 <sup>+</sup>	1.45
Qwt Sus. Sediment ( $\text{g L}^{-1}$ )	0.12 <sup>+</sup>	0.87
Qwt pH	9.09	8.63
Qwt EC ( $\mu\text{S cm}^{-1}$ )	26.66 <sup>+</sup>	12.25
Qwt D:M ( $\mu\text{eq}$ )	6.91	1.36
Qwt DSi ( $\mu\text{M}$ )	22.18	20.76
Qwt ASi ( $\mu\text{M}$ )	9.87	205.78
Qwt $\delta^{30}\text{DSi}$ (‰)	0.41	-0.25
Qwt $\delta^{30}\text{ASi}$ (‰)	-0.47	-0.22
Bulk Rock $\delta^{30}\text{Si}$ (‰)	-0.18	0.00 <sup>+</sup>

820 <sup>+</sup>Data previously published by Hawkings et al. (2016).

821

822 **Figure 1: Location of Kiattuut Sermiat and Leverett Glacier, from Hawkings et al. (2016).** Water  
823 samples were collected at the locations marked in black from proglacial streams, as in Lawson et al.  
824 (2014) and Hawkings et al. (2016). Hydrological monitoring was completed at points of stable  
825 bedrock (white markers).

826 **Figure 2: Hydrological and Geochemical Time series for LG (black, A) and KS (red, B).** Vertical  
827 black shading in A shows outburst events recorded during the melt season (Kohler et al., 2017) and  
828 vertical red shading in B shows the “Spring Event” at KS (Hawkings et al., 2016).

829 **Figure 3: Major ion ratio time series.** Ratio of Divalent/Monovalent ions (D:M) Differences in the  
830 seasons (i.e. Late and Early Season) are defined by differences in hydrological and geochemical data  
831 in the proglacial river.

832 **Figure 4: Na-normalised molar ratio mixing diagrams.** Silicate and carbonate endmembers taken  
833 from Gaillardet et al. (1999) and references within, using data from small rivers draining one single  
834 lithology.

835 **Figure 5: Silicon isotope composition results.** A) The relationship between  $\delta^{30}\text{DSi}$  and D:M ion  
836 ratio. B)  $\delta^{30}\text{DSi}$  and discharge for both catchments. C and D) Time series of  $\delta^{30}\text{DSi}$  and  $\delta^{30}\text{ASi}$  for KS  
837 and LG. Vertical blue shading in C indicates opening of the subglacial drainage system (“Spring  
838 Event”, Hawkings et al. (2016)) and outburst events in D. Bulk bedrock  $\delta^{30}\text{Si}$  is indicated by the  
839 horizontal dashed line, with the shading representing 2SD. All error bars represent 2SD of external  
840 errors (0.08 for  $\delta^{30}\text{DSi}$  and 0.14 for  $\delta^{30}\text{ASi}$ ).

841 **Figure 6: Diagram to illustrate conceptual model of subglacial hydrology development.**  
842 Simplified development of subglacial hydrology at LG as the melt season progresses. Panel A  
843 represents processes during the Early season of LG (before the first outburst event), B represents the  
844 system after the first outburst event and subglacial water influences the proglacial stream chemistry,  
845 and C represents the system during the late season once the snowline has retreated further and  
846 outburst events occur more often. The smaller size of KS limits the subglacial hydrology development  
847 to A and B, without the connection of isolated subglacial water by supraglacial lake drainage.

848 **Figure 7: Major ion relationship plots.** A)  $\text{Ca}^{2+} + \text{Mg}^{2+}$  versus  $\text{SO}_4^{2-}$  B)  $\text{Ca}^{2+} + \text{Mg}^{2+}$  versus  $\text{HCO}_3^-$  C)  
849  $\text{HCO}_3^-$  versus  $\text{SO}_4^{2-}$ . Early Season at KS is defined as before the subglacial system connected (Day  
850 157, Hawkings et al. (2016)), Transition Period is the hydrologically unique period after the  
851 subglacial connection and Late Season is post Day 169. Early season at LG is defined at prior to first  
852 outburst event/subglacial connectivity at day 170 and late season is defined as day 170 onwards. Grey  
853 dashed lines show main relationships found by Tranter et al. (2002) when studying an Alpine valley  
854 glacier. The table outlines the regression slopes, intercepts and  $R^2$  values for each relationship.

855 **Figure 8: Modelled Percentage ASi required for measured  $\delta^{30}\text{DSi}$  composition.** Panels A and B  
856 show the calculated  $f_a$  values according to Equation 7, with A (black) showing the time series at LG  
857 and B (red) showing the time series at KS.  $\epsilon$  values represent the varying fractionation factors used in  
858 the model. Panels C (LG) and D (KS) show the concentration of ASi required to produce the  
859 measured DSi, compared to the measured ASi concentrations over the melt season.

860 **Supplementary Figure 1: Estimated lake Residence Time at KS.** Residence time of the proglacial  
861 lake at KS estimated by calculating the turnover rate of the lake using the estimated water volume of  
862 the lake and river discharge.

863 **Supplementary Figure 2: Silicon three Isotope plot.** All samples analysed during the study are  
864 included with red dashed lines showing the 95% prediction band and grey lines showing the 95%  
865 confidence band. The black regression line has a gradient of 0.5118 ( $p < 0.0001$ ,  $R^2 = 0.9946$ ), showing  
866 mass dependent fractionation

867 **Supplementary Figure 3: Relationship between  $\delta^{30}\text{DSi}$  composition and saturation index of  
868 amorphous silica.** Saturation indices calculated using Geochemists Workbench Software and plotted  
869 against  $\delta^{30}\text{DSi}$  for both catchments. Data has been split into the defined hydrological periods of the  
870 melt season to show the temporal trend.

871 **Supplementary Figure 4: Comparison of geochemical data with previously published data at  
872 LG.** Records of discharge, D:M ratio and  $\text{SO}_4^{2-}$  from LG in 2009 (Hindshaw et al., 2014) and 2015  
873 (this study).

874 **Supplementary Figure 5: Sulphate Mass Fraction Timeseries.** Calculated SMF for KS and LG  
875 over the melt season, with the seasons defined by hydrogeochemical data.

876 **Supplementary Figure 6: Relationship between SPM concentrations and dissolved silicon**  
877 **isotope composition.**



# Supplementary Tables

- 1 **Supplementary Table 1: Selected ionic concentrations from Leverett Glacier 2015, corresponding to timepoints in which silicon isotope measurements**
- 2 **were made.**

<b>Decimal Day</b>	<b>Na<sup>+</sup> (μeq)</b>	<b>K<sup>+</sup> (μeq)</b>	<b>Mg<sup>2+</sup> (μeq)</b>	<b>Ca<sup>2+</sup> (μeq)</b>	<b>Al<sup>3+</sup> (μeq)</b>	<b>SO<sub>4</sub><sup>2-</sup> (μeq)</b>	<b>NO<sub>3</sub><sup>-</sup> (μeq)</b>	<b>HCO<sub>3</sub><sup>-</sup> (μeq)</b>	<b>F<sup>-</sup> (μeq)</b>	<b>Cl<sup>-</sup> (μeq)</b>
128.4	53.84	36.79	98.64	349.25	-	229.58	0.35	271.80	2.11	30.98
135.4	41.04	26.68	61.27	215.01	1.66	118.64	0.39	200.01	0.93	20.05
140.8	56.16	38.59	41.59	126.20	4.06	75.20	0.44	256.49	1.22	37.1
145.4	64.46	39.38	73.43	252.03	-	117.65	0.35	286.19	1.53	19.94
148.4	46.21	30.20	52.09	183.94	5.59	88.86	0.33	203.98	1.08	1487
153.4	88.11	48.69	52.26	262.81	2.03	161.33	0.15	260.72	3.14	24.90
160.4	67.06	38.06	42.93	220.05	12.62	122.06	0.31	222.43	1.90	18.24
165.4	49.13	25.40	24.07	123.15	2.50	59.27	0.08	145.82	1.18	14.58
171.4	55.46	28.05	20.89	102.54	1.56	47.75	0.18	143.14	1.07	12.95
174.7	44.83	16.96	13.71	58.35	1.98	24.71	0.16	87.43	1.31	18.68
178.4	65.73	29.85	19.74	87.76	20.78	45.21	0.08	148.22	1.43	7.25
183.4	66.15	37.57	20.31	83.19	22.94	37.66	0.16	159.93	1.54	6.24
186.4	50.78	24.61	14.56	65.33	-	26.79	0.19	119.10	0.92	6.34
189.4	47.50	20.29	13.47	60.33	139.81	23.70	0.15	111.80	0.94	3.48
191.4	48.82	22.10	15.99	75.44	15.09	31.13	0.18	124.23	0.98	4.05
196.3	36.56	16.10	13.40	65.53	16.01	25.13	0.13	99.53	0.85	4.59
200.4	44.98	20.55	12.40	73.24	61.02	38.46	0.09	105.85	0.98	4.83
207.4	52.17	23.93	13.88	89.36	58.96	45.75	0.10	128.46	1.46	2.53
208.4	59.20	26.68	14.14	106.63	69.25	54.22	0.10	146.83	1.35	3.09
210.3	53.30	24.34	16.37	95.11	78.68	47.20	0.12	136.60	1.38	2.64

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7 **Supplementary Table 2: Selected ionic concentrations from Kiattuut Sermiat 2013, corresponding to timepoints in which silicon isotope measurements**  
 8 **were made.**

