1	Impact of glacial activity on the weathering of Hf
2	isotopes - observations from Southwest Greenland
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33 **Abstract** Data for the modern oceans and their authigenic precipitates suggest 34 incongruent release of hafnium (Hf) isotopes by chemical weathering of the 35 continents. The fact that weathering during recent glacial periods is associated 36 with more congruent release of Hf isotopes has led to the hypothesis that the 37 incongruency may be controlled by retention of unradiogenic Hf by zircons, 38 and that glacial grinding enhances release of Hf from zircons. Here we study 39 the relationship between glacial weathering processes and Hf isotope 40 compositions released to rivers fed by land-terminating glaciers of the 41 Greenland Ice Sheet, as well as neighbouring non-glacial streams. The 42 weathered source rocks in the studied area mostly consist of gneisses, but also 43 include amphibolites of the same age (1.9 Ga). Hafnium and neodymium 44 isotope compositions in catchment sediments and in the riverine suspended 45 load are consistent with a predominantly gneissic source containing variable 46 trace amounts of zircon and different abundances of hornblende, garnet and 47 titanite.

48 Glacially sourced rivers and non-glacial streams fed by precipitation and 49 lakes show very unradiogenic Nd isotopic compositions, in a narrow range ( $\varepsilon_{Nd}$ 50 = -42.8 to -37.9). Hafnium isotopes, on the other hand, are much more radiogenic and variable, with  $\varepsilon_{Hf}$  between -18.3 and -0.9 in glacial rivers, and 51 even more radiogenic values of +15.8 to +46.3 in non-glacial streams. 52 53 Although relatively unradiogenic Hf is released by glacial weathering, glacial 54 rivers actually fall close to the seawater array in Hf-Nd isotope space and are 55 not distinctly unradiogenic.

56 Based on their abundance in rocks and sediments and their isotope 57 compositions, different minerals contribute to the radiogenic Hf in solution 58 with a decreasing relevance from garnet to titanite, hornblende and apatite. 59 Neodymium isotopes preclude a much stronger representation of titanite, hornblende and apatite in solution, such as might result from differences in 60 61 dissolution rates, than estimated from mineral abundance. The strong contrast 62 in Hf isotope compositions between glacial rivers and non-glacial streams 63 results mostly from different contributions from garnet and zircon, where 64 zircon weathering is more efficient in the subglacial environment.

65 A key difference between glacial and non-glacial waters is the water-rock 66 interaction time. While glacial rivers receive continuous contributions from 67 long residence time waters of distributed subglacial drainage systems, non-68 glacial streams are characterized by fast superficial drainage above the 69 permafrost horizon. Therefore, the increased congruency in Hf isotope 70 weathering in glacial systems could simply reflect the hydrological conditions 71 at the base of the ice-sheet and glaciers, with zircon weathering contributions 72 increasing with water-rock interaction time.

73

## 74 **1. Introduction**

75 Glacial weathering processes promote high silicate weathering rates over 76 Pleistocene glacial-interglacial cycles (e.g., Vance et al., 2009). Glacial grinding 77 of rock substrate produces fine-grained rock powder with large surface area, 78 which can be exposed to weathering in a range of settings (e.g., Anderson, 79 2007). In addition to surface area, soil age is an intrinsic factor for silicate-80 weathering rates, with rates decreasing rapidly with time of exposure (e.g., 81 Taylor and Blum, 1995). Taken together, glacial-interglacial cycles combine to 82 yield high time-integrated silicate weathering rates as reactive soil substrate is 83 produced during glacial periods and weathered effectively in intervening 84 interglacials (Foster and Vance, 2006). These interactions may be reflected in 85 the Pb isotope evolution of seawater in the northwestern Atlantic (Foster and 86 Vance, 2006; Gutjahr et al., 2009; Kurzweil et al., 2010). Other approaches to 87 study variations in glacial-interglacial weathering rates, such as oceanic 88 <sup>10</sup>Be/<sup>9</sup>Be ratios, on the other hand, suggest little change (von Blanckenburg et 89 al., 2015).

90 Continental weathering conditions also affect the seawater evolution of Hf 91 isotopes on Pleistocene and longer time-scales (Piotrowski et al., 2000; van de 92 Flierdt et al., 2002; Gutjahr et al., 2014; Dausmann et al., 2015; 2017). These 93 variations can, however, also reflect changes in weathered source rocks rather 94 than the degree of weathering congruency (Chen et al., 2012). The information 95 Hf isotopes hold has not been fully accessible to date due to our limited 96 understanding of their behaviour during weathering. Early studies of iron-97 manganese crusts and nodules, which record ambient seawater isotope

98 compositions for radiogenic isotopes, suggested that the generally incongruent 99 release of Hf isotopes during weathering is diminished during times of 100 continental glaciation (Piotrowski et al., 2000; van de Flierdt et al., 2002). A 101 cause for the incongruency of Hf isotope release is the retention of 102 unradiogenic Hf isotopes in weathering-resistant zircon (Patchett et al., 1984), 103 with more efficient release during glacial times due to glacial comminution of 104 rocks and the production of glacially strained surfaces (Piotrowski et al., 2000; 105 van de Flierdt et al., 2002). This concept has recently been reinforced by 106 observations on the Hf isotope composition in dispersed marine iron-107 manganese phases extracted from sediments that span the last deglaciation of 108 North America (Gutjahr et al., 2014). In addition, Gutjahr et al. (2014) inferred 109 that a change from a relatively congruent release of Hf isotopes during the Last 110 Glacial Maximum to a more incongruent release shortly afterwards could be 111 linked to the transition from a dominantly cold-based to a warm-based 112 Laurentide Ice Sheet.

113 A complementary mineralogical control, namely the release of radiogenic Hf 114 from preferentially weathered accessory minerals with high Lu/Hf ratios, can also affect the incongruency in Hf isotope weathering (Bayon et al., 2006; 115 116 Godfrey et al., 2007; Chen et al., 2011; 2013a,b). Thus, studies of the dissolved 117 load of rivers specifically invoke preferential weathering of apatite and titanite 118 or garnet, depending on the weathering lithologies (Bayon et al., 2006; Godfrey 119 et al., 2007). Hafnium released during weathering may, hence, become more 120 congruent with increasing soil age, as the accessory minerals are depleted. This 121 mechanism has, however, not been evaluated conclusively to date (e.g., Ma et 122 al., 2010; Bayon et al., 2016). An effect on dissolved Hf from the dissolution of 123 radiogenic accessory minerals, which also carry radiogenic Pb, appears to be at 124 odds with observations from the North Atlantic as there is no co-evolution of 125 seawater Pb and Hf - isotope compositions (Gutjahr et al., 2014).

In addition to glacial activity, mineralogy and soil age, the release of Hf isotopes has also been suggested to depend on run-off conditions and temperature (Bayon et al., 2012; 2016; Rickli et al., 2013). High run-off seems to promote the release of radiogenic Hf, as observed in catchments with different source lithologies in Switzerland (Rickli et al., 2013). Hafnium isotopes in the clay fraction of river and shelf sediments, mostly reflecting
released Hf during weathering, are positively correlated with precipitation and
temperature in catchments of various sizes and lithology from around the
globe (Bayon et al., 2016).

135 A currently unclear aspect of seawater Hf isotope compositions is the 136 relative overall homogeneity between  $\varepsilon_{Hf}$  = -2 in the Northwest Atlantic and  $\varepsilon_{Hf}$ 137 = +6 in the North Pacific (Rickli et al., 2009; Zimmermann et al., 2009a). This 138 narrow range cannot be easily reconciled with the variable riverine Hf isotope 139 compositions reported thus far (Bayon et al., 2006; Godfrey et al., 2007; Chen 140 et al., 2013b; Rickli et al., 2013; Merschel et al., 2017) and a short seawater residence time of Hf (Chen et al., 2013b; Filippova et al., 2017), similar to that 141 142 of Nd (< 500 yr, Siddall et al., 2008).

143 In summary, the interplay of environmental parameters – such as soil age, 144 glacial activity, temperature and precipitation - and mineralogical properties of 145 weathered rocks - in particular the availability of specific accessory minerals -146 is likely to govern the Hf isotope compositions of rivers. But the relative 147 significance of these aspects is not well constrained to date. In this study, we 148 seek to characterise the hydrological and mineralogical controls on the 149 congruency in Hf isotope release in the subglacial and proglacial environment of the Russell and Leverett Glaciers in West Greenland (Fig. 1). To this end, we 150 151 have characterized the weathered source rocks and derived sediments, and 152 compare them to the dissolved riverine isotope compositions, specifically in 153 glacially fed rivers and non-glacial streams.

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### 155 2. Study area

156 The studied rivers and streams are situated within the proglacial zone of the 157 Greenland Ice Sheet (GRIS), inland from Søndre Strømfjord, near the town of 158 Kangerlussuag on the west coast of Greenland (Fig. 1). Glacial waters were 159 sampled in July 2006 from the two major rivers in the region, Akuliarusiarsuup 160 Kuua and Quinnguata Kuusua, which merge at Kangerlussuaq to form the 161 Watson River. In addition, four non-glacial streams (GR11, 12, 13 and 15) and 162 a further glacial stream (GR9) were sampled. A time series of 11 different 163 samples was obtained from the main river draining Leverett Glacier in July 164 2009 (Table 1). For the purpose of this study, glacial rivers refer to those that 165 are directly fed by ice sheet melting, as opposed to non-glacial streams, which 166 are not directly linked to the ice sheet. The discharge of the latter is derived 167 from direct precipitation as well as drainage from shallow lakes, which make 168 up 5–10% of the surface area around Kangerlussuaq (Willemse, 2002).

169 The study area is dominated by amphibolite facies gneisses belonging to the 170 Ikertôq complex of the Nagssugtoqidian fold belt. The protoliths initially 171 formed at 3 to 2.7 Ga and were metamorphosed at 1.9 Ga (Fig 1., Henriksen et 172 al., 2000). Amphibolite layers and lenses within the gneisses are thought to 173 represent metamorphosed remnants of dolerite dykes (Escher et al., 1976). To 174 the southeast, the Nagssugtoqidian fold belt is bordered by the Archean Craton 175 of Greenland, where a felsic intrusion related to the Qôrqut granite (2.5 Ga) and granulite facies gneisses outcrop (Henriksen et al., 2000). Although the 176 177 felsic intrusion may be of some relevance as a source lithology for weathering 178 at, for example, the Leverett site (see section 5.4), this is unlikely to be the case 179 for the cratonic gneisses given their spatial occurrence (Fig. 1).

180 The area is characterized by an Arctic climate with a mean annual temperature of -5.7 °C (1973-1999). Seasonal variations are pronounced, with 181 an average of -19.8 °C in January and of +10.7 °C in July (Cappelen et al., 2001). 182 Annual precipitation amounts to only 149 mm (1973-1999, Cappelen et al., 183 184 2001), compared to 300 mm of evapotranspiration (Hasholt and Sogaard, 185 1978). In the unglaciated area permafrost is continuous, with an active layer 186 thickness of 0.1–2.5 m (Tatenhove, 1996). Most of the deglaciated area of the 187 study has been ice-free since at least  $\sim 6.8$  ka (Fig. 1, Levy et al., 2012 and 188 references therein). The non-glacial streams GR11, 12 and 13 are situated to 189 the west of the Ørkendalen moraines. These moraines delimit the largest areal extent of the GRIS between ~6.8 ka and the late  $19^{\text{th}}$  century, at a distance < 2 190 191 km from the current Ice Sheet margin. Stream GR15, flowing eastwards from a 192 non-glacial lake to a small lake with glacial inflow, drains sediments very close 193 to the Ice Sheet margin (see Fig. 2a in Levy et al., 2012). This stream is, 194 however, mostly within the Ørkendalen moraines and, therefore, probably not 195 influenced by the weathering of more recently exposed glacial material.

