# Constraints on the source of reactive phases in sediment from a major Arctic river using neodymium isotopes

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#### Abstract :

Riverine suspended particulate matter (SPM) is essential for the delivery of micronutrients such as iron (Fe) to the oceans. SPM is known to consist of multiple phases with differing reactivity, but their role in the delivery of elements to the oceans is poorly constrained. Here we provide new constraints on the source and composition of reactive phases in SPM from the Mackenzie River, the largest sediment source to the Arctic Ocean. Sequential leaching of SPM shows that river sediments contain labile Fe phases. We estimate the labile Fe flux is substantial (0.21(+0.06,-0.05) Tg/yr) by quantifying Fe concentrations in weak leaches of the SPM. The labile Fe phase hosts a considerable amount of rare earth elements (REE), including neodymium (Nd). We demonstrate that the labile Fe phase and dissolved load have radiogenic Nd isotope ratios that are identical within uncertainty, but up to 8 epsilon units distinct from the silicate phase. We interpret this as evidence for dynamic cycling between Fe-oxide phases in SPM and the river water, demonstrating the high reactivity of the labile Fe phase. Nd isotope and elemental molar ratios suggest that a significant amount of labile Fe- and Nd-bearing phases are derived from Fe-oxides within the sedimentary source rock rather than silicate mineral dissolution. Thus, sedimentary rock erosion and weathering provides an important source of labile Fe, manganese (Mn) and by extension potentially other trace metals. Our results imply that both past and future environmental change in the Arctic, such as permafrost thaw, may trigger changes to the supply of reactive trace metals. These results demonstrate that a re-evaluation of sediment reactivity within rivers is required where uplifted sedimentary rocks are present.

#### Highlights

► Assessment of reactive phases in suspended particulates in a major Arctic river. ► Neodymium isotopes show equilibration between Fe-Mn oxides and dissolved loads. ► Reactive iron flux from the Mackenzie River is large; 0.21(+0.06,-0.05) Tg/yr. ► Sedimentary rock weathering provides an important source of labile trace metals.

Keywords : iron oxides, bioavailable, neodymium isotopes, Arctic

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# 1. Introduction

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54 Rivers are conduits for the transfer of particulate and dissolved material derived from weathering to the oceans (Gaillardet et al., 1999). Weathering plays a critical role in 55 56 the cycling of elements at the Earth's surface and is a key regulator of the Earth's 57 climate over geological timescales (Walker et al., 1981). Rivers also transport 58 essential nutrients to the oceans sustaining marine primary productivity (e.g., 59 Deutsch and Weber, 2012). Changes to the flux of nutrients that limit productivity in 60 the oceans have been linked to large scale perturbations in the carbon cycle and 61 past climate change (Martin, 1990; Vincent and Berger, 1985). The flux of 62 suspended particulate matter (SPM) sustains the supply of certain elements to the 63 oceans and plays an important role in bio-geochemical cycling (Jeandel and Oelkers, 64 2015; Jones et al., 2012).

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66 Micronutrients (e.g., Fe), the rare earth elements (REE, particularly Nd) and many 67 other trace elements have low solubility in natural waters. In the dissolved load (<0.2 µm) these elements are predominately associated with nanoparticulate and colloidal 68 69 phases (Gaillardet et al., 2014, and references therein). Their concentrations in river 70 dissolved loads are low (ppb to ppt, Gaillardet et al., 2014); concentrations in SPM 71 are several orders of magnitude higher (Jeandel and Oelkers, 2015). Salt-induced 72 flocculation of many such nonmobile elements in estuaries further removes most of 73 these low concentration elements prior to reaching the oceans (Elderfield et al., 74 1990).

76 Basalts and volcanic particulates are important sources of key bio-limiting nutrients 77 (e.g. Fe) as they are easily weatherable and enriched in those elements (Jones et al., 2012). However, the importance of sedimentary rocks has not been similarly 78 79 evaluated. Marine sediments contain phases, including Fe-Mn oxyhydroxides with 80 elevated REE concentrations, which are formed from precipitation and exchange 81 with seawater and pore fluids (e.g., Chester and Hughes, 1967). Marine sedimentary 82 rocks tectonically emplaced on the continents retain some of these characteristics 83 (Hindshaw et al., 2018; Jang et al., 2020). This results in a reservoir of inherited 84 phases which can be more reactive than silicate minerals and may supply a greater 85 amount of bioavailable Fe to riverine sediments (Hindshaw et al., 2018). For 86 instance, amorphous Fe (oxy)hydroxides (e.g. ferrihyrite) contained within riverine 87 sediments are easily reducible and potentially bioavailable (Bhatia et al., 2013, 88 Hawkings et al., 2018).