Decimal Day	Na <sup>+</sup> (μeq)	K <sup>+</sup> (μeq)	Mg <sup>2+</sup> (μeq)	Ca <sup>2+</sup> (μeq)	Al <sup>3+</sup> (μeq)	SO <sub>4</sub> <sup>2-</sup> (μeq)	NO <sub>3</sub> <sup>-</sup> (μeq)	HCO <sub>3</sub> <sup>-</sup> (μeq)	F <sup>-</sup> (μeq)	Cl <sup>-</sup> (μeq)
111.59	47.92	29.14	63.53	442.50	5.10	66.13	0.07	477.37	19.01	24.51
124.48	45.32	28.32	53.57	404.70	-	70.94	0.23	417.08	18.15	22.96
139.50	45.22	29.46	54.62	378.04	-	75.97	0.23	387.05	17.65	23.91
149.50	46.24	29.47	56.42	397.38	-	82.29	0.32	398.28	18.39	26.13
150.50	42.33	27.46	51.81	373.51	6.64	82.68	0.32	363.79	19.03	25.91
155.52	42.37	27.34	49.55	358.36	7.24	73.93	0.30	359.86	16.12	24.21
159.50	36.44	24.16	42.55	296.01	-	59.01	0.26	303.08	12.14	21.92
162.63	30.68	21.43	36.56	264.80	-	50.45	0.26	268.78	11.86	19.41
170.90	24.47	18.84	28.39	224.87	17.44	29.96	0.19	251.22	9.11	12.73
172.68	22.96	18.03	27.04	224.71	19.92	26.37	0.16	244.61	8.57	11.29
178.50	21.14	17.23	26.89	227.25	-	22.92	0.15	250.96	7.44	9.39
181.50	19.95	16.37	25.15	226.12	-	22.27	0.12	246.50	8.09	9.24
194.52	18.88	14.46	25.91	218.87	-	20.43	0.09	240.61	8.43	7.47
205.54	15.01	11.96	21.04	179.89	-	14.43	0.08	200.92	6.53	4.99
207.50	13.18	10.44	17.51	147.3	-	13.76	0.09	161.89	6.71	4.99
212.53	14.65	11.22	19.34	181.91	13.53	15.05	0.06	198.67	7.41	5.26
220.63	13.56	10.52	17.92	167.17	11.73	15.04	0.05	180.46	7.82	5.17
221.6	13.14	11.01	16.93	164.53	-	14.15	0.03	191.17	6.68	4.57

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12 **Supplementary Table 3: Summary of Kiattuut Sermiat silicon isotope results from 2013.**

Decimal Day	Discharge (m <sup>3</sup> s <sup>-3</sup> )	pH	Suspended Sediment (g L <sup>-1</sup> )	DSi (μM)	δ <sup>30</sup> DSi (‰)	δ <sup>30</sup> DSi error * (2σSD ‰)	ASi (wt%)	δ <sup>30</sup> ASi (‰)	δ <sup>30</sup> ASi error * (2σSD ‰)	Monovalent/ Divalent ions (μeq L <sup>-1</sup> )
111.6	0.7	8.29	0.04	40.59	1.01	0.05				6.57
124.5	2.6	7.79	0.04	39.29	0.79	0.18				6.22
139.5	3.6	8.08	0.11	36.60	0.71	0.08				5.79
149.5	7.0	7.94	0.25	38.45			0.24	-0.56	0.05	5.99
150.5	7.1	8.02	0.15	36.14	0.69	0.11				6.09
155.5	17.9	8.23	0.16	34.62	0.64	0.09				5.85
159.5	34.5	8.86	0.18	28.33			0.37	-0.54	0.05	5.59
162.6	27.6	8.85	0.13	26.76	0.47	0.10				5.78
170.9	34.4	9.19	0.13	22.70			0.05	-0.38	0.07	6.05
172.7	33.3	9.25	0.11	21.84	0.26	0.11				6.14
178.5	18.5	9.38	0.13	20.69			0.36	-0.42	0.06	6.62
181.5	19.5	9.38	0.12	20.90	0.25	0.11				6.92
194.5	43.3	9.46	0.10	20.68	0.38	0.08	0.16	-0.48	0.06	7.34
205.5	42.9	9.08	0.09	16.49	0.57	0.02				7.45
207.5	37.7	9.08	0.10	16.00			0.26	-0.42	0.06	6.98
212.5	36.2	9.13	0.10	16.49	0.41	0.09	0.24			7.78
220.6	29.9	9.48	0.08	16.79	0.16	0.14				7.67
221.6	29.9	9.34	0.09	15.92			0.18	-0.56	0.05	7.67

13 \*Propagated internal error (2σ SD, ‰)

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18 **Supplementary Table 4: Summary of Leverett Glacier silicon isotope results from 2015.**

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Decimal Day	Discharge (m <sup>3</sup> s <sup>-3</sup> )	pH	Suspended Sediment (g L <sup>-1</sup> )	DSi (μM)	δ <sup>30</sup> DSi (‰)	δ <sup>30</sup> DSi error * (2σSD ‰)	ASi (wt%)	δ <sup>30</sup> ASi (‰)	δ <sup>30</sup> ASi error * (2σSD ‰)	Monovalent/ Divalent ions (μeq L <sup>-1</sup> )
128.4	-	8.98	-	53.66	0.85	0.04	-	-	-	4.94
135.4	-	7.88	-	36.01	0.87	0.04	0.85	0.05	0.05	4.08
140.8	-	8.35	-	38.86	-		1.41	0.21	0.03	3.42
145.4	-	7.90	-	38.47	0.65	0.03	1.40	-0.16	0.03	3.13
148.4	5.4	7.72		35.63	-	-	-	-	-	3.09
153.4	13.0	7.85	0.54	37.00	0.66	0.04	1.03	-	-	2.30
160.4	17.5	7.80	0.53	31.51	0.60	0.02	0.75	-	-	2.34
165.4	25.1	7.64	0.56	24.13	0.37	0.03	0.86	-0.19	0.03	1.98
171.4	73.2	8.01	1.18	20.24	0.02	0.08	0.64	-	-	1.48
174.7	90.0	7.63	0.72	15.05	-0.23	0.02	-	-	-	1.17
178.4	122.7	8.41	1.37	24.85	-0.18	0.03	0.69	-0.20	0.04	1.12
183.4	215.1	9.18	2.09	27.69	-0.52	0.02	0.39	-0.15	0.03	1.00
186.4	207.2	8.33	0.85	19.75	-0.41	0.03	0.70	-	-	1.14
189.4	294.5	8.76	0.97	17.79	-0.41	0.03	0.83	-	-	1.09
191.4	360.9	9.24	1.05	21.41	-0.22	0.03	0.59	-0.27	0.04	1.29
196.3	230.1	8.58	0.66	16.29	-0.55	0.04	0.64	-	-	1.50
200.4	311.3	8.86	0.66	19.70	-0.14	0.03	0.85	-0.32	0.03	1.31
207.4	301.9	9.53	0.70	22.02	-	-	0.54	-0.16	0.04	1.36
208.4	270.5	9.62	0.78	24.83	-0.09	0.03	0.31	-	-	1.41
210.3	236.3	9.51	0.83	23.63	-0.06	0.03	0.97	-0.11	0.03	1.44

20 \*Propagated internal error (2σ SD, ‰)

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23 **Supplementary Table 5: Summary of Measured and Calculated values for Mass Balance Calculations at LG.**

	Up to Day 210	After Day 210*	Total*	24
Q (L)	$1.27 \times 10^{12}$	$6.04 \times 10^{11}$	$1.88 \times 10^{12}$	25
% of Q for melt period	67.8	32.2	100	26
DSi Flux ( $\mu\text{Mol}$ )	$2.65 \times 10^{13}$	$1.26 \times 10^{13}$	$3.90 \times 10^{13}$	27
ASi Flux ( $\mu\text{Mol}$ )	$2.65 \times 10^{14}$	$1.25 \times 10^{14}$	$3.90 \times 10^{14}$	28
Total Si Flux ( $\mu\text{Mol}$ )	$2.91 \times 10^{14}$	$1.38 \times 10^{14}$	$4.29 \times 10^{14}$	29
% of DSi Flux for melt period	9.1	9.1	9.1	30
% of ASi Flux for melt period	90.9	90.9	90.9	31
				32
Qwt $\delta^{30}\text{DSi}$ (‰)	-0.25	+2.20	NA	33
Qwt $\delta^{30}\text{ASi}$ (‰)	-0.22	-0.22	NA	34
Total $\delta^{30}\text{Si}$ (‰)	-0.21	+0.44	0.00	

35 \* All calculated values, except for Q, based on the percentage of Q measured before and after Day 210.