196 Recent studies have indicated that the Leverett and Russell Glaciers 197 experience strong seasonal variations in subglacial hydrology similar to those 198 observed in smaller alpine glaciers, whereby an inefficient distributed drainage 199 network transforms into an efficient channelized network of drainage channels 200 as the summer progresses (Bartholomew et al., 2010; Chandler et al., 2013). 201 Such changes have also been observed in other outlet glaciers of the GRIS, 202 suggesting that they are a common feature (Bhatia et al., 2011; Palmer et al., 203 2011; Meierbachtol et al., 2013).

204

# 205 **3. METHODS**

# 206 **3.1. Water and solid samples**

# 207 3.1.1 Glacial rivers (excluding Leverett River) and non-glacial streams

208 Both glacial and non-glacial samples were collected in July 2006 (Fig. 1, Table 209 1). Previous studies have reported Li and Mg isotope compositions of these 210 samples, also providing data on sediment load, river chemistry, pH and 211 temperature (Wimpenny et al., 2010; 2011). Some of the glacially fed rivers 212 were sampled far from the ice sheet, and thus receive small contributions from 213 non-glacially sourced waters (e.g., GR2, GR7, Fig. 1) with limited effects on 214 glacial river chemistry. Some of the non-glacial samples were coloured (GR11 215 and GR12), possibly reflecting high organic contents. Sample GR10 was taken 216 close to the harbour and contains a significant seawater component.

At each sample location approximately 15 l of river water were collected for Hf and Nd isotope analysis. Each water sample was filtered ( $0.2 \mu m$ ) within 12 h of sampling using a Sartorius frontal filtration unit. Suspended particulate material was also kept for analysis. Hafnium and neodymium were subsequently enriched from the water by co-precipitation with Fe (e.g., Rickli et al., 2009).

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# 224 **3.1.2 Leverett Glacier time series**

The main river draining the Leverett Glacier was sampled 11 times between the 6<sup>th</sup> and the 28<sup>th</sup> of July 2009 (Table 1, Fig. 1). Samples were taken every second day (excluding the 16<sup>th</sup> of July), alternating between  $\sim$ 8:30 and  $\sim$ 17:30 local time. The sampling location was  $\sim$ 1 km downstream from the glacier 229 mouth, in a stretch of turbulent flow (Fig. 1). Hence, samples taken at the edge 230 of the river are considered representative of the bulk water chemistry. In 231 addition, water samples were taken from a proglacial lake close to the river 232 sampling site, as well as from two supraglacial streams.

233 Water samples were collected in acid-cleaned 15 l HD-PE carboys pre-rinsed 234 with river water. Samples were filtered (0.45 µm) within 24 hours of sampling 235 using a peristaltic pump, clean HD-PE tubing and a filter holder made from 236 polypropylene. A second filtration was performed after the initial filtration to 237 ensure complete removal of suspended load in this high suspension river (> 2 238 g/l, see section 4.1). An unfiltered aliquot of Leverett River samples was kept 239 to determine Hf and Nd isotope compositions and concentrations in the 240 suspended sediment load. This, however, implied small but well constrained 241 corrections for the measured isotopes to account for dissolved elemental 242 contributions (<0.02  $\epsilon$ Nd, <0.2  $\epsilon$ <sub>Hf</sub>). Dissolved Nd and Hf for isotope analysis 243 were enriched from 10 to 20 l of filtered water by co-precipitation with Fe (e.g., 244 Rickli et al., 2009). A separate 1 l aliquot was kept for elemental analysis (Hf, 245 Sm, Nd) and acidified to pH < 2 with double distilled HCl.

Complementary data - including pH, temperature, runoff, suspended sediment amounts, solute concentrations and Sr/Ca isotope data - have previously been published (Bartholomew et al., 2010; Hindshaw et al., 2014). This data is used here to characterize river chemistry and hydrological conditions.

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### 252 **3.1.3 Solid samples**

253 Hafnium and neodymium isotopes and concentrations (Sm, Nd, Hf by isotope 254 dilution) have been measured on undissolved sample aliquots from Hindshaw 255 et al. (2014). These samples include: (i) powders of two orthogneisses (Ro2, 256 Ro4), two amphibolites (Ro1, Ro3) and 17 mineral separates of these rocks, 257 which were collected close to the sampling location on the Leverett River (Fig. 1); (ii) seven sediments from the proglacial environment of Leverett Glacier, 258 259 including the river bank at the dissolved load sampling location (Sed 1, 2; LC 260 sand), a side moraine (SM Sed), a proglacial lake (PGL 1) and dirt cones on 261 Leverett Glacier (MH 1, 2, Fig. 1). Note that, to avoid confusion between the

notation of Wimpenny et al. (2010) and Hindshaw et al. (2014), the rock
samples of Hindshaw et al. (2014) are relabelled from GR01 to Ro1, etc.

In addition, Hf and Nd systematics were measured for two zircon separates from Ro2 and Ro4, a rutile separate from Ro3 and four garnet and one apatite separate from catchment sediments (Table 2). Measurements of suspended particulate matter from the Leverett River, Akuliarusiarsuup Kuua and Quinnguata Kuusua, and a bulk rock analysis of an Archean granite sample originating from southwest of Quinnguata Kususa (ggu 415961, Fig. 1), complement the data.

271 Mineral separates were obtained from the  $< 425 \mu m$  fraction using heavy 272 liquids and magnetic separation to enrich minerals, followed by handpicking 273 under a binocular microscope (see Hindshaw et al., 2014). The separates, 274 excluding apatite, were initially leached for 30 minutes in 6M HCl at 80°C to 275 remove potential surface contamination, and subsequently rinsed twice with 276 MQ. All solid samples, except the zircon and apatite separates, were digested 277 on a hotplate at 120 °C for at least two days in a mixture of concentrated HF 278 and HNO<sub>3</sub> (28 M, 14 M, 4:1). Zircons were leached in the same mixture for four 279 hours and the weight loss was monitored to calculate elemental 280 concentrations. The apatite separate was dissolved in 7M HNO<sub>3</sub> to avoid 281 digestion of silicate inclusions.

282 After evaporation to dryness, all solid samples were repeatedly dissolved 283 and dried in 6M HCl to eventually yield clear solutions in 6ml of 6M HCl. 284 Isotope compositions (Hf, Nd) and elemental concentrations by isotope 285 dilution (Hf, Sm and Nd) were usually obtained on separate fractions of these 286 stock solutions. In the case of minerals, however, Sm/Nd concentrations as 287 well as Hf and Nd isotopes were obtained on the same solution fraction (spiked with a tracer enriched in <sup>150</sup>Nd and <sup>149</sup>Sm), whereas Hf concentrations were 288 289 determined separately.

290

# **3.2. Ion chromatography and procedural blanks**

Ion chromatographic procedures for the purification of Hf and Nd followed
previously detailed methods (Rickli et al., 2009; 2013) based on earlier work
(Patchett and Tatsumoto, 1980; Pin and Zalduegui, 1997; Münker et al., 2001).

295 Total procedural blanks amount to < 30 pg of Hf and Nd for the procedures used for isotope measurements, and to < 2 pg and < 7 pg for concentrations. 296 297 Average blank levels for Hf and Nd isotope measurements were < 0.05% of the 298 sample sizes. Some trace element poor mineral separates had elevated blank 299 contributions of 0.2 - 0.5% for Hf and 0.1 - 0.3% for Nd. The procedural blank 300 for Nd concentrations was < 0.1%. In the case of Hf concentrations, it was <301 0.1% for solid samples, but ranged between 0.3 and 1% for waters. No blank 302 corrections were applied to isotope and concentration data.

303

# 304 **3.3. Elemental concentrations of Hf, Sm and Nd**

Elemental concentrations of Hf, Sm and Nd for river waters, and hotplate
digests of solid samples, were measured by isotope dilution following
previously outlined methods (Rickli et al., 2009; Table 1, 2). Reproducibility of
Sm, Nd and Hf concentrations by isotope dilution was better than 1% for river
waters and rocks (replicate measurement of Lev10 and BCR-2).
Samarium/neodymium ratios reproduced to within 0.3 ‰.

Hafnium concentrations were also measured by isotope dilution on di-Lithium tetraborate fused rock powders and catchment sediments. Major elements - Si, Al, Fe, Ti, Ca, Mg, K, Na, P, Mn - and Sr concentrations for these fused samples were presented in Hindshaw et al. (2014). Fusion will completely dissolve highly resistant minerals like zircon, whereas they can remain largely unaffected during hotplate digestion.

317 Individual mineral grains in sediments from the proglacial environment of 318 Leverett Glacier (Fig. 1b) were also analysed for trace elements by LA-ICP-MS 319 using a Thermo Element XR connected to a 193 nm Resonetics ArF Excimer 320 laser (ETH Zurich). Some of these sediments were collected from the same 321 sites as in Hindshaw et al. (2014), although they represent separate sample 322 aliquots. Coordinates of sampling locations and further details are given in 323 Appendix S1. Sediments were mounted in EPOXY blocks and polished to 1 µm. 324 The laser was operated in a Laurin Technic S155 ablation cell with a spot size 325 between 20 and 43  $\mu$ m, a frequency of 4-5 Hz and a laser power density of 2 326 J/cm<sup>2</sup>. Electron microprobe (EMP) data were used as internal standards for all 327 measured minerals. NIST SRM610 was used for external standardisation and

GSD-1G glass as a secondary standard (Jochum et al., 2011a; 2011b). Raw data
were reduced off-line using the SILLS software (Guillong et al., 2008).
Uncertainties, on repeat measurements of GSD-1G at concentrations of 40-50
ppm, usually range between 2 and 3% and are < 6% for all elements. For the</li>
low Hf and Nd concentrations in many of the analysed minerals, uncertainties
are < 20 % at 5 ppm and up to 100 % close to the limit of detection (0.01 - 0.4</li>
ppm). Trace element concentrations are given in Appendix S1.

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# 336 **3.4. Isotope analysis of Hf and Nd**

337 Hafnium and neodymium isotopes were measured by MC-ICP-MS, either at the 338 University of Bristol (Neptune) or at ETH Zurich (Neptune Plus). Instrumental 339 mass bias correction followed Vance and Thirlwall (2002) in the case of Nd, and used a natural <sup>179</sup>Hf/<sup>177</sup>Hf ratio of 0.7325 for Hf. The external 340 341 reproducibility of the mass spectrometric analysis was monitored in each 342 session by repeated measurements of La Jolla for Nd and JMC 475 for Hf (n  $\geq$ 343 15), and the measured averages were used to renormalize the sample data to 344 the respective literature values (Nowell et al., 1998; Thirlwall, 1991). For Nd 345 isotopes, measured at concentrations > 50 ppb in most cases, external 346 reproducibility is < 0.2  $\varepsilon_{Nd}$  (2 SD). For Hf isotopes external reproducibility 347 depends on available Hf for analysis, varying between < 0.3  $\varepsilon_{Hf}$  for rocks, 348 sediments and many minerals, to 0.5 - 0.6  $\varepsilon_{Hf}$  for most riverine dissolved and 349 suspended sediment samples. Internal errors were  $\leq$  0.6  $\epsilon_{Hf}$  for all but 6 350 samples (2 SEM, Table 1). More details of the uncertainties in Hf isotopic 351 determination are given in Appendix S2, leading to the general conclusion that 352 internal and external errors are, to a good approximation, identical for Hf 353 isotopes.

Hafnium and neodymium isotopic compositions are expressed in epsilon units, as deviations from the Chondritic Uniform Reservoir (Jacobsen and Wasserburg, 1980; Bouvier et al., 2008).

357

### 358 **3.5. Mineral abundances**

359 Mineral abundances were obtained by point counting on thin section360 microphotographs combined with SEM back- scattered images and EDS

element maps using the ImageJ software. SEM analyses were carried out at the
University of Bristol (Hitachi S-3500N microscope equipped with Thermo
Noran energy dispersive spectrometer) and at ETH Zurich (Jeol JSM-6390LA
instrument equipped with Thermo Fisher Ultradry EDS detector coupled to a
Thermo Fisher Noran System 7).