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90 Since Fe-oxides are enriched in REEs, their origin can be traced using radiogenic Nd 91 isotopes ( $^{143}Nd/^{144}Nd$ , expressed as  $\epsilon Nd$ ), a source tracer in riverine and marine 92 sediments (Goldstein and Jacobsen, 1988; Goldstein and Hemming, 2003). Since 93 there is limited fractionation of the Sm/Nd ratio in particulate phases during 94 weathering and the decay constant is long compared to the timescales of recent 95 weathering (Babechuk et al., 2014), ɛNd is only controlled by changes in source. Nd in the dissolved load typically has  $\varepsilon$ Nd greater (more radiogenic) than in the SPM in 96 97 rivers with abundant sedimentary rocks in their catchments (Goldstein and 98 Jacobsen, 1987; Hindshaw et al., 2018). This offset is attributed to dissolved Nd 99 sourced from the preferential weathering of marine precipitates and other reactive 100 components contained within the parent rock (Goldstein and Jacobsen, 1987).

101 Nonetheless, a clear link between these reactive phases and dissolved riverine102 chemistry has not been shown.

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104 With knowledge of the source composition, ɛNd is a valuable tool for tracing the 105 origin of reactive phases from sedimentary rock weathering and their transport in 106 river systems. We apply this tracer to a large Arctic river system. The Mackenzie 107 River, as the first and third largest source of suspended sediment and water, 108 respectively, to the Arctic Ocean (Macdonald et al., 1998, Holmes et al., 2002), 109 provides a good representation of basin-scale fluxes. It is dominated by shale 110 weathering with a low abundance of volcanic or mafic rocks and limited influence 111 from the weathering of crystalline rocks (Millot et al., 2003, Fig. 1B). This makes the 112 Mackenzie an ideal location to investigate the mobilisation of reactive, micronutrient-113 bearing phases during shale weathering on a large scale. 114 115 In this study we characterize the ɛNd of the dissolved load (<0.2 µm and

ultrafiltrates) and suspended sediment (sequential extractions and residue) in the
Mackenzie and key tributaries over two years (2017, 2018) sampled within a few
days of the peak discharge after ice-break up. We show that the εNd is within
uncertainty between the dissolved load and easily leachable phases in the SPM. We
use REE patterns to constrain the mineralogy of the leachable phases. We quantify
the concentrations of major and some trace elements in leachable phases and
estimate the flux of labile Fe from the Mackenzie to the Arctic Ocean.

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124 **2. Materials and Methods** 

Full analytical and procedural methods are given in Supplementary Text 1 and
summarised briefly here. All analysis was carried out at the University of Cambridge,
Department of Earth Sciences.

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130 **2.1 Study area and sample collection** 

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The Mackenzie River system is large (Fig. 1, area 1.78x10<sup>6</sup> km<sup>2</sup>). The weathering of sedimentary rocks dominates the dissolved and SPM loads of the river (Calmels et al., 2007; Horan et al., 2019; Huh et al., 2004; Millot et al., 2003) despite the large surface area of crystalline rocks (29.2% of the basin geology, Fig. 1b). Less than 1% of the basin is underlain by volcanic rocks (Fig. 1b). The river is characterised by a sharp peak in discharge each year after the seasonal melting (freshet) of river ice in May-July (Fig. 2). Samples were collected at this period of maximum flux.

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140 The Mackenzie mainstem and two large tributaries which enter at the delta, the Peel 141 River and the Arctic Red River, were sampled at 3 main sites (Fig. 1) in early June in 142 2017 and 2018. Suspended sediment samples were taken at different depths in the 143 channel using a 5L Van-Dorn type depth sampler (following Hilton et al., 2015). 144 Discharge was quantified using an acoustic-Doppler current profiler (ADCP, Rio 145 Grande II (1200 kHz), Teledyne Instruments, Fig. 2, Supplementary Table S4). 146 Additionally, in 2018 a small Peel River tributary and meltwater from the surface 147 Arctic Ocean (seawater diluted by sea and river ice melt) were sampled. Dredged 148 bed load, bank sand and rock fragments were also collected.

River samples were filtered at 0.2  $\mu$ m, collecting both water and sediment. Filtered waters collected for major cation and REE concentrations were acidified using distilled HNO<sub>3</sub> to pH 2 in acid cleaned bottles. In 2018, ultrafiltration was carried out on <0.2  $\mu$ m filtrates from the Mackenzie (Middle Channel) and the Peel River, using dialysis membranes with molecular cut off weights of 10 kDa and 1 kDa (Supplementary Text 1.2). Approximately 10 L of <0.2  $\mu$ m water was collected for the analysis of Nd isotopes using a Fe co-precipitation method (Hindshaw et al., 2018).