36

37 **Supplementary Table 6: Details of ASi concentrations taken from selected proglacial debris in front of LG.**

Sample Number	Date of Collection	Location	ASi (%)
1	30/5/2015	N67 03.909 W050 09.795	0.02
2	30/5/2015	N67 03.912 W050 09.789	>0.01
6	2/6/2105	N67 03.758 W050 12.093	0.01

38

Figure 1

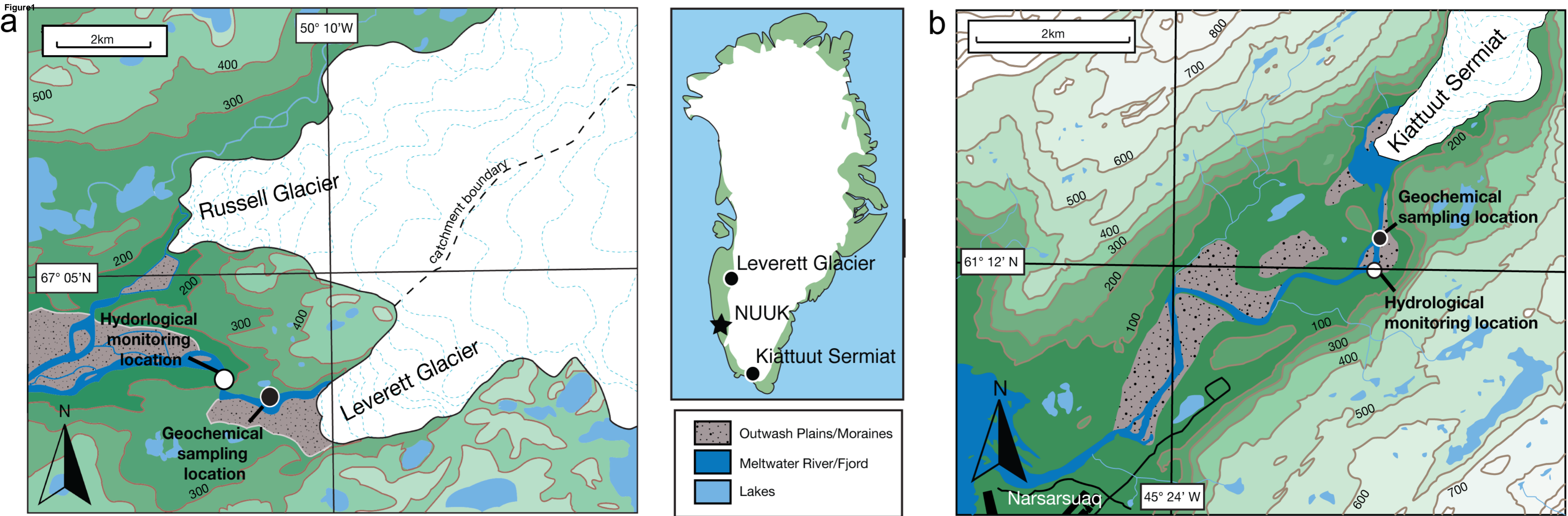


Figure2

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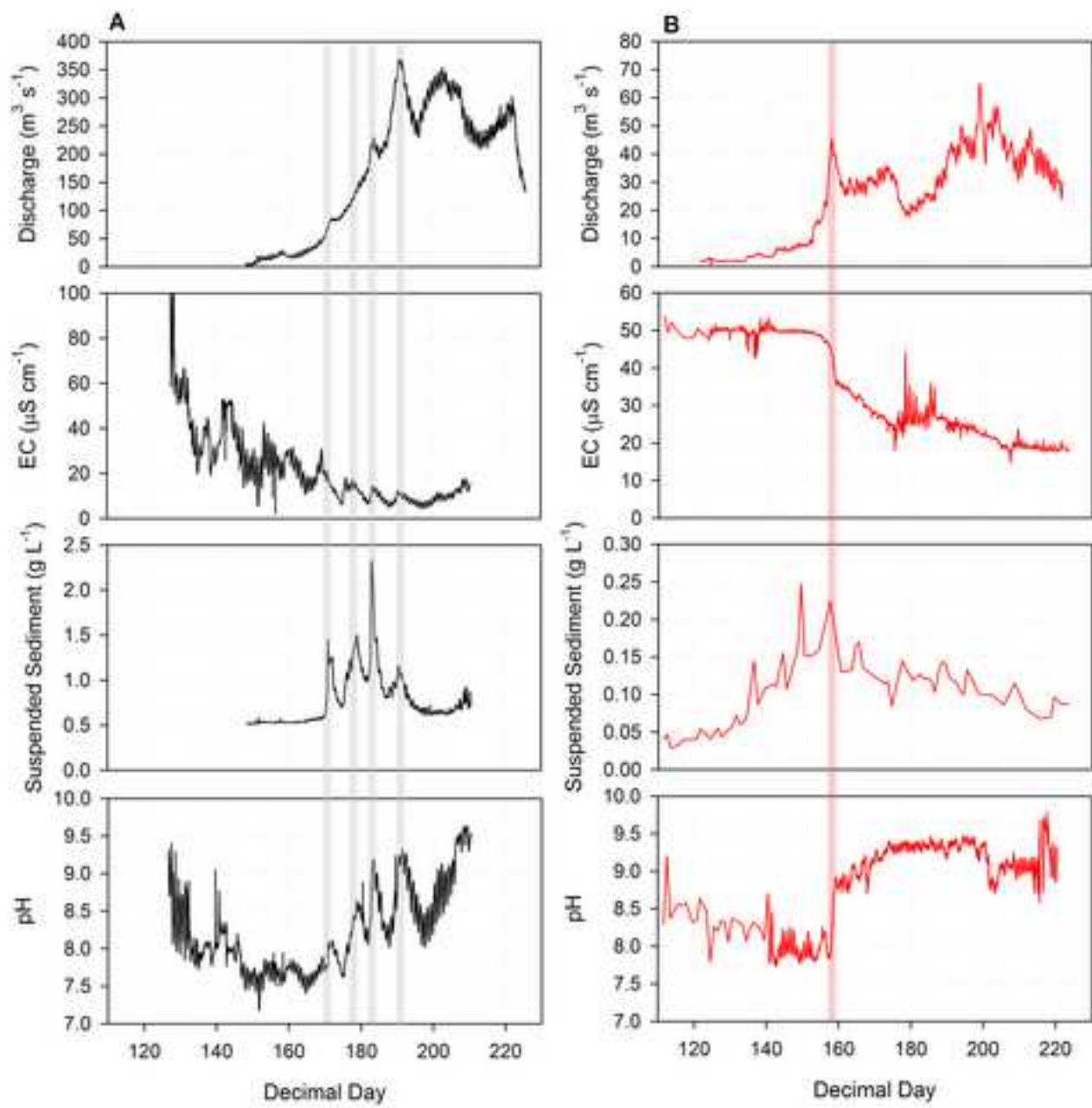