366

# 367 **4. Results**

### 368 **4.1. Hydrochemistry**

Hydrochemical data, including runoff from Leverett Glacier in July 2009 (Fig.
2), have previously been published and discussed (Wimpenny et al., 2010;
Bartholomew et al., 2011; Hindshaw et al., 2014). The key results relevant for
the discussion of the new data on Hf and Nd isotopes are briefly summarized
here.

374 The discharge of Leverett Glacier varied strongly during summer 2009 (Fig. 375 2), reflecting seasonal variation in insolation and corresponding surface ice 376 melt. Discharge from Leverett Glacier was  $< 6 \text{ m}^3/\text{s}$  prior to the start of June 377 and then increased to a maximum value of 317 m<sup>3</sup>/s on 16<sup>th</sup> July, before 378 gradually declining again to 42 m<sup>3</sup>/s on 3<sup>rd</sup> September (Bartholomew et al., 379 2011). Four distinct discharge pulses punctuate the rising limb of the 380 hydrograph, coinciding with transient increases in suspended sediment concentrations and electrical conductivity. The last pulse started on 3<sup>rd</sup> July, 381 382 just before the samples from this study were collected (Fig. 2, the first three 383 discharge pulses are not shown). The discharge pulses have been linked to the 384 sudden drainage of lakes on the glacier surface, which are known from satellite 385 observations (Bartholomew et al., 2011). No discharge data are available for 386 the other reported glacial and non-glacial samples.

Total dissolved solids (TDS), calculated as the sum of major cations (Ca, Mg, Na, K), major anions (HCO<sub>3</sub>-, SO<sub>4</sub><sup>2-</sup>, Cl<sup>-</sup>) and Si, show clear differences for glacial rivers versus non-glacial streams (Table 1). Glacial waters yield values between 160 and 470  $\mu$ mol/l. In contrast, non-glacial streams yield much higher values between 851 and 5043  $\mu$ mol/l, more similar to the lake (TDS = 8273  $\mu$ mol/l). Total suspended sediment (TSS) ranges between 2.3 and 7.4 g/l in Leverett River and between 0.4 and 0.8 g/l in all other glacial samples. In contrast, it is very low in non-glacial streams (< 0.014 g/l) due to calm water</li>
flow. From field observations it is also clear that lake and supraglacial waters
are virtually suspension free, although no measurements are available.

397 Calcium/sodium ratios are low in all river samples, ranging between 0.6 and 398 3.9 (all ratios are reported on a molar basis, Table 1), indicating relatively 399 small contributions from carbonate weathering (e.g., Gaillardet et al., 1999). 400 Elevated K is characteristic of glacial samples, as observed before (e.g., 401 Anderson et al., 1997), resulting in K/Na ratios of 0.54 to 1.22 compared to 402 ratios between 0.03 and 0.48 in non-glacial samples. Magnesium, on the other 403 hand, is more abundant in non-glacial streams, which define a relatively 404 distinct field in major cation compositions (Fig. 3a). Bicarbonate is the 405 dominant anion, contributing a mole fraction > 0.83 in glacial and 0.67-0.84 in 406 non-glacial samples (Fig. 3b). Chlorine fractions are elevated in non-glacial 407 streams, which also leads to a clear distinction of glacial and non-glacial water 408 based on anions. Overall, the major cation and anion chemistry of the lake is 409 similar to the non-glacial streams. All waters are under-saturated with respect 410 to most minerals except, depending on the river or stream, in iron oxides and a few schist silicates (kaolinite, K-mica) (Wimpenny et al., 2010; Hindshaw et al., 411 412 2014).

413

## 414 **4.2. Dissolved hafnium and neodymium**

415 Concentration data are only available for the Leverett time series. Hafnium and 416 neodymium concentrations are highest at the start of the sampling campaign, 417 at the time of the meltwater pulse, yielding concentrations of 42.7 and 4984 418 pmol/l, respectively. The concentrations of both elements are negatively 419 correlated with discharge reflecting dilution by increased meltwater over the 420 course of the season (r = -0.86, Table 1, Fig. 2). Concentrations of many 421 elements also show diurnal cycles due to increasing runoff over the course of 422 the day (Hindshaw et al., 2014). Hafnium isotopes from the Leverett River 423 show strong variability among the first three observations, between  $\varepsilon_{Hf}$  = -18.3 424 and -9.1. Subsequently they are relatively constant, in the range of -12.4 to -9.9 425 (Fig. 2). Further glacial samples overlap with the observations from Leverett River, but also show more radiogenic signatures up to -0.9. Non-glacial streams
are distinctly more radiogenic in Hf with values of +15.8 to +46.3 (Fig. 4a).

Dissolved Nd isotopes are essentially invariant in the Leverett River at  $\varepsilon_{Nd}$  = -38.5, within the range observed for other glacial samples ( $\varepsilon_{Nd}$  = -39.5 to -38, Table 1). Non-glacial streams are more heterogeneous (-42.8 to -38.7). Stream GR14 from close to Sisimut (see Fig. 1 inset) is not included in these isotope ranges for Hf and Nd, as its dissolved Nd isotope composition suggests drainage of somewhat different source lithologies ( $\varepsilon_{Nd}$  = -30.4).

Supraglacial streams are very dilute in Hf and Nd, precluding the measurement of Hf isotopes (Table 1). Neodymium isotopes in these waters are more radiogenic than the rivers and yield values of -34.6 and -32.4. In contrast, the lake sample is characterized by unradiogenic isotope compositions of -42.3 in Nd and of -15.7 in Hf, respectively.

439

## 440 **4.3. Mineral abundances**

Estimates of mineral abundances in the analysed rocks and a catchment sediment sample (JM1 – termed Sed 3 here) have been reported previously (Hindshaw et al., 2014). Applying the same methods (see section 3.5), two additional sediment samples - Soil 1 and PGL 2 - were analysed in this study in order to further constrain mineral compositions of sediments and potential variations (Table 3).

447 Sed 3 is a fine sand collected from the riverbank at the Leverett River water 448 sampling location (location 1, Fig. 1b), similar to Sed 1 and 2 in Hindshaw et al. 449 (2014) in terms of origin and grain-size. Soil 1 is finer grained and was 450 collected beneath a few cm thick layer of soil with a grassy cover, north of the 451 Leverett River sampling location (location 2, Fig. 1b). PGL 2 comes from the 452 near shore ground of the same proglacial lake as PGL 1 of Hindshaw et al. 453 (2014 - location 3, Fig. 1b). Overall, these three samples are very similar in 454 their mineralogical composition (Table 3). Plagioclase (40 – 45 vol.%), quartz 455 (34 to 38 vol.%), K-feldspar (8 - 17 vol.%) and minor biotite are largely 456 derived from gneisses similar to Ro4 and possibly the Archean granite (see Fig. 457 4b in Hindshaw et al., 2014). Plagioclase, quartz and K-feldspar jointly 458 comprise > 84 vol.% of the minerals. The occurrence of garnet (1.5 - 2.6 vol.)

and hornblende (5.9 - 9.5%), as well as Ca-rich plagioclase, points to a small
contribution from garnet amphibolites. Hornblende is only present at
accessory levels in the characterized gneisses, contrasting with abundances of
45% and 68% in the amphibolites. Observed accessory minerals in sediments
include clino- and orthopyroxene, biotite, titanite, epidote, magnetite, ilmenite,
apatite and zircon.

465 The grey gneiss sample Ro4 has been reanalysed by point-counting here in 466 order to better constrain its accessory mineral content and to obtain an 467 estimate of a zircon free weathering endmember (section 5.5.2). Apatite 468 abundance corresponds to  $\sim 0.15$  vol.%, titanite to  $\sim 0.13$  vol.% and zircon to 469  $\sim 0.02$  vol.%. The corresponding wt.% are  $\sim 0.17$ ,  $\sim 0.17$  and  $\sim 0.034$ . The 470 analysis revealed that epidote and chlorite show strong abundance variations. 471 While the initial analysis gave an estimate of 1 vol.% chlorite and 7 vol.% 472 epidote (Hindshaw et al., 2014), an estimate over six times the initial area, 473 corresponding to 500 mm<sup>2</sup>, yields 6 vol.% chlorite and 2 vol.% epidote. Epidote 474 - which also occurs as veins - and mostly chlorite replace primary biotite and 475 minute amounts of hornblende. Element maps show the existence of epidote 476 (~ 80 %) and epidote-allanite (~ 20 %) with < 5 wt.% REE deduced from EMP 477 analyses. Similarly, REE-poor and REE-rich phosphates are observed possibly 478 resulting from metamorphic depletion of primary apatite minerals (Harlov et 479 al., 2002).

For elemental and isotope budgets (sections 5.3 - 5.5) mineral weight percent is more useful than modal abundance. The density of catchment sediments is ~ 2.7 g/cm<sup>3</sup> based on mineral abundances, which is used for the conversion. Table 3 also lists weight percent ranges for minerals in sediments.

# 485 4.4. Hafnium and samarium/neodymium isotope systematics in 486 sediments, rocks and mineral separates

487

## 488 4.4.1 Concentrations

Hafnium and neodymium concentrations are homogenous in hotplate digestsof suspended load, spanning a range of 1.3-2.7 and 29.3-33.9 ppm, respectively

491 (Table 2, Fig. 5). Catchment sediments from the proglacial area of Leverett are

more heterogeneous (mostly in the range of 2.4-7.5 ppm for Hf and 16.2-25.5
ppm for Nd) as are the bulk rocks (Hf: 0.7-3.7 ppm; Nd: 5.1-27.3 ppm). Among
the catchment sediments Sed 1 is peculiar, with distinctly high concentrations
in Hf and Nd (10.4 and 40.3 ppm, respectively). Hafnium concentrations
measured in hotplate digests are between 42% and 91% of those obtained on
fused powders. The implications of these differences in Hf concentrations are
further discussed in section 5.5.2.

499 Mineral separates rich in Nd include titanite (772 and 1110 ppm), apatite 500 (636 ppm) and to a lesser degree epidote, zircon, chlorite and hornblende 501 (vielding up to 17.3-99.4 ppm, Table 2, Fig. 5). Hafnium is mostly contained in 502 zircon separates (> 9000 ppm) and is also distinctly high in titanite (16.5 and 503 46.4 ppm). All further non-feldspar mineral separates - hornblende, garnet, 504 clinopyroxene, epidote, chlorite - yield Hf concentrations between 0.3 and 3.2 505 ppm. Feldspar separates are low in Hf and Nd ranging between 0.05 and 0.65 506 ppm and 0.2 and 2.6 ppm, respectively.

507 In most cases, concentrations observed in single sedimentary mineral 508 grains obtained by LA-ICP-MS overlap with the range observed in mineral 509 separates (Fig. 5, Appendix S1). Hafnium in feldspar separates (0.05 to 1.5 510 ppm) is, however, somewhat higher than in measured sediment grains (mostly 511 < 0.05 ppm). An analogous statement holds true for Nd in K-feldspar, where Nd 512 amounts to 2.6 ppm in the separate and is < 0.2 ppm in single grains. These 513 discrepancies possibly reflect, at least in part, the impact of inclusions in 514 feldspars, which raise observed concentrations to higher values for mineral 515 separates.

Samarium/neodymium and Hf/Nd ratios of sedimentary grains also overlap
with mineral separates (Fig. 5, Appendix S1). Furthermore, Sm/Nd ratios
suggest that sedimentary plagioclase is largely derived from gneisses whereas
hornblende is sourced from mafic rocks, consistent with the previous notion
based on mineral abundances in bulk rocks and plagioclase chemistry (section
4.3; Hindshaw et al., 2014).

Biotite (Hf < 0.05 ppm, Nd < 1.5 ppm), orthopyroxene (Hf < 0.06 ppm, Nd <</li>
0.6 ppm), ilmenite (Hf < 0.9 ppm, Nd < 0.2 ppm) and oxides (Hf, Nd < 0.05</li>
ppm) are depleted in REE and Hf (Appendix S1) and only occur as accessory

525 minerals. Hence, these minerals are irrelevant for elemental budgets in rivers 526 and sediments and will not be considered further in the discussion. Apatite, on 527 the other hand, is rich in REE and could affect sedimentary and riverine 528 dissolved Nd budgets, and potentially dissolved Hf isotope compositions 529 (Bayon et al., 2006).

530

# 531 **4.4.2 Isotopes**

532 Neodymium isotopes and Sm/Nd ratios in suspended load and catchment 533 sediments are confined to the range between the grey (Ro4,  $\varepsilon_{Nd}$ = -39) and the 534 pink orthogneiss (Ro2,  $\varepsilon_{Nd}$  = -31.4, Table 2, Fig. 4a). The suspended load is 535 virtually constant in  $\varepsilon_{Nd}$  (-36.8 to -35.7) and on average ~3 units more 536 radiogenic than Ro4. Catchment sediments are mostly between -34.2 and -33.3, but also include one sample with a signature similar to Ro4 and the rivers 537 538 (MH1, -38.3). Hafnium isotopes show the least radiogenic values in felsic bulk 539 rocks ( $\epsilon_{Hf}$  = -59.9 to -55.1), and are progressively more radiogenic in catchment 540 sediments (-54.4 to -47) and the suspended load (-46.4 to -34.6). The latter is 541 offset from the dissolved load by at least ~20  $\epsilon_{Hf}$  (Fig. 4b). The two 542 amphibolites (Ro1, Ro3) are very distinct in Nd-Hf isotope space, yielding  $\varepsilon_{Hf}$  > 543 +26 and  $\varepsilon_{Nd}$  > -6.7, respectively.

544 The four rock samples Ro1 to Ro4 yield ages for the Sm/Nd isotope system 545 ranging between 1746 and 1887 Myr, consistent with the literature (Table 4, 546 Henriksen et al., 2000). With the exception of Ro1, the errorchrons are, 547 however, characterized by mean square weighted deviations  $\geq$  140. Analysed 548 mafic rocks and constituent minerals have Sm/Nd ratios  $\geq 0.267$  and are 549 shifted towards more radiogenic  $\varepsilon_{Nd}$  at a given Sm/Nd ratio, resulting in higher 550 initial <sup>143</sup>Nd/<sup>144</sup>Nd ratios for their isochrons (Fig. 4a, Table 4). The garnet 551 separates from PGL 2 are also characterized by high Sm/Nd ratios, but appear 552 to be genetically linked to the gneisses rather than the sampled amphibolites 553 (Fig. 4a). High Sm/Nd ratios and radiogenic Nd are also observed for titanite 554 and hornblende within the gneisses, and in chlorite and epidote of Ro4. The 555 measured Archean granite is less radiogenic than all other samples, yielding an 556  $\varepsilon_{Nd}$  of -47.3.

557 Consistent with the bulk rock Hf isotope compositions, mineral separates 558 from the amphibolites are typically more radiogenic than those from the 559 gneisses. Radiogenic Hf isotope compositions are, however, observed in 560 gneissic titanite, epidote and hornblende (Table 2, Fig. 4b). The most notable 561 minerals for the Hf systematics are the zircons in the gneisses – unradiogenic 562 and with high Hf concentrations – and the extremely radiogenic garnets and 563 apatites, yielding  $\varepsilon_{\rm Hf}$  between +872 and +1940.

564

565 **5. Discussion** 

# 566 5.1. Potential for external sources of hafnium and rare earth elements in 567 rivers and streams

In principle, a significant fraction of the Hf and REEs in the sampled waters could be derived from external sources, rather than from weathering within the catchment. Precipitation as well as snow and ice-melt are insignificant sources of Hf and Nd, yielding a maximum contribution of 0.03% and 0.01% for the sampled lake. The calculation assumes element/Cl<sup>-</sup> ratios similar to surface seawater of the neighbouring Labrador Sea (Filippova et al., 2017) and precipitation as the only source of Cl<sup>-</sup> in the catchment (e.g., Yde et al., 2014)

575 Another potentially important external source is wind-blown dust from 576 outside the catchment (Tepe and Bau, 2015). The samples likely to be affected most by external dust are the supraglacial streams, where any deposited dust 577 578 might be less diluted compared to the sub- and proglacial area. Neodymium 579 isotopes in two supraglacial streams are about 5  $\varepsilon$ -units more radiogenic than 580 the glacial samples (Table 1). This could imply addition of some external Nd to 581 the catchment, but could also reflect a specific mineral assemblage exposed to 582 weathering on the glacier resulting from sorting by wind *within* the catchment. 583 Mineral sources of radiogenic Nd are abundant in the catchment rendering this 584 interpretation entirely plausible (Fig. 4a).

585 Nevertheless, a small impact on dissolved isotopes in supraglacial streams 586 due to partial dissolution of external dust cannot be precluded. Any impact on 587 Leverett river waters is, however, unlikely since they are more concentrated in 588 Hf and Nd than supraglacial streams by at least a factor of 69 and 116, 589 respectively (Table 1). A potential source of REEs in Western Greenland is

590 Asian dust (Tepe and Bau, 2015). The more radiogenic signatures in supraglacial streams could be accounted for by  $\sim 15$  % Nd from such dust, 591 assuming a corresponding average Nd isotope composition of  $\sim$  -6 (Zhao et al., 592 593 2014). A similar dust-derived Nd contribution to the Leverett River would 594 correspond to a Nd fraction < 0.12 %, and would imply a true weathering 595 signal that is only marginally less radiogenic, by < 0.04  $\varepsilon_{Nd}$ . Furthermore, the 596 observed dissolved isotope compositions are entirely consistent with sampled 597 rocks and sediments being the sole sources of dissolved Nd and Hf and no 598 external sources are implied in any of the observations (section 5.3 - 5.5).

599

# 600 **5.2. Carbonate versus silicate weathering**

601 Solutes in glacial and non-glacial waters could reflect different mineralogical 602 sources because the weathering process beneath the GRIS is different from the 603 proglacial area in many respects, including water-rock interaction time, 604 regolith/soil exposure age and access to atmospheric oxygen (Yde et al., 2010; 605 Wimpenny et al., 2011; Hindshaw et al., 2014; Scribner et al., 2015). Although 606 the weathered rocks are likely very similar, the different conditions may 607 induce differences in mineral weathering reactions. For example, glacial 608 activity could lead to continuous supply of trace carbonates from the bedrock, 609 resulting in larger carbonate weathering contributions for glacial waters (e.g., 610 Anderson et al., 2000). Hafnium/calcium ratios in Leverett River samples exceed  $3 \times 10^{-7}$  (Table 1), and are about two orders of magnitude higher than 611 612 in the leached carbonate fraction of natural carbonate rocks ( $\sim 7 \times 10^{-9}$ , Rickli 613 et al., 2013). These dissolved ratios are most likely much lower than the 614 weathering released Hf/Ca ratios as a result of stronger adsorption of Hf onto 615 suspended load. The observed variations in Hf isotopes between glacial and 616 non-glacial waters, as well as temporal changes in Leverett River, therefore 617 reflect changes in the weathering of different silicate minerals.

618

## 619 **5.3. Hafnium and neodymium in sediments**

The major element composition of catchment sediments and their plagioclase
grains suggests large contributions from grey relative to pink orthogneisses
and amphibolites (Hindshaw et al., 2014). The catchment sediments can,

hence, be modelled as Ro4 complemented with additional minerals like garnetand hornblende at their observed abundance in sediments (Table 3).

625 Catchment sediment Nd isotope compositions are consistent with 10 wt.% 626 hornblende and 0.25 wt.% titanite added to Ro4, whereas the same additions 627 scaled by a factor of 0.55 yield compositions similar to the suspended load (Fig. 628 6a). Titanite enrichments in catchment sediments relative to Ro4 are 629 supported by elevated Ti concentrations (Hindshaw et al., 2014). Smaller 630 effects in Nd isotope compositions are also modelled for garnet (~ 0.4  $\varepsilon_{Nd}$  for 4 wt.%) and apatite (~ 1  $\varepsilon_{Nd}$  for 0.2 wt.%) addition. There is, however, no 631 632 evidence for apatite enrichment in sediments relative to Ro4.

633 In terms of the Nd budget, the modelled addition of titanite and hornblende 634 to Ro4 produces a Nd concentration of about 29 ppm, versus observed values 635 in catchment sediments that are mostly between 16.2 and 25.5 ppm (Table 2). Higher Nd concentrations are, however, observed in the suspended load (29.3-636 637 33.9 ppm, Table 2). The catchment sediments are probably depleted in REErich accessory minerals like allanite and monazite relative to the suspended 638 639 load and Ro4, likely as a result of prevailing small sizes of these mineral grains 640 (e.g., Garzanti et al., 2008).

641 As mentioned earlier, Hf isotope compositions in sediments cover a nearly continuous range, from  $\varepsilon_{Hf}$  = -54.4 in Sed 2 to -34.6 in a river suspended load 642 sample (GR5, Fig 4b). Titanite and hornblende enrichment in sediments also 643 644 shifts Hf isotope compositions to more radiogenic values compared to Ro4 (Fig. 6b, solid black line). The shift is, however, not sufficient to explain the Hf 645 646 systematics: suspended load usually falls above the mixing line between Ro4 647 and titanite/hornblende in  $\varepsilon_{Hf}$  versus  $\varepsilon_{Nd}$  space, whereas catchment sediments are less radiogenic. The deviations from the mixing line can be reproduced by 648 649 variable additions of zircon and garnet (Fig. 6b), using observed Hf 650 concentrations as an additional constraint on zircon abundances (Fig. 6c). The 651 calculations suggest that Hf isotopic variability in catchment sediments reflects 652 variable abundances of garnet, ranging between  $\sim 1.1$  and  $\sim 4$  wt.%, coupled to variable zircon abundances of  $\sim 0.06 - 0.7$  wt.% in excess of Ro4. The large 653 654 range of Hf concentrations in catchment sediments, between 10.4 and 2.4 ppm, 655 is controlled by zircon. Isotopic variations, however, also relate to variable

656 garnet abundances. A mixing calculation that only involves variation in 657 amounts of zircon (Fig. 6c, grey dashed line) produces a larger range in Hf 658 isotope compositions than observed. It should be noted that the Hf 659 concentrations used in this modelling relate to those accessible in hotplate 660 digestions, so that they represent a fraction of the constituent zircon (see 661 section 5.5.2).

For river suspended loads a mixing model consistent with the isotope and concentration data is presented in Fig. 6b and c. Lower Hf concentrations in some of the suspended sediments imply a lower abundance of zircon than in Ro4. This is accounted for by including removal of zircons to model these observations, reflected by the negative signs in zircon ranges (Fig. 6b and c).

667

### 668 **5.4. Sources of REE in rivers and streams**

669 Dissolved Nd in glacial river samples is isotopically very similar to the grey 670 orthogneiss Ro4 (Table 1, 2; Fig. 4a). This observation suggests that the Nd 671 isotope budget of glacial rivers is likely dominated by the nearly congruent 672 dissolution of such gneisses. Clearly less radiogenic Nd isotope compositions 673 than in Ro4 are, however, observed in the lake and two proglacial streams (Fig. 674 6d). Accessory minerals rich in REE, such as allanite and monazite, are 675 probably more represented in these samples. In support of this interpretation, 676 a missing unradiogenic Nd pool is implied in the Nd isotope composition of the 677 bulk Ro4 relative to the measured minerals, with an approximate Nd isotope 678 composition of -45.6 (Table 2). Selective weathering of unradiogenic Nd has 679 been observed previously on glacially produced sediments, although the 680 mineralogical source was not identified (Andersson et al., 2001).