Figure3

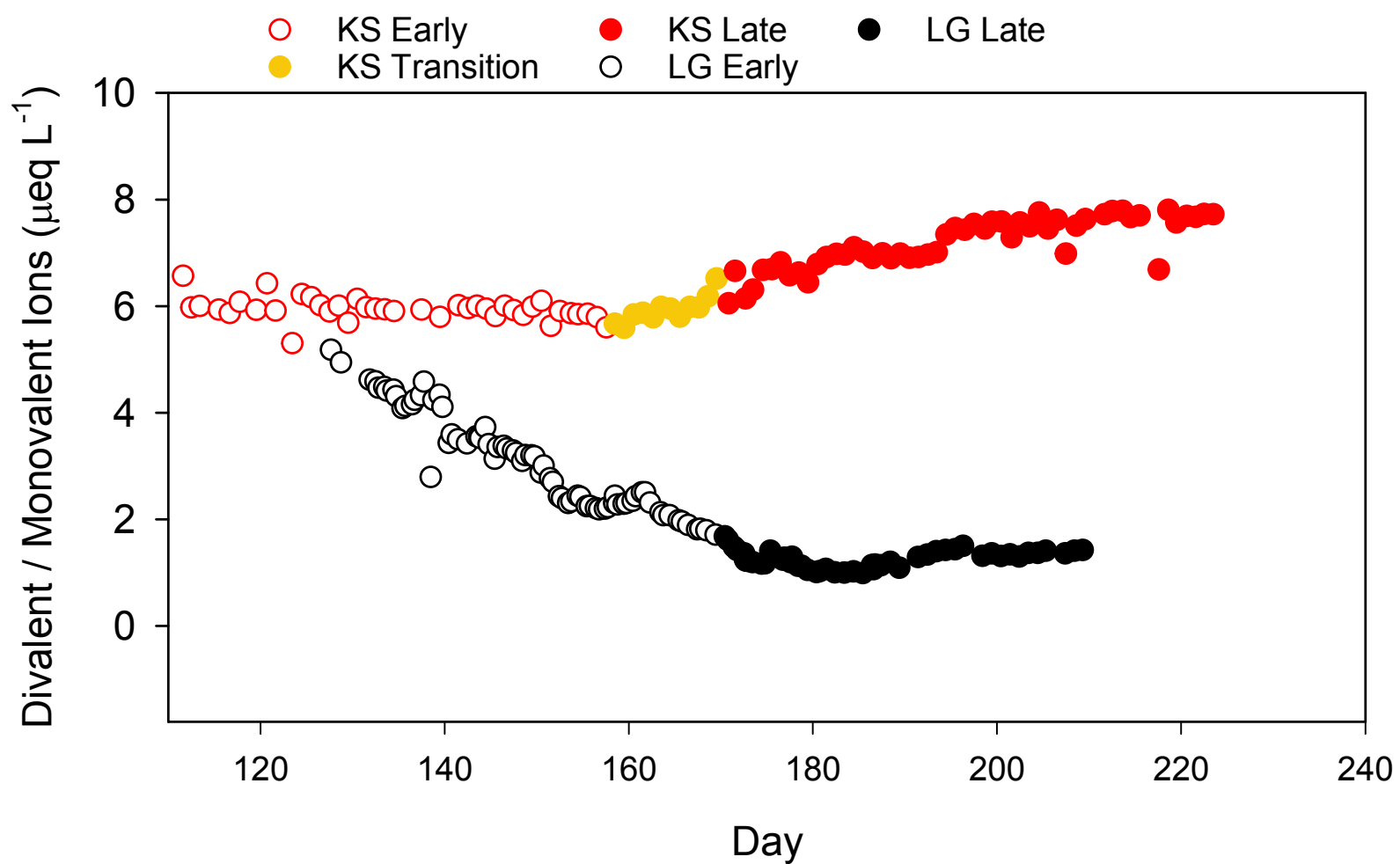




Figure 4

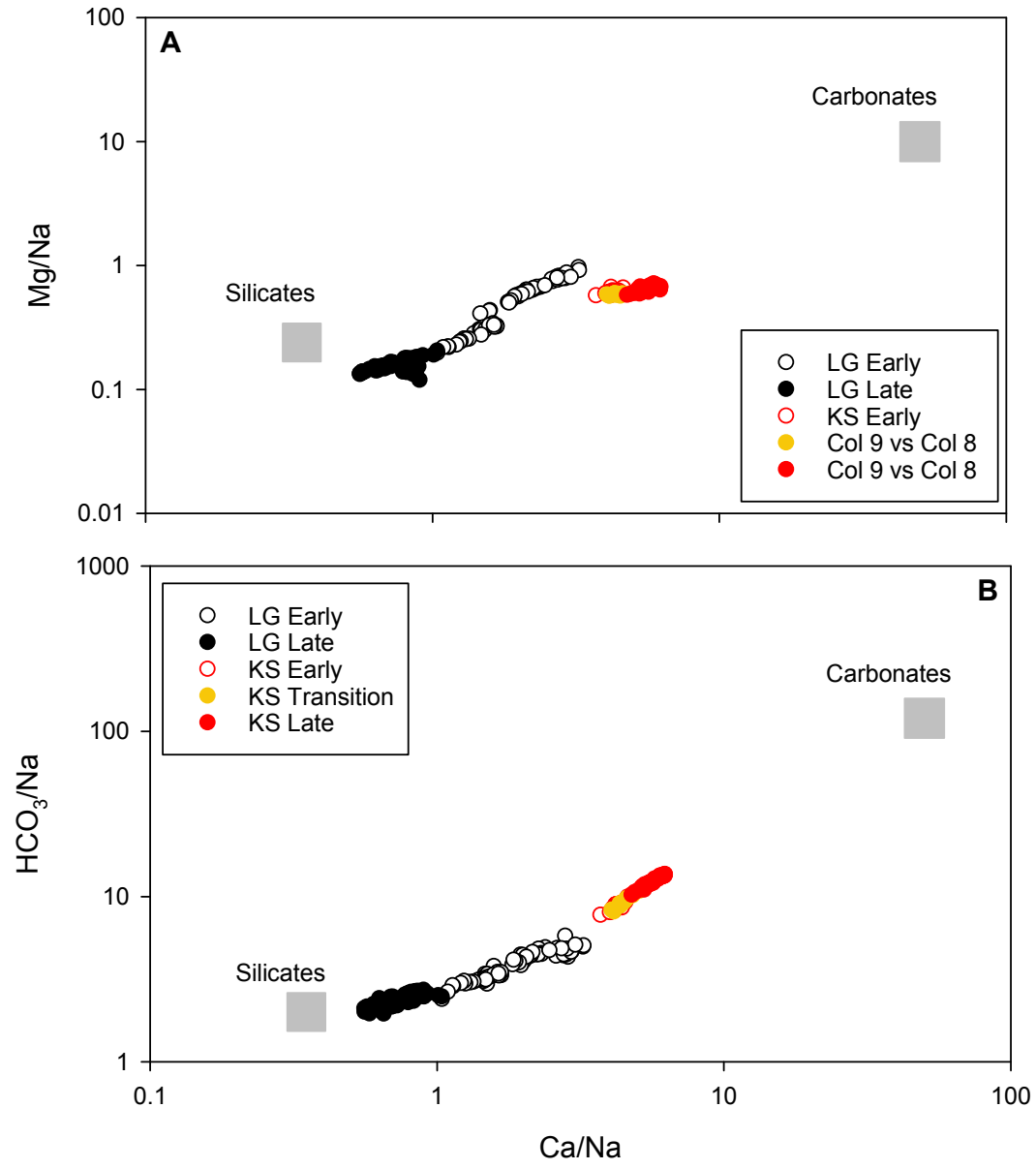


Figure5  
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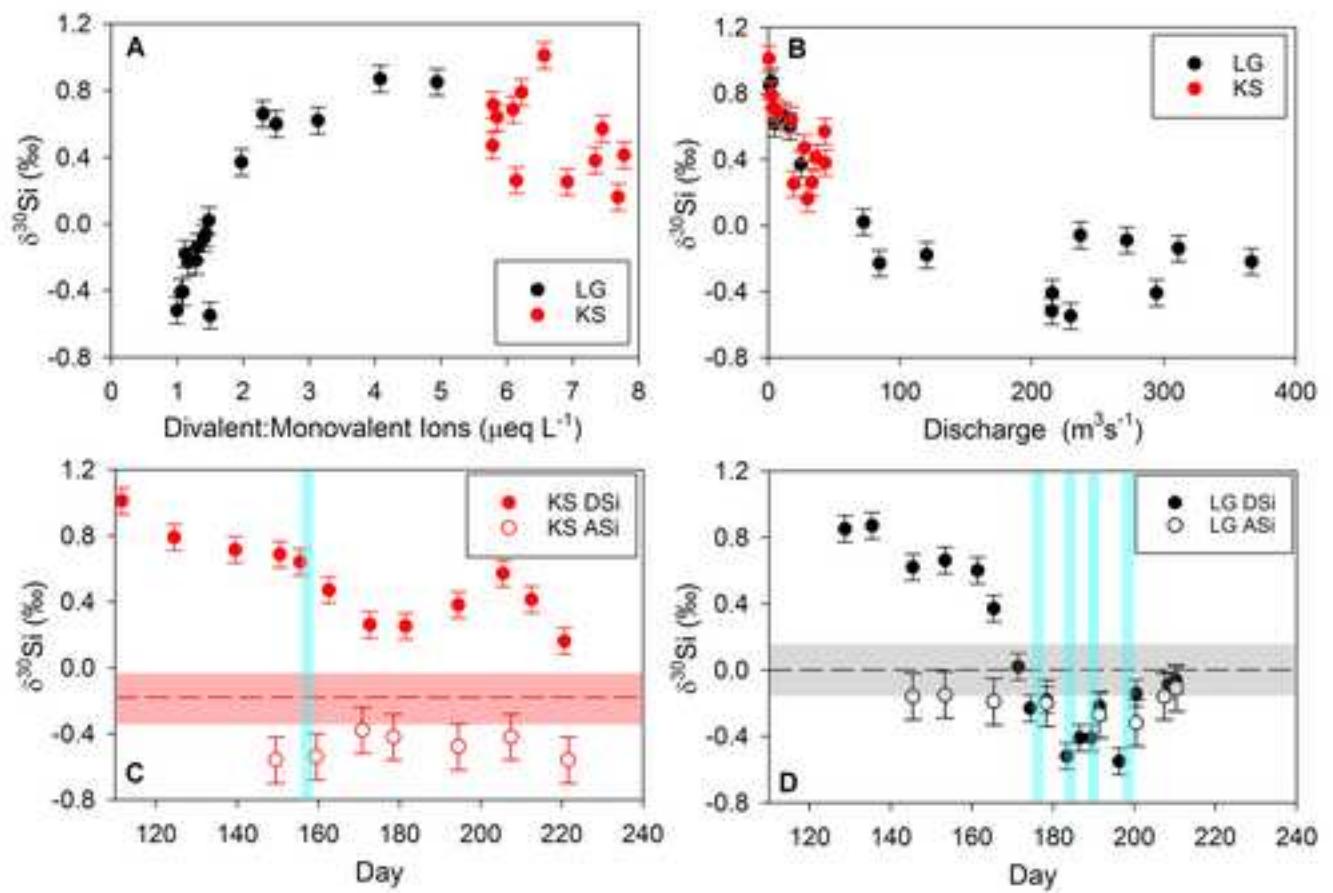


Figure 6

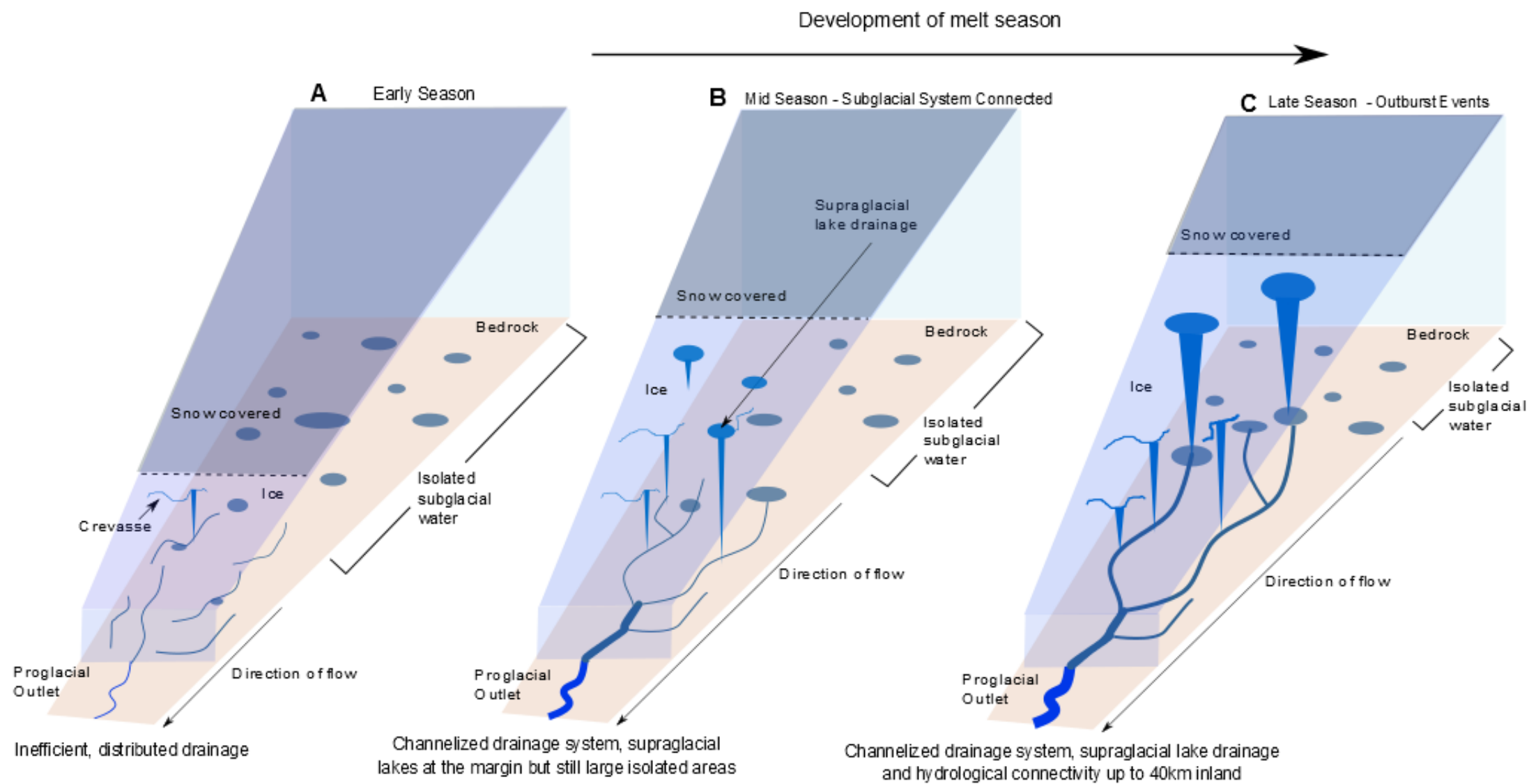
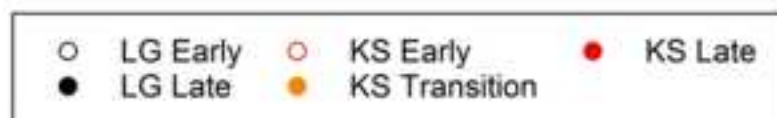
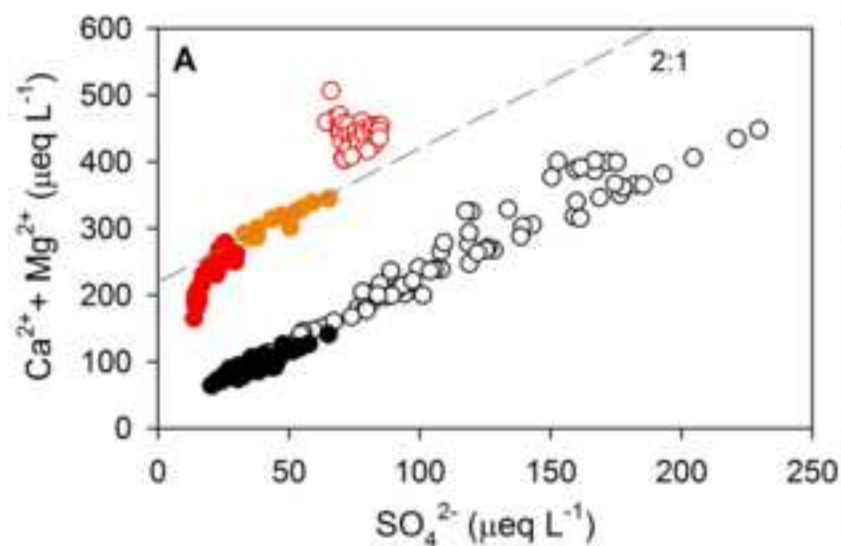


Figure 7

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		$\text{HCO}_3^- / \text{SO}_4^{2-}$	$(\text{Ca}^{2+} + \text{Mg}^{2+}) / \text{SO}_4^{2-}$	$(\text{Ca}^{2+} + \text{Mg}^{2+}) / \text{HCO}_3^-$
KS Early	Gradient	-1.58	-0.32	0.81
	Intercept	514	465	120
	R <sup>2</sup> Value	0.1359	0.0076	0.9109
KS Transition	Gradient	0.82	1.82	1.50
	R <sup>2</sup> Value	0.4735	0.8585	0.8224
KS Late	Gradient	5.53	0.04	1.05
	Intercept	114	109	-5.69
	R <sup>2</sup> Value	0.7634	0.8259	0.9927
LG Early	Gradient	0.95	1.92	1.52
	Intercept	95.4	38.1	-52.3
	R <sup>2</sup> Value	0.6461	0.9354	0.8231
LG Late	Gradient	1.17	1.52	0.83
	Intercept	80.2	38.8	-7.57
	R <sup>2</sup> Value	0.4941	0.8586	0.768