Another possible source of unradiogenic Nd are Archean granites ( $\varepsilon_{Nd}$  = -681 682 47.3). However, if such granites were an important source of dissolved Nd, one 683 would expect a clear shift towards less radiogenic Nd in Quinnguata Kuusua, 684 given that it flows across the intrusion (GR7, Fig. 1). Rather, its Nd isotope 685 composition of -39 is similar to samples from Akuliarusiarsuup Kuua and the 686 Leverett River, which argues against Archean granites as a significant lithology 687 controlling dissolved Nd isotope compositions. The Nd isotope composition of 688 Quinnguata Kuusua is apparently acquired in the subglacial environment and the interaction with the intrusion is too short to affect the dissolvedcomposition.

691

# 692 **5.5. Sources of hafnium in rivers and streams**

693 The significance of a mineral for the dissolved Hf in rivers and streams will 694 depend on its abundance, its Hf concentration and its susceptibility to 695 weathering. In a simple illustrative case a mineral is abundant, rich in Hf and 696 dissolving easily and will, hence, yield a large contribution to the dissolved Hf. 697 The isotopic impact of a mineral also depends on the isotopic contrast relative 698 to other minerals. Minerals with high Lu/Hf ratios, such as apatite and garnet, 699 have highly radiogenic isotope signatures, so that even small contributions 700 from their dissolution will substantially affect the dissolved Hf signatures.

In sections 5.5.3 and 5.5.4, we identify the sources of Hf in solution from isotope systematics in  $\varepsilon_{Hf}$  versus  $\varepsilon_{Nd}$  space and compare them with expectations from dissolution rates, given the observed mineral abundances, elemental concentrations and isotope compositions. But before doing so, we discuss the potential relevance of mineral inclusions (5.5.1) and the implications of a zircon free Ro4 for dissolved Hf isotope compositions (5.5.2).

707

### 708 5.5.1 Mineral inclusions

709 Observed mineral inclusions include apatite in clinopyroxene, hornblende and 710 plagioclase, as well as quartz and clinopyroxene in garnet. Most likely, there 711 are further minerals present as inclusions not mentioned here. The mineral 712 digests could, hence, have a certain bias towards these inclusions, such that the 713 mass balance calculations could be affected and misleading to some degree. 714 The effect is expected to be small for two reasons: (1) estimates of mineral 715 abundances also include observed mineral inclusions so that there should be 716 no underestimation of accessory minerals such as apatite; (2) the key minerals 717 used in the calculations show consistent isotope and concentration systematics 718 where several measurements are available - garnets, zircons, titanite, 719 hornblende (Table 2) – and are also consistent with the literature for apatite 720 (e.g., Barford et al., 2004).

721 Feldspars separates are likely biased by inclusions since all separates yield 722 higher concentrations than observed on single mineral grains by LA-ICP-MS 723 (Table 2, Appendix S1). The weathering of these inclusions is likely to be 724 strongly coupled to the weathering of the host, as the inclusions are typically 725 only a few micrometres. Hence, it is justified to use the measured 726 concentrations and isotope compositions of these separates, although the 727 derived Hf and Nd will not reflect feldspar weathering in a strict sense. 728 Measured feldspar isotope compositions are not extreme, so that a strong bias 729 from mineral inclusions can be precluded (Table 2).

The model calculations use average concentrations and isotope compositions for separates of the same mineral – 5 for garnets, 2 for hornblende, titanite and zircon – also increasing their robustness. Note that, for hornblende, only the values for those in amphibolites are used (Ro1 and Ro3), because they are represented in the sediment (section 4.3).

735

### 736 **5.5.2 Estimate of a zircon-free weathering endmember**

737 The incongruency in Hf isotope weathering is thought to reflect a combination 738 of the retention of unradiogenic Hf in zircons and the preferential weathering 739 of high Lu/Hf phases (e.g., Bayon et al., 2006; Godfrey et al., 2007; Chen et al., 740 2013a). A constraint on the Hf isotope composition of a model zircon-free 741 weathering endmember in the studied catchment is therefore valuable to 742 compare the relative magnitude of these two phenomena. A large fraction of 743 the Hf in the studied rocks and catchment sediments is accessed through 744 hotplate digestion (42%-90%, Table 2), compared, for instance, to a fraction of 745 1.1% in young gneisses from the Swiss Alps (Verzasca catchment, Rickli et al., 746 2013). This means that zircons are easy to digest in the studied catchment, 747 probably as a result of their age and the accumulated fission damage. Hotplate 748 digests thus do not reflect the zircon-free Hf isotope compositions. Instead, an 749 estimate of the zircon free Ro4 is derived from observed mineral abundances 750 using two different but not fully independent mass balance considerations (see 751 Appendix S3). The best estimate of this zircon free endmember yields a Hf 752 concentration of 0.18 ppm and an  $\varepsilon_{Hf}$  of -5. Extreme estimates correspond to 753 0.17 ppm / -1.2  $\varepsilon_{Hf}$  and 0.22 ppm / -16  $\varepsilon_{Hf}$ , respectively. It could be argued from these data alone that the glacial rivers result from the relatively congruent dissolution of the zircon free Ro4 only, but this is not a plausible interpretation given the large effect of garnet on dissolved riverine  $\varepsilon_{\rm Hf}$  (section 5.5.3).

758 The estimate of the zircon free Hf concentration in Ro4 conflicts with 759 models that explain the seawater array in terms of a consistent partitioning of 760 Hf between zircons versus other upper crustal minerals. These models suggest 761 that only  $\sim 65\%$  of the Hf is zircon-bound (Chen et al., 2011; van de Flierdt, 2007), whereas it seems to be ~ 95% in Ro4 and ~ 99% in the Verzasca 762 763 catchment (Rickli et al., 2013). The underlying model assumptions, namely that 764 there is a close correspondence between Hf isotopes released by continental 765 weathering and the seawater array (van de Flierdt et al., 2007) or a constant 766 incongruent weathering effect in  $\varepsilon_{Hf}$  at any given  $\varepsilon_{Nd}$  (Chen et al., 2011), are not 767 well constrained to date. Although this may be the case on a continental scale, 768 it is certainly not the case for small catchments, which yield highly variable 769 riverine  $\varepsilon_{Hf}$  at near constant  $\varepsilon_{Nd}$  (compare for instance data in Bayon et al., 770 2006; Godfrey et al., 2007 and Rickli et al., 2013).

771

### 772 **5.5.3 Sources of dissolved hafnium from isotope systematics**

773 To start with, we note that the relevance of allanite and monazite for dissolved 774 Hf isotopes is not fully constrained since no mineral separates were measured 775 for their Hf isotope compositions and concentrations. The mineral abundance 776 of allanite is unlikely to exceed  $\sim 0.12$  wt.% based on REE concentrations in 777 sediments, assuming Nd concentrations of  $\sim 21'000$  ppm as measured in a 778 sedimentary allanite grain (Appendix S1) in the range of literature data (e.g. 779 Gromet and Silver, 1983; Boston et al., 2017). Monazite abundance could be at 780 most ~0.025 wt.% assuming Nd concentrations of 10 wt.%. Given typical Hf 781 concentration in allanite < 0.7 ppm and in monazite < 1 ppm (Rubatto et al., 782 2006; Boston et al., 2017), allanite may yield an effect on dissolved Hf similar 783 to apatite, which is discussed below. Monazite, on the other hand, can hardly be 784 significant for dissolved Hf signatures.

Insights can be gained into the relative significance of apatite, titanite,garnet and hornblende as sources of radiogenic Hf in solution if they are added

to a model accessory mineral-free Ro4. The corresponding Hf concentration and isotope composition is estimated from feldspar, chlorite and epidote abundances in Ro4 and yields a value of  $\varepsilon_{Hf}$  = -38 and 0.15 ppm Hf. The corresponding Nd isotope composition, representing the same minerals and the REE-rich accessory minerals allanite and monazite, can also be derived from mass balance ( $\varepsilon_{Nd}$  = -42.7, Nd = 24.3 ppm).

793 At the observed abundance, not taking into account individual mineral 794 weathering rates (see section 5.5.4), hornblende, garnet, titanite and apatite 795 contribute significantly to radiogenic Hf isotopes in solution (Fig. 6d, dashed 796 black lines). The influence on dissolved signatures decreases from garnet to 797 titanite, hornblende and apatite. Combining only apatite, titanite and 798 hornblende with the accessory mineral free Ro4 produces a trajectory, which 799 yields Hf and Nd isotope compositions similar to glacial waters, although 800 slightly too radiogenic in  $\varepsilon_{Hf}$  and  $\varepsilon_{Nd}$  (Fig. 6d, solid black line). The possibility 801 that the slightly different  $\varepsilon_{Nd}$  is due to low dissolution rates of horblende is 802 discussed later (section 5.5.4). The offset in  $\varepsilon_{Hf}$  is unexpectedly low, since 803 garnets are common in the catchment and should have a significant impact on 804 all water samples. Some zircon likely compensates for garnet weathering: 805 Adding 2 wt.% garnet and 1/500 of this amount of zircon to the mixing 806 trajectory at the Nd isotope composition of Ro4 (black circle) yields Hf isotope 807 composition in the range of glacial waters. Similarly, non-glacial streams can be 808 modelled by a combination of 3 wt.% garnet and 1/10'000 this amount of 809 zircon, or 2.5 wt% garnet and no zircon (Fig. 6d, lines with arrows).

The model does not claim that only zircon can compensate for radiogenic Hf from garnet. However, the zircon-derived amount of Hf in the model that reproduces glacial waters is 0.4 ppm. This is nearly three times the Hf concentration of the accessory mineral free Ro4. Hence, most of the compensation must be accomplished by zircon.

Two further findings should be mentioned here. Firstly, very efficient subglacial weathering of zircon, meaning that most of the hotplate digestion accessible Hf is released, can be precluded since radiogenic accessory minerals could not compensate for this process. A complete dissolution of the measured Ro4 would imply unrealistic garnet contributions of ~10 wt.% to explain glacial samples (Fig. 6d, grey dashed line). Secondly, it also seems unlikely that
hornblende, titanite or apatite are strongly overrepresented in the dissolved
load as this would result in more radiogenic dissolved Nd isotope compositions
than observed (Fig. 6d).

824

# 825 **5.5.4 Mineral weathering rates**

826 Mineral weathering reactions are confined to mineral surfaces (e.g., Brantley 827 and Olsen, 2014). As a result dissolution rates scale with surface area and small 828 minerals will contribute much to the weathering flux. In the subglacial 829 environment all minerals are likely abundant in the fine fraction as a result of 830 glacial grinding. Hence, differences in dissolution rates relating to surface area 831 are reduced. This could be different in non-glacial streams since much of the 832 weathered substrate can be relatively coarse. Larger garnet contributions in 833 these streams could therefore reflect small grain sizes of garnet relative to 834 other minerals in the proglacial zone, but this is not consistent with observed 835 garnet sizes in catchment sediments.

836 Figure 7 shows laboratory based surface area-normalised dissolution rates 837 as a function of pH for many relevant minerals in the catchment (Guidry and Mackenzie, 2003; Palandri and Kharaka, 2004 and references therein). 838 839 Conditions correspond to 25°C and solutions far from mineral saturation. At 840 the neutral to alkaline pH relevant to the sampled waters (Table 1, pH = 6.8 -841 9.3) apatite dissolution rates are similar to, but slightly faster than, the 842 dissolution of almandine garnet. Titanite dissolution is probably similar to 843 garnet (see Morton, 1984; Bateman and Catt, 1985). Laboratory based 844 dissolution rates observed for hornblende are variable (Sverdrup, 1990; 845 Frogner and Schweda, 1998; Golubev et al., 2005). In field studies hornblende 846 appears to be relatively stable (Colman, 1982), so that the lower laboratory-847 based rates are possibly more representative. The dissolution rates for garnet, 848 apatite and titanite, hence, suggest that the relative abundances of these 849 minerals should be reflected in dissolved Hf as their dissolution rates are 850 relatively similar at neutral to alkaline pH. Hornblende, on the other hand, may 851 be underrepresented relative to apatite, titanite and garnet. At conditions close 852 to mineral saturation, dissolution rates decrease (e.g., Brantley and Olsen,

853 2014), which provides an explanation for the different Hf isotope compositions

in glacial and non-glacial waters as discussed below (section 5.6.1.).