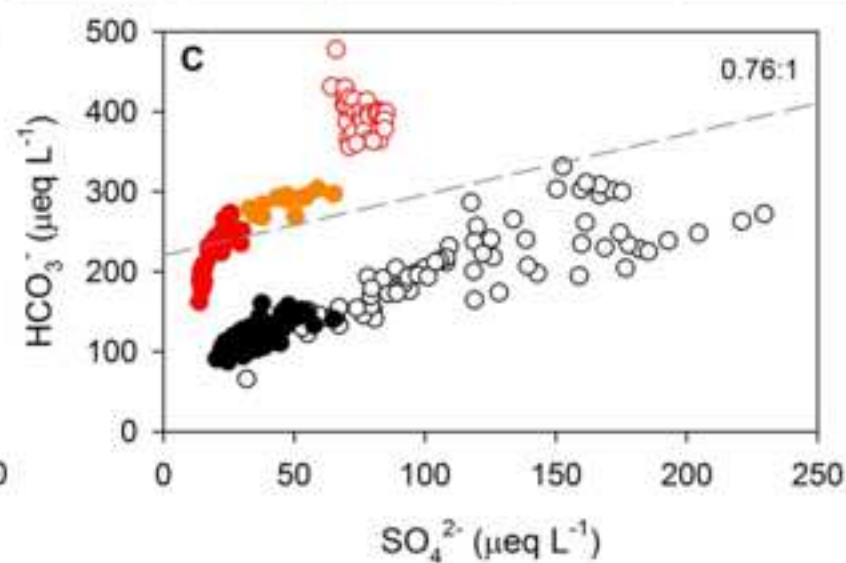
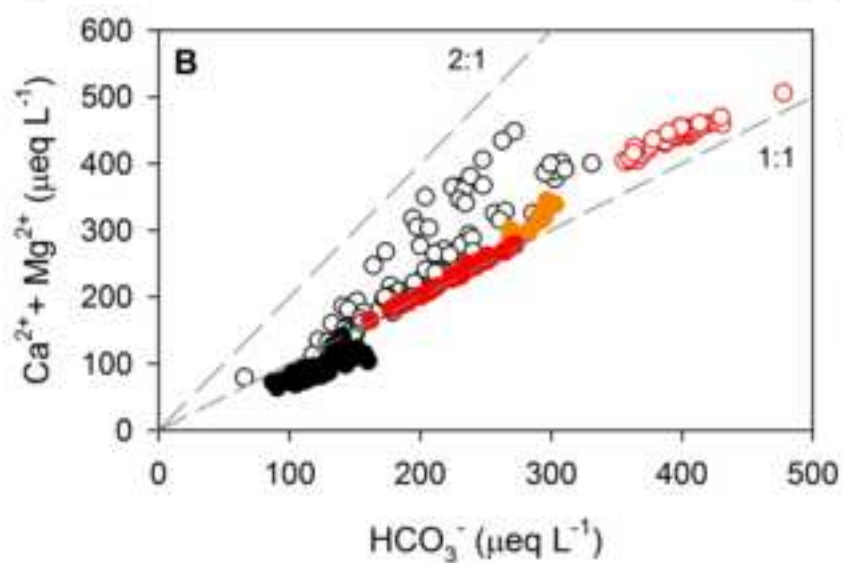
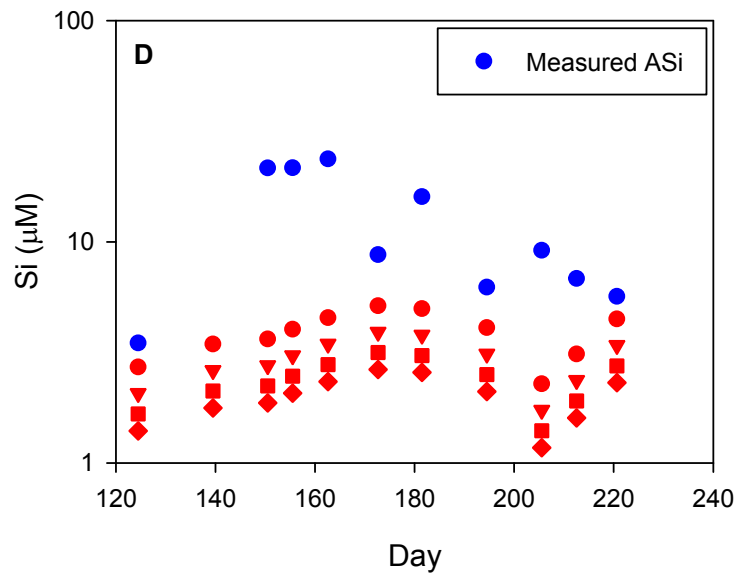
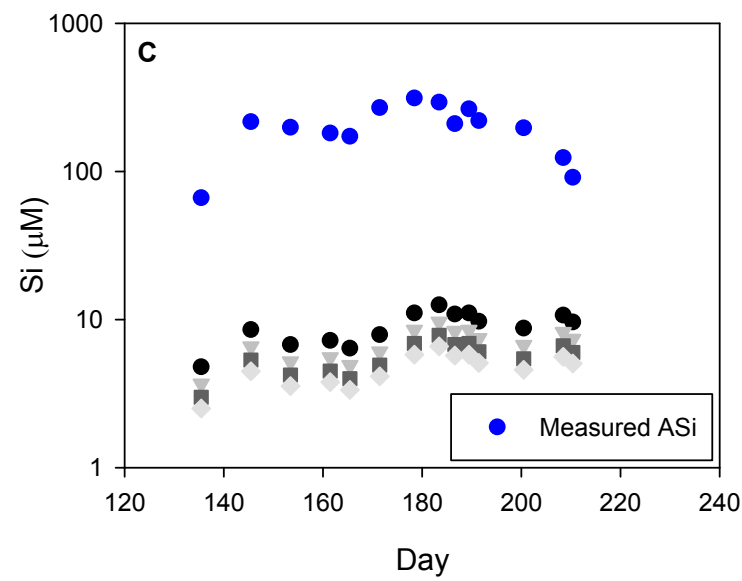
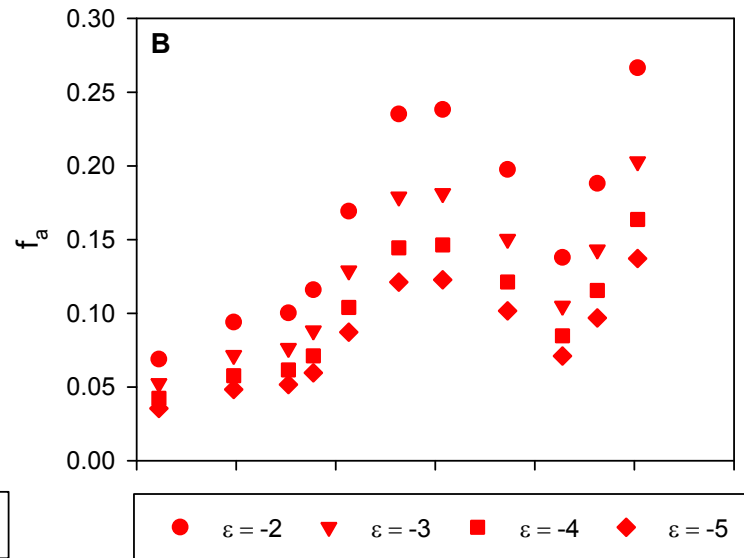
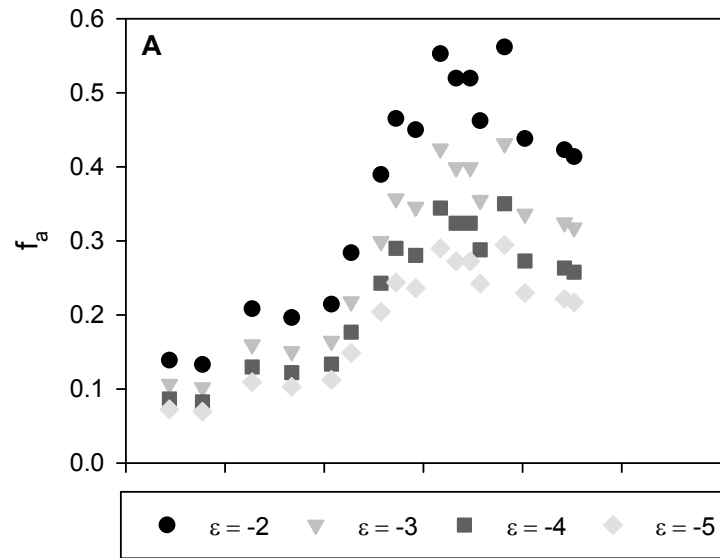
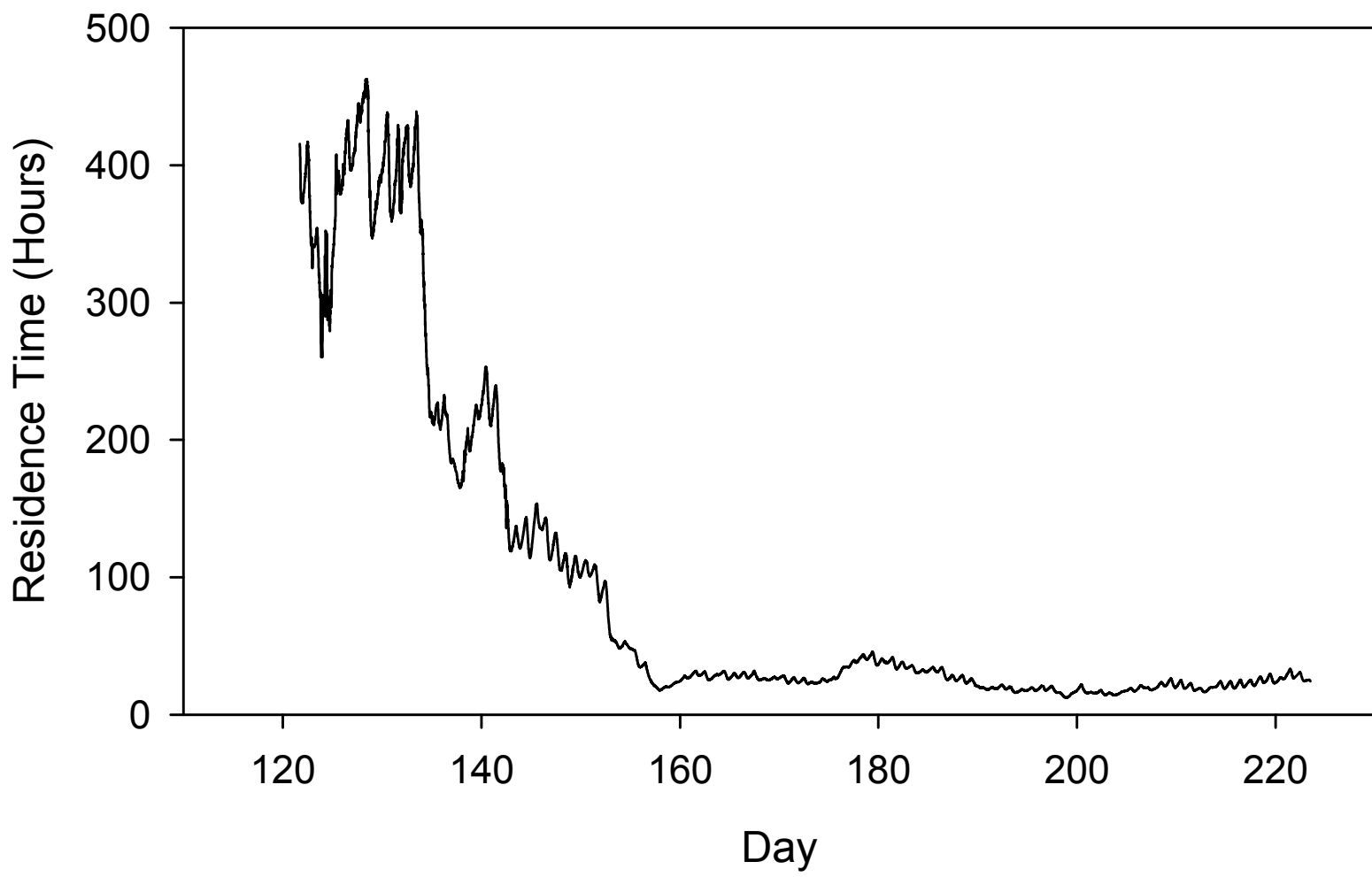
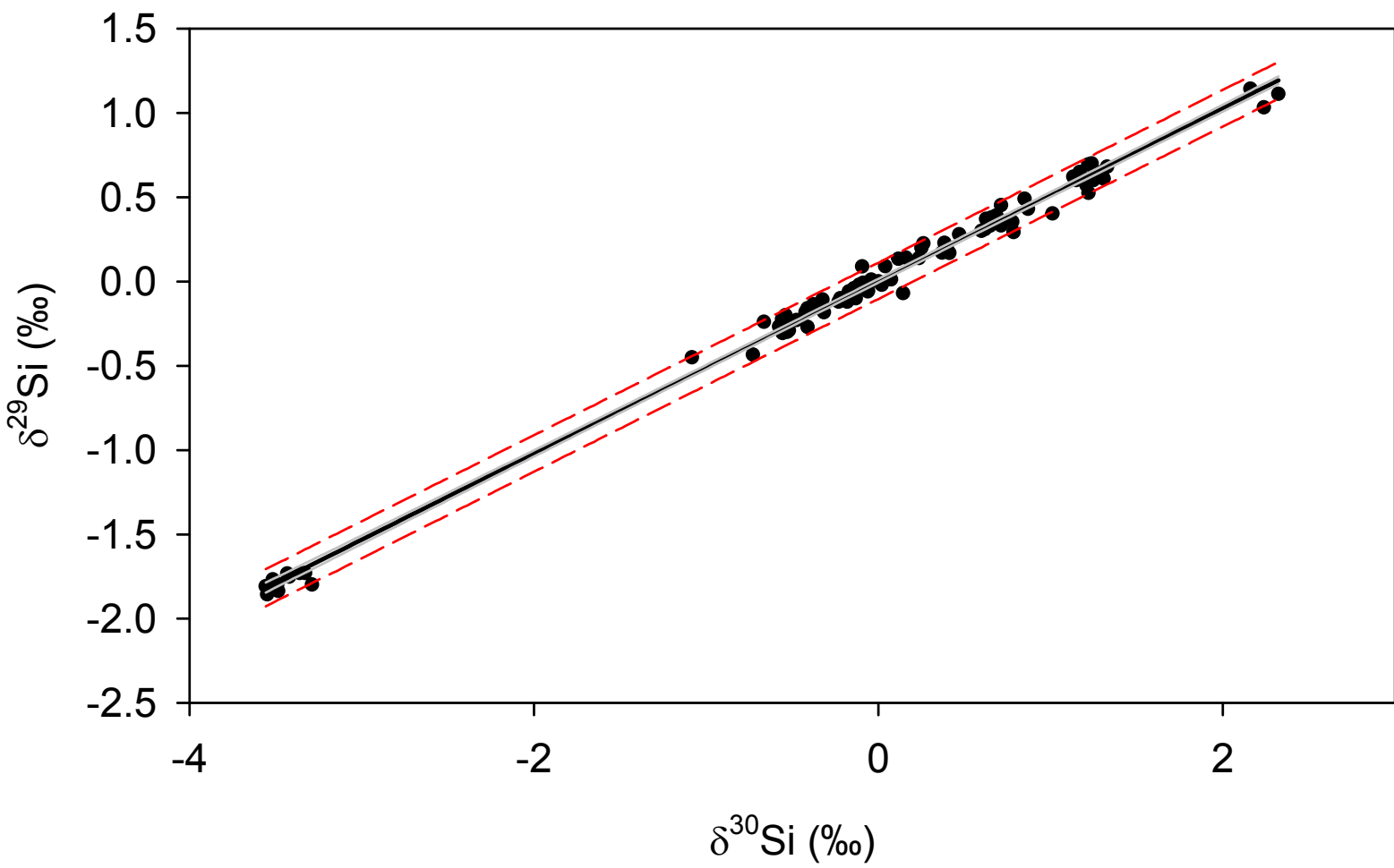
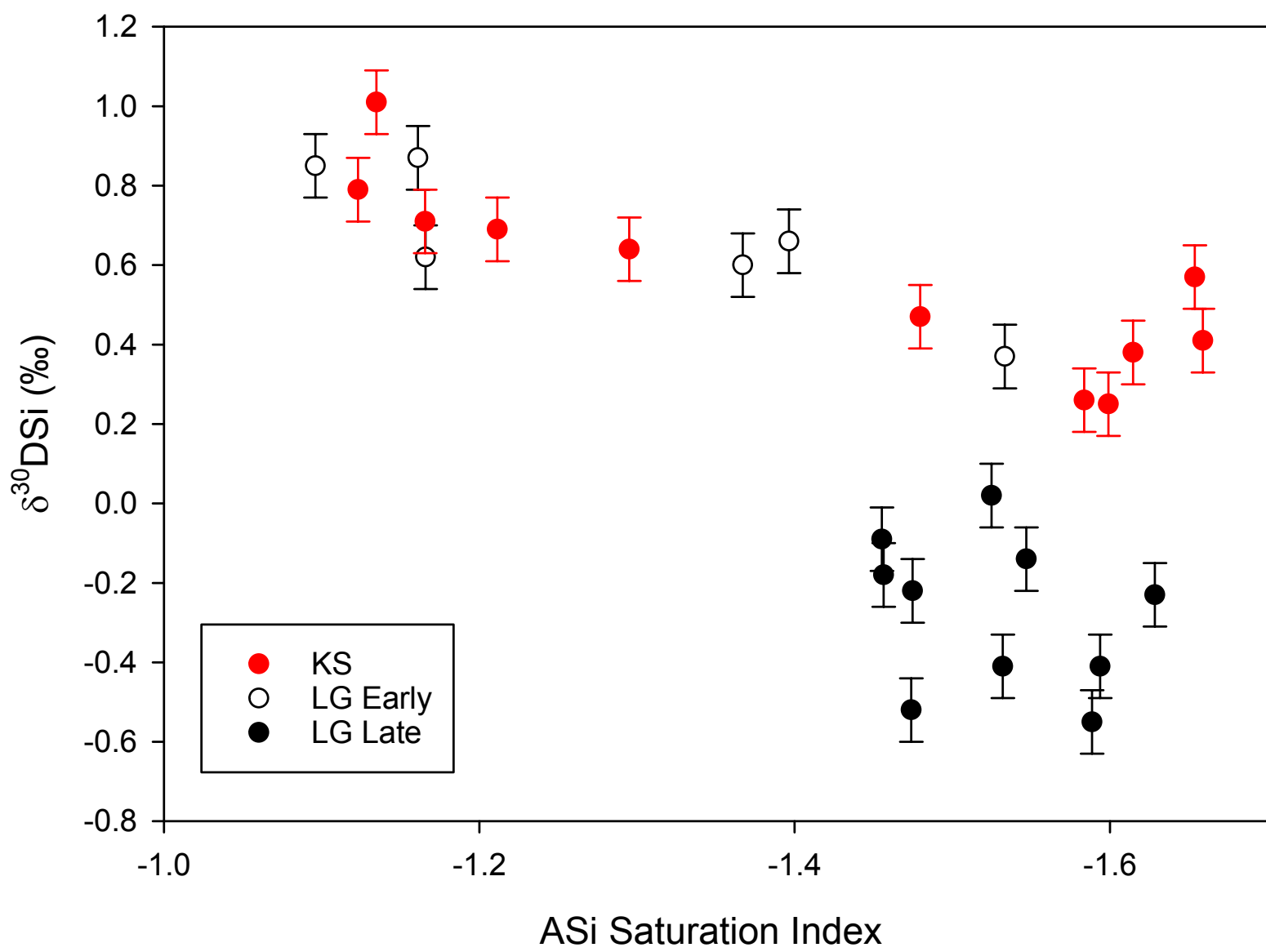