855

# 856 **5.6. Controls on dissolved hafnium isotopes**

857 The two most striking features regarding dissolved Hf isotope compositions 858 are the large variations at the beginning of the Leverett time series (Fig. 2) and 859 the large isotopic contrast between glacial and non-glacial samples (Fig. 4). 860 Previous work on radiogenic Sr and Pb has highlighted two key aspects, which 861 can affect the congruency in released isotopes relative to weathered source rocks, namely regolith exposure age (e.g., Blum and Erel, 1997; Bullen et al., 862 1997; Harlavan et al., 1998), and water-rock interaction time (Bullen et al., 863 864 1996; Hindshaw et al., 2014; Arendt et al, 2016). The data obtained in this 865 study provide some insights into the significance of water-rock interaction 866 time (5.6.1) and regolith exposure (5.6.2) for the release of Hf isotopes, given 867 the constraints on landscape exposure to weathering from moraine ages (Fig. 868 1).

869

### 870 **5.6.1. Water-rock interaction time**

Water-rock interaction times are very variable for waters discussed here. 871 Subglacial water in distributed systems can interact with sediments for more 872 873 than half a year, from the cessation of the channelized system in 874 August/September until the next melting season (Chu et al., 2016). These 875 waters can yield high dissolved concentrations (Graly et al., 2014) and, likely, 876 saturation with respect to reactive minerals. The influence of less reactive 877 minerals will therefore increase, since they will continue to dissolve. In 878 contrast, fresh melt in summer will only take up to 4 days to reach the outlet of 879 the Leverett Glacier from inland moulins (Chandler et al., 2013). Similar time 880 spans will also apply for the rock-interaction time of non-glacial streams 881 before they join the large glacial rivers.

The first two dissolved Hf isotope compositions of Leveret River were obtained during the final meltwater pulse event of the 2009 season, during which seven surface lakes drained and expelled long-term stored subglacial meltwater from the bed of the ice sheet (Fig. 2a, Bartholomew et al., 2011).

886 This caused up-glacier expansion of the fast/efficient channelized system at the 887 expense of a slow/inefficient distributed drainage system. The flushing of basal 888 waters, which have interacted with the subglacial sediment for an extended 889 period of time, is consistent with the enrichment of the bulk runoff in solutes 890 and elevated electrical conductivity, starting with the water pulse and ending 891 at around the 9<sup>th</sup> of July (Fig. 2b, e.g., Cuffey and Paterson, 2010; Bhatia et al., 892 2011; Hindshaw et al.; 2014). Such waters, which do not reflect very recent 893 melt, are referred to as delayed waters (e.g., Bhatia et al., 2011). The 894 unradiogenic Hf isotope composition of the first two samples, therefore, 895 reflects higher proportions of delayed waters compared to later sampling 896 occasions, when recent glacial melt draining to the bed through crevasses, 897 moulins and englacial channels increasingly constituted the water flux of the 898 river. The large contrast between the first two observations, whereby the 899 second seems more influenced by delayed waters than the first, may imply 900 isolated water reservoirs with heterogeneous  $\varepsilon_{Hf}$  compositions beneath the 901 glacier, reservoirs that are accessed gradually by the extending channelized 902 draining system.

903 The clear difference in Hf isotope compositions between glacial and non-904 glacial waters probably also results from different average water-rock 905 interaction times. The channelized drainage system beneath the Leverett 906 Glacier expands inland over the course of the melting season (Bartholomew et 907 al., 2010; Chandler et al., 2013), which implies a continuous source of delayed 908 waters (e.g., Bhatia et al., 2011; Chandler et al., 2013). In contrast, the Hf 909 isotopic composition of non-glacial streams results from the short timescale 910 interaction of these waters with the rock and regolith substrate, as they flow in 911 the shallow zone above the permafrost horizon. More radiogenic Hf isotope 912 compositions in non-glacial compared to glacial waters are thus an expression 913 of their short water-rock interaction time, whereby the released Hf isotope 914 signal is more influenced by weathering of reactive and radiogenic minerals, 915 especially garnet. Glacial waters, on the other hand, are more controlled by 916 weathering of unradiogenic and less reactive minerals, in particular zircon (Fig. 917 6d). The strong contrast in Hf concentrations of supraglacial melt (0.06 to 0.2 918 pmol/l) and the Leverett River (13.9 to 42.7 pmol/l; Table 2) also suggests that

the Hf isotope composition of delayed waters will exert a strong control on thebulk isotope composition of the river.

921

# 922 **5.6.2. Regolith exposure age**

923 It could be argued that the availability of freshly ground and strained mineral 924 surfaces, produced continuously through the active sliding of warm-based 925 glaciers (Cuffey and Paterson, 2010), results in a bias towards unradiogenic Hf 926 from zircons (Piotrowski et al., 2000; van de Flierdt et al., 2002). Non-glacial 927 streams, on the other hand, would potentially lack this source as strained 928 surfaces anneal over time in the proglacial area. Two arguments can be made 929 against this interpretation. In a detailed study of Sr isotopes at Leverett Glacier 930 in July 2009, Hindshaw et al. (2014) documented a clear change in Sr isotope 931 composition at the transition from a distributed to a channelized subglacial 932 drainage system. Although a mobile element like Sr, which is abundant in a 933 range of minerals with variable weathering susceptibility, could be more prone 934 to respond to water-rock interaction time, it seems unlikely that the underlying 935 processes for Sr and Hf isotopes are completely different. In addition, the lake 936 sample measured in this study is most likely strongly influenced by weathering 937 of long-exposed proglacial material, as it is situated west of the Ørkendalen 938 moraines, which are 6.8 kyrs old (Levy et al., 2012). Nevertheless, the lake yields a Hf isotope composition similar to glacial samples (Fig. 6d), probably as 939 940 a result of the long residence time of water in the lake (c.f., Anderson et al., 941 2001). Although, high Cl<sup>-</sup> concentrations in streams indicate that non-glacial 942 streams are fed by lakes, their Hf isotope composition is much more 943 radiogenic. This probably reflects removal of Hf from lakes, possibly through 944 adsorption onto diatoms (Stichel et al., 2012a) and an overprint of Hf isotope 945 signatures in streams. In support of this interpretation, Si concentrations are 946 very low in the lake (Table 1) and diatoms constitute some of the deposited 947 sediment (Willemse, 2002).

948

## 949 **5.7 Implications**

The new observations clearly show that glacial weathering processes increasethe congruency in Hf isotope weathering compared to the glacial forefield. Such

952 a direct influence of ice-sheets on the release of Hf isotopes was previously 953 only inferred indirectly from observations in marine ferromanganese 954 precipitates (Gutjahr et al., 2014; Piotrowski et al., 2000; van de Flierdt et al., 955 2002). The more congruent Hf signals in glacially sourced waters compared to 956 the non-glacial streams support the interpretation that more crustal-like  $\varepsilon_{Hf}$  in 957 the North Western Atlantic since  $\sim 3$  Ma relates to the onset of northern 958 hemispheric glaciation (van de Flierdt et al., 2002). The glacial rivers studied 959 here are, however, not distinctly unradiogenic in Hf in general terms, since 960 they fall close to the seawater array in Hf-Nd isotope space (Fig. 4b).

961 For the studied area, extended water-rock interaction time in subglaical 962 settings is likely more relevant for the release of unradiogenic Hf than the 963 effect of glacial grinding (section 5.6). A strong influence of strained zircon 964 surfaces on dissolved Hf isotopes also seems inconsistent with observations on 965 a longer time scale (Gutjahr et al., 2014). If the availability of freshly ground 966 rock substrate was crucial for the release of unradiogenic Hf, it would probably 967 lead to a co-evolution of  $\varepsilon_{Hf}$  and Pb isotopes in seawater during the deglacial 968 since both isotope systems would be governed by the exposure and weathering 969 of glacial material and a short oceanic residence time (20 to 30 yr for Pb in the 970 Atlantic, Henderson and Maier-Reimer, 2002). While efficient zircon 971 weathering would gradually decrease in the newly forming glacial forefields -972 reflecting zircon annealing - weathering of newly exposed allanite, apatite and 973 titanite with radiogenic Pb and Hf isotopes (Harlavan et al., 1998; Harlavan and 974 Erel, 2002; Bayon et al., 2006), would be expected to intensify. Recent 975 observations indicate, however, a  $\sim 5$  kyr earlier change towards more 976 radiogenic Hf in the Northwest Atlantic after the last glacial maximum than is 977 observed for Pb isotopes (Gutjahr et al., 2014). Although seawater Pb isotope 978 evolution is governed by the exposure of radiogenic minerals in Pb in glacial 979 forefields (Harlavan et al., 1998; Foster and Vance, 2006), Hf isotopes appear to 980 be mostly responsive to changes in subglacial hydrology (this study, Gutjahr et 981 al., 2014). This does not preclude the release of highly radiogenic Hf during the weathering of glacially produced sediments: The non-glacial streams studied 982 983 here are very radiogenic, which supports the notion that glacially sourced 984 sediments could be the cause for radiogenic Hf isotopes in the Kalix River985 (Chen et al., 2013b).

986 In general, currently available dissolved Hf and Nd isotopes of rivers do not reproduce seawater compositions, which are characterized by relatively 987 988 homogenous  $\varepsilon_{Hf}$  in the open ocean and a relatively systematic relationship with 989 ε<sub>Nd</sub> (Godfrey et al., 2009; Rickli et al., 2009; 2010; 2014; Zimmermann 2009a; 990 Stichel et al., 2012a,b). Larger variability in  $\varepsilon_{Hf}$  is observed in semi-enclosed 991 basins like the Baltic and the Labrador Sea (Chen et al., 2013b; Filippova et al., 992 2017). Riverine Hf isotope compositions are, in contrast to the open ocean, 993 highly variable at a given Nd isotope composition (c.f., this study; Bayon et al., 994 2006; Godfrey et al., 2007; Zimmermann et al., 2009b; Chen et al., 2013b; Rickli 995 et al., 2013). The discordance between seawater Hf-Nd isotope compositions 996 and rivers is unexpected given recent estimates of the seawater residence time 997 of Hf (Chen et al., 2013b; Filippova et al., 2017), which is most likely shorter 998 than that of Nd (< 500 yr, Siddall et al., 2008). This could suggest that a further 999 significant source of Hf in seawater is more homogeneous than rivers, possibly 1000 representing an exchange process at the ocean boundary as observed for Nd 1001 (Lacan and Jeandel, 2005). Clay sized sediments are indeed rather homogenous 1002 in  $\varepsilon_{Hf}$  (Bayon et al., 2016) implying that the release of Hf from fine sediments 1003 deposited in the ocean could be the sought-after source (Albarède et al., 1998). 1004 Alternatively, unradiogenic riverine fluxes in Hf may be balanced by radiogenic 1005 fluxes on a larger scale, leading to overall homogenous riverine inputs to the 1006 oceans. If that turns out to be the case, it would justify models that interpret 1007 the seawater array as a result of a consistent partitioning of Hf between zircon 1008 and other minerals in the upper continental crust (van de Flierdt et al., 2007; 1009 Chen et al., 2011) and a constant effect of incongruent weathering for the 1010 zircon free crust on large scales (Chen et al., 2011). It would also support the 1011 idea, that the "clay array" mostly reflects weathering released Hf (Bayon et al., 1012 2016), without any significant contributions of Hf from primary minerals 1013 including zircon.

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- 1015

### 1016 **6. Conclusions**

1017 This study represents the first comprehensive attempt to understand the 1018 hydrological and mineralogical factors that affect and control the release of Hf 1019 isotopes into the hydrosphere in a high latitude setting characterized by glacial 1020 activity. To this end, we have measured Hf and Nd isotopes and concentrations 1021 in catchment rocks, corresponding mineral separates, catchment sediments, 1022 riverine suspended load as well as glacially fed rivers and non-glacial streams. 1023 These analyses are complemented by the mineralogical characterization of 1024 sediment samples from the catchment, providing an estimate of the mineralogy 1025 of weathered material.