Figure8



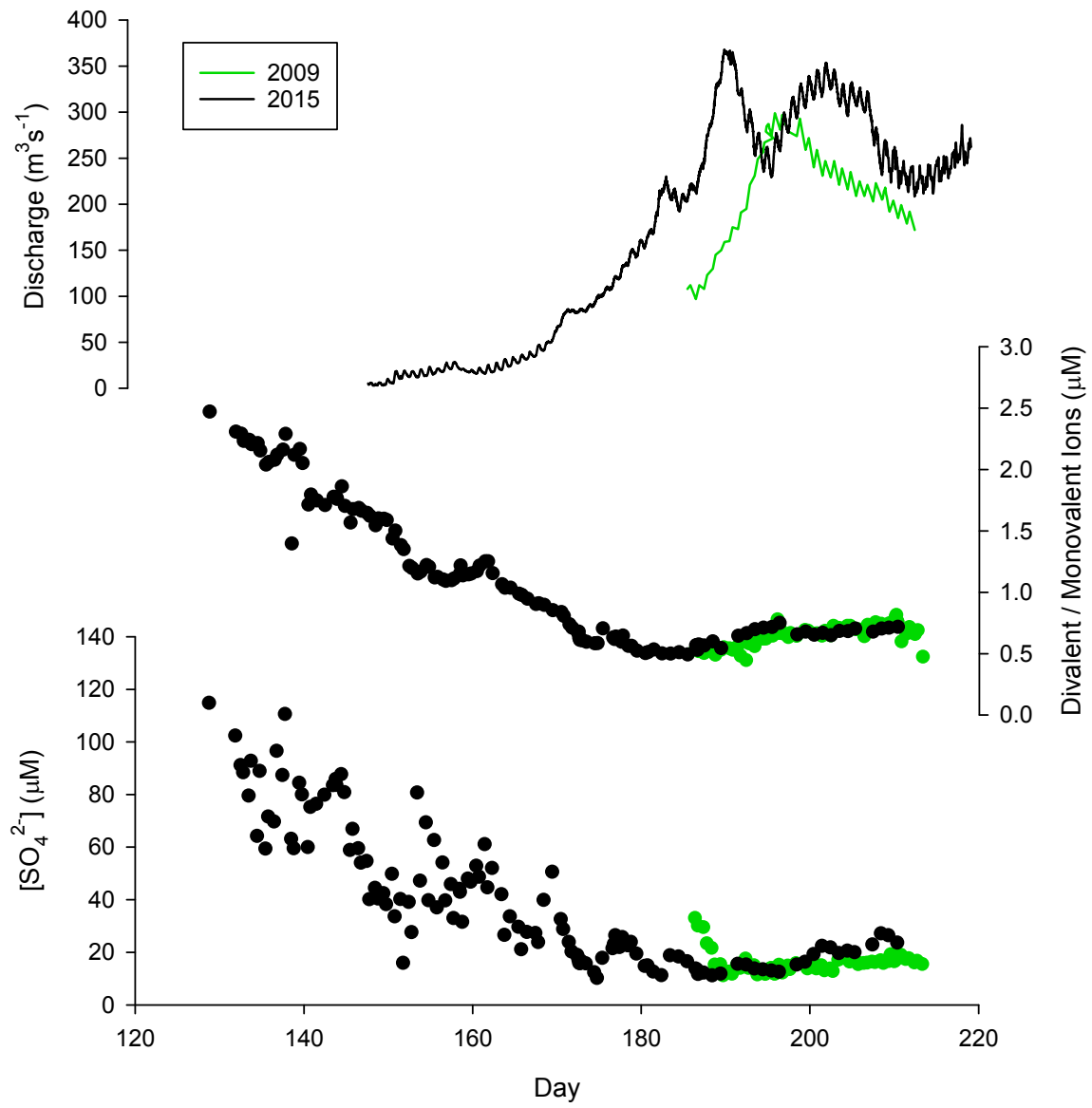


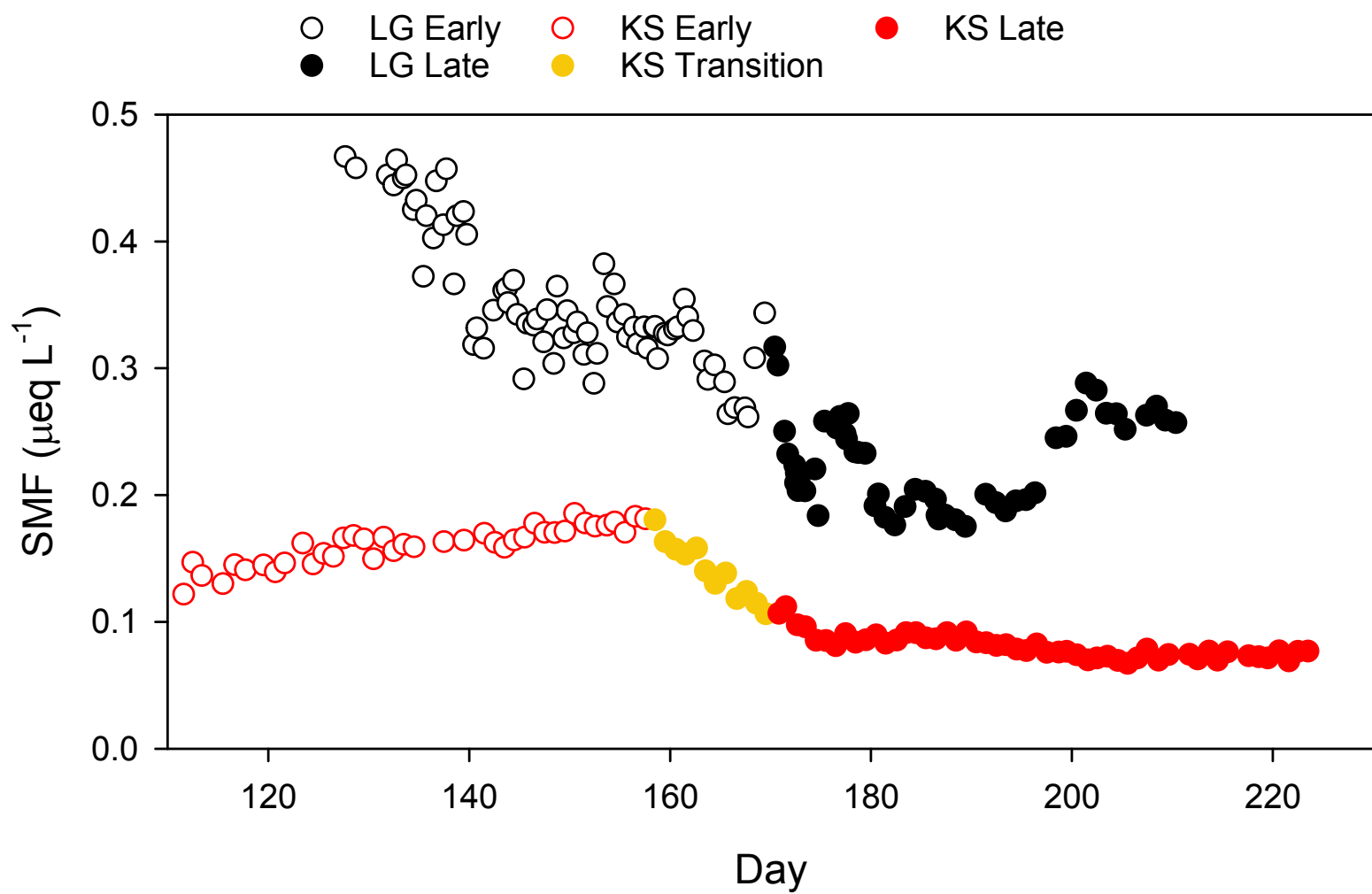


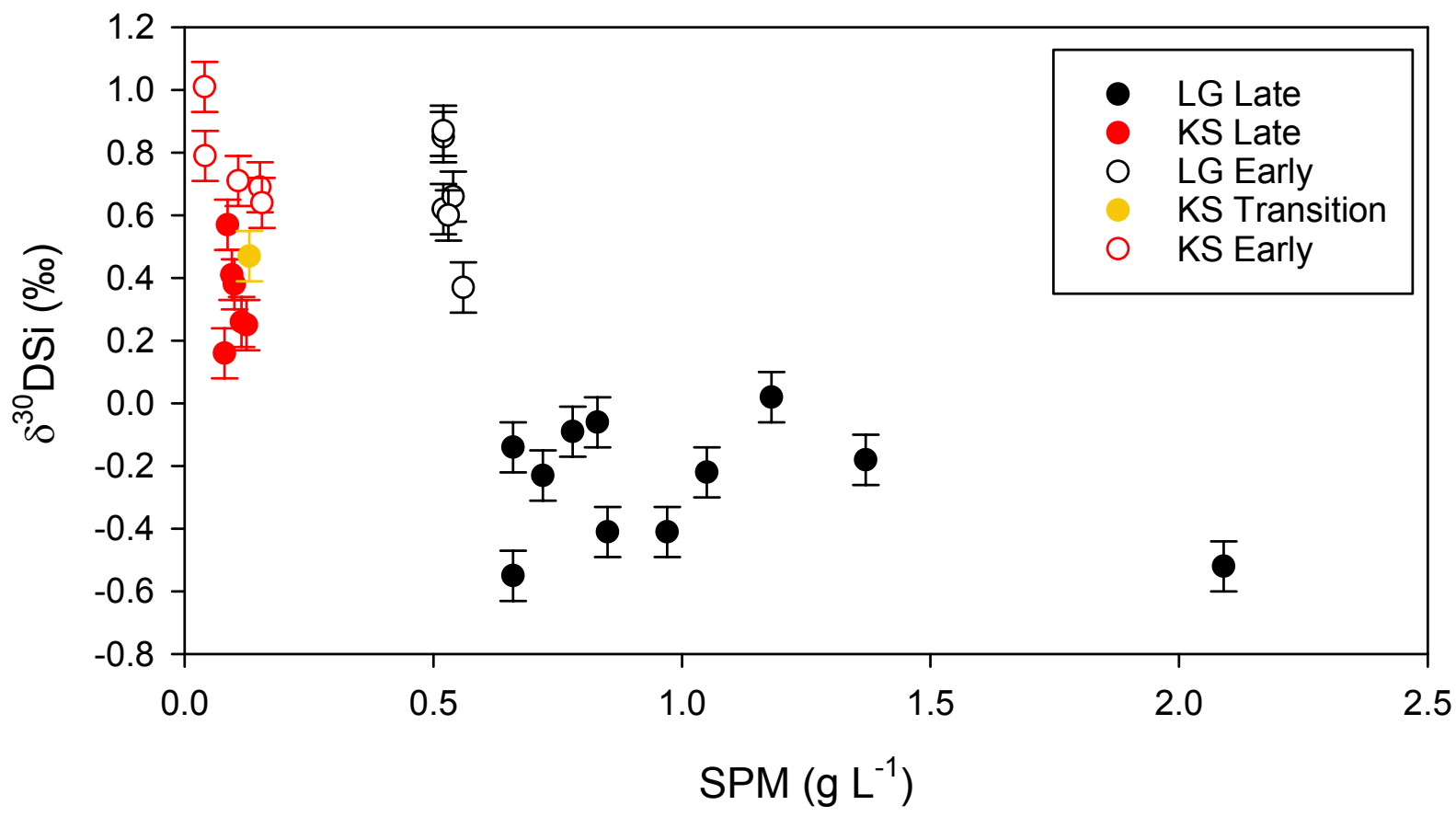




SI Figure4







## Equations

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