1026 Hafnium and neodymium isotope compositions of catchment sediments and 1027 riverine suspended load are well reconciled with the measured Hf and Nd 1028 isotope compositions and concentration in bulk rocks and mineral separates. 1029 Specifically, their Nd isotope composition is consistent with a typical gneissic 1030 composition in the catchment, slightly enriched in titanite and including 1031 amphibolite derived hornblende. The Hf isotopic range in sediments 1032 additionally requires variable proportions of highly radiogenic garnets and 1033 variable trace amounts of unradiogenic zircon.

Dissolved Hf and Nd isotopes in waters show very different characteristics. Neodymium isotopes closely mirror the isotope composition of representative gneisses in the catchment ( $\varepsilon_{Nd} = -39$ ) and cover a small range in all glacially fed and non-glacial waters ( $\varepsilon_{Nd} = -42.8$  to -37.9). Dissolved Hf isotopes, on the other hand, are shifted to much more radiogenic values than bulk gneisses ( $\varepsilon_{Hf}$ = -55.1) yielding values close to the seawater array for glacial rivers ( $\varepsilon_{Hf} = -18.3$ to -0.9) and between +15.8 and +46.3 for non-glacial streams.

1041 Two hypotheses can explain the strong contrast in dissolved Hf isotopes in 1042 glacial and non-glacial waters. First, it could imply a larger influence of zircon 1043 weathering in glacial rivers due to the availability of glacially strained zircon 1044 surfaces. Second, it could reflect different average water-rock interaction times 1045 and, as a result, different contributions from reactive/radiogenic versus 1046 weathering resistant/unradiogenic minerals. Glacial rivers receive continuous 1047 contributions from long interacting waters of distributed subglacial drainage 1048 systems, whereas non-glacial streams are characterized by fast superficial 1049 drainage above the permafrost horizon. The implied short interaction time

1050 increases the significance of garnet over zircon weathering, mostly as a result 1051 of the chemical inertness of zircons relative to garnet. Although it may be 1052 premature to draw a firm conclusion as to which process is more dominant, the 1053 current observations favour water-rock interaction time. The sampled lake, for 1054 instance, is chemically closely related to non-glacial streams, yet shows a Hf 1055 isotope signature in the range of glacial rivers, probably due to the long water 1056 residence time in the lake. Similarly, increasingly warmer ice sheets produced 1057 less congruent weathering in Hf isotopes in the early stages of the last 1058 deglaciation (Gutjahr et al., 2014), probably due to more efficient subglacial 1059 drainage and a shortening of water-rock interaction times.

1060

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1374Table 1: Chemical and physical characteristics, as well as Nd and Hf isotopic1375compositions, of the sampled rivers, streams and the lake. pH, T, Ca, Mg, Na, K,1376Si,  $HCO3^-$ ,  $SO4^{2^-}$ , Cl<sup>-</sup> and TSS have been compiled from the literature, where the1377exact coordinates of sampling locations can also be found (Hindshaw et al.,13782014; Wimpenny et al., 2010; 2011). External errors on isotope ratios are1379discussed in section 3.4.

1380

Table 2: Elemental concentrations (Hf, Sm, Nd) and isotope compositions (Hf,
Nd) of bulk rocks, minerals, catchment sediments and riverine suspended load.
Hafnium concentrations for bulk rocks and catchment sediments are reported
for hotplate digests (HP) and fused powders (see section 3.3). External errors
on isotope ratios are discussed in section 3.4.

1386

Table 3: Mineral abundances in rocks and sediments. Compositions of rocks
Ro1 - Ro3 and Sed 3 are from Hindshaw et al. (2014). Ro4 has been reanalysed
(see section 4.3).

1390

Table 4: Ages for the four studied rocks based on the Sm/Nd isotope system.Ages were calculated using Isoplot (Ludwig, 2012).

1393

1394 Figure 1: (a) Water sampling sites and geology underlying the Kangerlussuag 1395 region of Southwest Greenland, also including moraines (Levy et al., 2012 and 1396 references therein). The inlet shows the study area within Greenland. (b) 1397 Detailed map of the sampling locations on Leverett Glacier. (X) denotes the 1398 Lake, (Y) the sampled supraglaical stream. Catchment sediments are from the 1399 Lake (X), the sampling location on Leverett River (1) and to the north of it (2), 1400 a proglacial lake (3), the end moraine (4), dirt cones on Leverett Glacier (5), a 1401 proglacial stream (6) and northern and southern lateral moraines (7, 8). 1402 Topography and geomorphology are based on Scholz and Baumann (1997). 1403 1404 Figure 2: Leverett time series of (a) discharge, (b) electrical conductivity (EC),

1404 Figure 2. Leverett time series of (a) discharge, (b) electrical conductivity (EC)

1405 (c) Hf and Nd concentrations and (e, f) isotope compositions in July 2009.

1406 Discharge and conductivity data are from Bartholomew et al. (2011).

1407

Figure 3: Composition of sampled waters in terms of major cations (a) andanions (b) in terms of molar abundances. (Data sources are given in the captionfor Table 1).

1411

Figure 4: Neodymium isotope composition vs molar Sm/Nd ratios for all water
and solid samples (a). Hafnium vs neodymium isotope compositions are shown
in (b) in the context of the terrestrial and the seawater array, including the
defining data (small grey symbols, Albarède et al., 1998; David et al., 2001;
Vervoort et al., 2011). Light blue symbols refer to the Leverett time series
samples, dark blue ones to further glacial samples collected elsewhere (see
Table 1).

1419

Figure 5: Hafnium and neodymium concentrations in bulk rocks, mineral separates, catchment sediments and riverine suspended load (a, b), along with molar ratios of Sm/Nd and Hf/Nd for these and water samples (c, d). Grey circles show results for laser analysis of single mineral grains from a variety of catchment sediments (Appendix S1). Hafnium bulk sample concentrations of fused powders are not shown. The feldspar separate from Ro4 represents a mixture of K-feldspar and plagioclase.

1427

1428 Figure 6: Mixing models to evaluate key minerals affecting the Hf and Nd 1429 isotope compositions observed in the riverine dissolved load, suspended load 1430 and in catchment sediments. All mixing calculations use average compositions 1431  $(Sm/Nd, \epsilon_{Nd}, \epsilon_{Hf})$  for titanite, hornblende derived from amphibolites and 1432 garnets. (a) Sm/Nd systematics in catchment sediments, suspended load and 1433 river waters. Neodymium isotopes in catchment sediments are consistent with 1434 the addition of 0.25 wt.% titanite and 10 wt.% hornblende to Ro4. The Sm-Nd 1435 systematics of the suspended load is also consistent with the addition of such a 1436 titanite-hornblende mixture, though at a smaller contribution (55%) relative to 1437 the catchment sediments . Also shown is the individual impact of each mineral 1438 added to Ro4. (b)  $\varepsilon_{Hf}$  vs  $\varepsilon_{Nd}$  for suspended load and catchment sediments. The 1439 variation in  $\epsilon_{Hf}$  is in agreement with variable amounts of garnet and zircon over

1440 and above the variations resulting from titanite and hornblende (see panel a). 1441 Implied garnet abundances are consistent with observations in catchment 1442 sediments, while zircon abundances are constrained by observed Hf 1443 concentrations in catchment sediments and suspended load. (c)  $\varepsilon_{Hf}$  vs 1/Hf for 1444 all sediments and the fitted relationships for the mixtures illustrated in b. 1445 Modelled sedimentary compositions are in good agreement with the 1446 observations, for which a linear regression is shown (grey line,  $r^2=0.82$ , p 1447 <0.001). Larger variations in  $\varepsilon_{Hf}$  would result for catchment sediments if their 1448 Hf isotope compositions and concentrations only reflected variable zircon 1449 abundances at a constant garnet abundance (grey dashed line). (d) Mixing models for dissolved Hf. Mixtures of different minerals with the accessory 1450 1451 mineral free Ro4 illustrate their significance for dissolved Hf isotopes. Adding 1452 titanite, hornblende and apatite to the accessory mineral free Ro4 defines the 1453 black solid trajectory, which, complemented by garnet and zircon, reproduces 1454 Hf in glacial and non-glacial waters (black arrows starting from the trajectory 1455 at an Ro4-like  $\varepsilon_{Nd}$ , black circle). Also shown is a mixture between the measured 1456 Ro4 and garnet (grey dashed line). Less radiogenic  $\varepsilon_{Nd}$  in some proglacial rivers 1457 and the lake likely reflect higher weathering contributions from allanite and 1458 monazite (section 5.4).

1459

1460 Figure 7: Surface-area normalised dissolution rates for minerals in the 1461 catchment. The rates are fitted to laboratory observations at 25°C far from 1462 mineral saturation (Guidry and Mackenzie, 2003; Palandri and Kharaka, 2004 1463 and references therein). Hornblende data is from Sverdrup et al. (1990, green fitted line), Golubev et al. (2005, green circles) and Frogner and Schweda 1464 1465 (1998, green diamonds). Apatite dissolution rates are nearly constant in a pH-1466 range from 6 to 10 (Chaïrat et al., 2007), so that the fit given here by Guidry 1467 and Mackenzie (2003) can be extended to the pH range of the figure. 1468 Dissolution rates of oligoclase under basic conditions are not available and 1469 albite rates are plotted instead. The shaded area depicts the pH range of 1470 studied waters.

1471

Sample	Туре	Date	Time	рН	Т	Ca	Mg	Na	K :	Si	HCO3	SO4 <sup>2-</sup>	Cl	TDS	TSS	Hf	Sm	Nd	εHf	2 SEM	εNd	2 SEM
					°C	µmol/l									g/l	pmol/l				int		int
Regional sampling - 2006																						
GR10	fjord	-		- 6.96	10.5	615.0	2940.0	>2000	586.0	26.6	266.0	) 1890.0	42300.0	48623.6			-	-	-		-40.82	0.15
GR11	non-glacial	15.7.06		- 6.81	10.0	105.0	86.3	99.1	15.6	76.1	358.0	) 17.2	94.0	851.3	0.004	Ļ	-	-	- +17.	9 0.3	-38.73	0.07
GR12	non-glacial	14.7.06		- 8.30	19.3	665.0	775.0	539.0	142.0	323.0	1740.0	) 263.0	596.0	5043.0	0.014	Ļ	-	-	- +15.	8 0.2	-42.80	0.07
GR13	non-glacial	15.7.06		- 7.65	14.2	148.0	116.0	136.0	40.5	6.9	532.0	) 22.0	105.0	1106.4			-	-	- +46.	3 0.5	-39.61	0.08
GR14 <sup>a</sup>	non-glacial	18.7.06		- 8.10	7.5	229.0	91.1	362.0	11.2	59.5	556.0	) 72.0	324.0	1704.8			-	-	- +13.	5 0.7	-30.40	0.07
GR15	non-glacial	16.7.06		- 8.25	12.3	181.0	202.0	171.0	82.7	4.2	784.0	28.4	119.0	1572.3			-	-	- +43.	3 1.4	-40.72	0.11
GR1	glacial	11.7.06		- 8.48	4.2	24.8	8.3	11.6	12.2	19.6	72.8	8 8.3	2.0	159.5	0.44	Ļ	-	-	7.	2 0.3	-39.22	0.08
GR2	glacial	11.7.06		- 7.91	4.2	32.9	11.0	29.9	18.2	21.6	98.5	5 13.4	6.0	231.5	0.60	)	-	-	12.	2 0.2	-39.30	0.07
GR4	glacial	12.7.06		- 7.11	6.0	40.8	11.6	32.2	20.2	25.7	119.0	) 19.5	4.3	273.3	0.85		-	-	13.	7 0.3	-39.18	0.12
GR5	glacial	13.7.06		- 7.18	3.0	39.2	11.4	30.2	18.6	24.8	145.0	) 16.8	5.2	291.2	0.40	)	-	-	13.	6 0.2	-39.07	0.10
GR7	glacial	13.7.06		- 7.30	8.6	42.4	13.7	35.4	21.6	28.7	133.0	) 19.3	4.1	298.2	0.55		-	-	14.	7 0.8	-38.97	0.12
GR8	glacial	14.7.06		- 7.20	5.5	40.6	14.4	37.7	22.9	42.1	151.0	) 17.9	3.2	329.8	0.44	Ļ	-	-	0.	9 0.6	-39.51	0.09
GR9	glacial	14.7.06		- 7.18	5.3	31.1	9.7	8.0	9.8	21.7	99.3	3 10.7	-	190.3			-	-	6.	6 0.4	-37.95	0.07
Leverett time series - 2009	1																					
Lev6	glacial	6.7.09	09:1	0 9.34	0.1	56.0	13.7	89.9	48.4	36.8	185.3	3 29.5	6.6	466.2	7.4	42.	7 689.	9 4984.	5 -13.	7 0.2	-38.62	0.08
Lev8	glacial	8.7.09	17:4	5 8.02	0.2	26.6	7.1	35.9	25.2	17.2	99.3	3 11.2	4.1	226.7	3.3	24.	4 457.	8 3310.	5 -18.	3 0.2	-38.35	0.07
Lev10	glacial	10.7.09	08:3	0 8.22	0.2	31.9	8.0	42.4	29.0	20.3	128.4	15.2	4.8	280.1	2.9	30.	5 564.	8 4065.	9 -9.	1 0.1	-38.49	0.09
Lev12	glacial	12.7.09	18:1	0 7.92	0.3	26.5	6.6	30.6	23.4	15.3	110.2	2 11.5	3.4	227.5	2.3	19.	2 357.	0 2585.	3 -11.4	4 0.2	-38.63	0.06
Lev14	glacial	14.7.09	08:0	0 8.21	0.2	30.9	8.3	35.2	23.8	20.2	113.2	2 14.0	3.2	248.7	3.2	15.	8 278.	7 2017.	1 -9.	9 0.4	-38.41	0.07
Lev18	glacial	18.7.09	08:1	0 8.25	0.1	36.2	8.5	39.0	25.5	20.9	104.6	5 15.6	4.4	254.8	3.0	19.0	376.	8 2651.	3 -11.	2 0.3	-38.51	0.07
Lev20	glacial	20.7.09	17:2	0 7.81	0.1	30.4	7.6	33.6	21.9	17.0	100.5	5 13.1	3.2	227.3		· 17.	4 370.	6 2666.	2 -12.4	4 0.2	-38.54	0.09
Lev22	glacial	22.7.09	08:1	5 8.08	0.1	35.7	8.4	37.6	24.6	19.1	109.1	17.1	4.1	255.7	3.3	20.	7 363.	4 2611.	0 -12.	3 0.2	-38.29	0.08
Lev24	glacial	24.7.09	17:5	0 7.88	0.1	33.5	8.7	36.7	23.9	17.9	99.9	9 15.4	4.4	240.4		- 17.	3 329.	1 2386.	5 -11.	8 0.3	-38.43	0.08
Lev26	glacial	26.7.09	08:1	0 8.08	0.1	35.9	8.3	37.3	23.6	19.9	110.7	/ 16.4	4.9	257.0	4.2	16.	7 319.	7 2305.	7 -11.	7 0.2	-38.52	0.07
Lev28	glacial	28.7.09	18:0	0 8.28	0.1	39.4	8.7	38.9	24.7	20.6	126.2	2 17.8	3.2	279.6	3.6	13.	9 266.	3 1909.	2 -11.	5 0.4	-38.40	0.09
CampLake	lake	17.7.09	16:0	0 8.93	14.0	834.0	1134.5	1008.5	400.1	0.0	3767.0	) 30.0	1099.0	8273.1		- 5.	5 92.	7 644.	1 -15.	7 0.5	-42.32	0.09
SGlacial23 <sup>b</sup>	supraglacial	23.7.09	14:0	0 5.31	0.1	0.0	0.0	0.4	0.1	0.1	NA	0.0	0.2	0.8		0.0	52.	0 12.	1		-32.40	0.20
SGlacial29 <sup>b</sup>	supraglacial	29.7.09	16:0	0												0.2	3.	1 16.	4		-34.56	0.11

<sup>a</sup> Sample from close to Sisimut

<sup>b</sup> Chemical composition reflects the average of two glacial samples from Hindshaw et al. (2014)

Table 1

	Hf	Hf	Sm	Nd	εHf	2 SEM	εNd	2 SEM
	ppm (HP)	(fused)				int		int
		( /				-		
Ro1 - Garnet Amphibolite	1.40	2.71	4.48	16.11	+26.4	0.2	-6.71	0.09
Garnet	0.63		1.39	4.04	+1308.3	0.3	+2.78	0.09
Hornblende	2.02		7.05	22.65	+1.6	0.3	-1.83	0.13
Plagioclase	0.28		0.39	1 27	-27.1	0.5	-2 71	0.13
i lagiociase	0.20		0.55	1.27	27.1	0.0	2.7 1	0.15
Ro3 - Amnhiholite	0.72	0.95	1 60	5 10	+273	04	+1 00	0.07
Clinopyroyene	0.72	0.55	0.41	1 1 2	- 27.5	0.4	+8 20	0.07
Enidoto	1.06		0.41	20.25	,, 177 0	0.4	+2 10	0.13
Horphondo	1.00		1 10	29.25	+122.0	0.5	+2.13	0.11
Plagioslasa	0.95		1.19	2.79	+0.0	0.4	+10.94	0.11
Plagiociase	0.05		0.25	0.07	+05.2	0.5	+0.22	0.20
Rutile	4.48		2.41	5.39	+39.2	0.5	+19.06	0.17
De 2. Orthe an eige	2.65	4.00	1.05	10 51	50.0	0.2	24.44	0.07
Roz - Orthogneiss	3.65	4.90	1.85	10.51	-59.9	0.2	-31.44	0.07
Chlorite	3.21		4.78	38.82	-59.8	0.3	-42.78	0.26
Epidote	2.18		15.59	99.42	-25.2	0.3	-30.44	0.07
K-feldspar	1.46		0.38	2.59	-61.4	0.1	-35.99	0.09
Plagioclase	0.66		0.05	0.24	-60.6	0.2	-29.82	0.16
Titanite	46.40		330.48	772.31	+20.9	0.1	+4.43	0.09
Zircon	10827.61		47.04	74.04	-61.4	0.1	-19.74	0.10
Ro4 - Orthogneiss	1.63	3.86	3.86	27.30	-55.1	0.2	-39.04	0.22
Chlorite	0.50		0.51	1.86	-50.3	0.7	-16.42	0.14
Epidote	1.73		14.17	34.53	-18.1	0.2	+2.11	0.14
Feldspar	0.08		0.04	0.22	-58.1	0.6	-33.03	0.15
Hornblende	2.23		8.67	33.35	-21.5	0.2	-17.63	0.12
Titanite	16.47		397.73	1109.51	+116.6	0.2	-5.34	0.13
Zircon	9130.48		5.88	18.90	-61.2	0.1	-32.13	0.20
Archean granite	0.97		2.01	19.99	-58.43	0.2	-47.33	0.06
Bulk catchment sediments <sup>a</sup>								
Sed 1 - river sediment	10.42	20.85	7.13	40.29	-53.1	0.1	-34.01	0.06
Sed 2 - river sediment	7.51	12.76	4.52	25.45	-54.4	0.1	-33.29	0.06
IC sand - river sand	2.59	5.09	2.96	17.06	-51.0	0.1	-33.79	0.08
SM Sed - side moraine	2.75	6.25	3.49	20.34	-47.0	0.1	-33.86	0.07
PGI 1 - proglacial lake	3 95	5 55	3 16	18 29	-52.4	0.2	-34 24	0.09
MH1 - dirt cone sediment	2 17	3 10	2 43	16.23	-53.0	0.2	-38.28	0.05
MH2 - dirt cone sediment	5.80	6 38	2.45	18 / 7	-53.0	0.2	-33 90	0.07
	5.00	0.50	5.21	10.47	55.2	0.1	55.50	0.00
Garnet PGL 2								
Garnet a	0.49		2 11	5 20	<b>⊥</b> 971 0	0.4	±7 //2	0.08
Carnet h	0.49		2.44	5.50	10/1.5	0.4	+7.42	0.08
Garnet o	0.40		2.70	0.00	+1247.4	0.4	+5.05	0.08
Gamet d	0.35		3.54	8.27	+1308.1	0.5	+3.72	0.06
Garnet d	0.26		2.90	5.73	+1940.2	0.5	+15.06	0.07
Anatita Sad 2 DCL 2	0.72		102.00		11100	<b>n</b> n	16 22	0.00
Apathe Sea 3, PGL 2	0.73		192.98	030.05	+1148.9	2.3	-10.33	0.09
Buik riverine suspension								0.07
Lev14	2.04		4.68	30.03	-43.3	0.3	-36.05	0.07
Lev18	2.16		4.65	29.36	-	-	-35.86	0.09
Lev20	2.28		4.86	30.67	-45.3	0.6	-35.78	0.08
Lev22	2.30		4.67	29.50	-45.3	0.6	-35.84	0.07
Lev24	2.72		4.77	30.00	-46.3	0.6	-35.81	0.08
Lev26	2.61		4.98	31.42	-46.4	0.5	-35.80	0.07
Lev28	1.92		4.63	29.33	-44.5	0.8	-35.96	0.09
GR1	1.39		5.26	33.89	-35.8	0.5	-35.74	0.11
GR2	1.59		5.15	32.27	-39.9	0.6	-35.85	0.07
GR4	1.42		4.68	29.90	-40.5	0.5	-36.81	0.09
GR5	1.46		4.65	30.01	-34.6	0.3	-36.83	0.07
GR8	1.26		5.16	33.31	-39.8	0.3	-36.43	0.09

<sup>a</sup> Some of the sediment labels in Table 2 of Hindshaw et al. (2014) were confounded. The correct labels for LC Sand, SM Sed, MH1 and PGL1 are SM Sed, PGL1, LC Sand and MH1. These misrepresentations have no effect on the discussions and conclusions in Hindshaw et al. (2014).

	vol. %							wt.%
	Ro1	Ro3	Ro2	Ro4	Sed 3	Soil 1	PGL 2	in sediments
Major minerals								
Clinopyroxene		22			<1	<1	2.2	<2.8
Garnet	4				1.5	2.6	2.6	2.1-3.7
Hornblende	68	45		<1	5.9	8.6	9.5	7.2-11.7
Scapolite		12			-	-	-	-
K-feldspar			24	44	8.3	16.9	7.5	7.3-16.3
Plagioclase	14	18	50	21	42.0	40.4	45.5	38.8-44.1
Quartz	10		22	27	33.8	37.6	34.3	33-36.8
Accessory minerals								
Apatite			1	~ 0.15	<1	<1	<1	<1.2
Biotite					<1	1.3	1.2	<1.4
Chlorite			2	6	-	-	-	-
Epidote		3	<1	2	<1	-	<1	<1.3
Ilmenite			1		<1	<1	1.0	<1.7
Orthopyroxene					<1	<1	<1	<1.2
Magnetite	1				-	<1	<1	<1.9
Titanite	3	<1	<1	~ 0.13	<1	-	<1	<1.3
Zircon			<1	~ 0.02	<1	<1	<1	<1.7

Table 3

	Age (Myr)	Initial 143Nd/144Nd	Uncertainty (2σ)	MSWD
Ro1 - Garnet Amphibolite	1855 ± 87	0.51025	0.00011	2.7
Ro3 - Amphibolite	1746 ± 200	0.51053	0.00028	140
Ro2 - Orthogneiss	1776 ± 320	0.50984	0.00030	2436
Ro4 - Orthogneiss	1887 ± 230	0.50970	0.00028	384

Table 4





Fig. 2



Fig. 3





Fig. 5





Fig. 7