158 To characterize the longer-term integrated composition of the Mackenzie River SPM 159 leachate compositions from 5 offshore sediment cores from the Mackenzie delta and 160 one from deeper in the Beaufort Sea were analysed (Natural Resources Canada, 161 Geology Survey of Canada's Marine Geoscience Collection, Supplementary Table 162 S5). Two silicate residue samples were analysed from two of the shallowest core 163 sites. The sedimentation rate at shallow sites in the Canadian Beaufort sea is high 164 (on the order of 2-3 m/ka, O'Regan et al., 2018), indicating that the range and depth 165 sampled within the cores (~10 cm) represents sediment that is likely several 166 decades old, and homogenises inter-annual and seasonal variability.

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# 168 2.2 Sequential extractions

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Approximately 50-100 mg of dry sediment was leached sequentially to extract: 1)
exchangeable phases (1M NH<sub>4</sub>Cl), 2) reactive Fe-Mn (oxy)hyroxides (a weak acidreductive leach of 5 mM hydroxylamine hydrochloride (HH)-3 mM Na-EDTA-1.5%
acetic acid buffered to a pH of ~4 with NaOH), 3) calcite and additional carbonates
(1.7M acetic acid), 4) dolomite, and any remaining authigenic components (e.g.,

175 crystalline Fe-Mn oxides, 1M HCl). The focus of this study is the weak acid-reductive 176 leach, hereafter referred to as the HH leach, targeting amorphous reactive Fe-Mn 177 oxides. This is a more dilute HH leach than previous methods (e.g. Bhatia et al., 178 2013; Chester and Hughes, 1967) carried out on non-decarbonated sediments, 179 following Blaser et al., 2016. This method minimizes dissolution of silicate phases 180 and avoids the loss of other reactive Fe phases during removal of carbonates. Due 181 to the buffering of this reaction in the presence of varying amounts of carbonate, it 182 represents a minimum estimate of the reactive Fe-Mn oxyhydroxides rather than a 183 fully quantitative leach. This leach will mobilise reactive carbonate phases and 184 potentially some REEs associated with phosphates. Residue sediment was digested 185 following lithium borate fusion (Supplementary Text S1.3).

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# 187 **2.3 Major and trace concentration analysis**

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Major and selected trace element (e.g. Fe, Mn) concentrations on acidified waters
were determined by ICP-OES (Agilent 5100, with a precision and accuracy of better
than 10% based on repeated analysis of certified water standards (Supplementary
Text 1.4.1).

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Leachate phases and dissolved sediment residues were similarly analysed for major
and some trace element concentrations by ICP-OES, with matrix-matched calibration
lines. External reproducibility (typically better than 10%) was monitored using
certified standards (Supplementary Text 1.4.1).

Mass balance was verified by comparison of the sum of concentrations in the
leachates and residue with bulk sediment and a leached USGS shale standard with
certified values (Supplementary Table 8). The sum reproduced expected values
usually within 10%, and always within 20%.

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#### 204 2.4 REE Analysis

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206 REE concentrations on filtered waters were measured by isotope dilution 207 (Supplementary Text 1.5). Nd (alongside REE) concentrations in sediments and 208 sediment leachates were determined using ICP-OES (Agilent 5100, Nd 209 concentrations > ~4 ppb only) and using a Thermo Element-XR ICP-MS with a 210 matrix-matched calibration line. Reproducibility was monitored using certified 211 standards and values were within ±10%. Two residue samples and all sequential 212 extraction steps from one SPM sample were determined by isotope dilution, as 213 described for filtered water samples (Supplementary Text 1.5). 214 215 2.5 Nd and Sr isotopes 216 217 Nd was separated from the leachate and residue solutions using established 218 chromatographic procedures; the light REE's were separated using TRUspec resin, 219 and Nd from Sm using LNspec resin. Radiogenic Sr isotope ratios (<sup>87</sup>Sr/<sup>86</sup>Sr) were 220 determined on a subset of water, sediment leachate and sediment samples. Sr was 221 isolated using SrSpec resin (following Hindshaw et al, 2018). Nd was separated from 222 water samples using two stages of cation exchange (Biorad AG50W-X8) and

223 LnSpec resin (Supplementary Text 1.1).

225	Nd and Sr isotopes were measured on a Thermo Neptune Plus MC-ICP-MS.
226	<sup>146</sup> Nd/ <sup>144</sup> Nd was normalised to 0.7219 using an exponential correction. Samples
227	were corrected to the accepted value of reference standard JNdi-1;
228	<sup>143</sup> Nd/ <sup>144</sup> Nd=0.512115 (Tanaka et al., 2000) which was repeatedly analysed at the
229	sample concentration throughout each measurement session. Two times the
230	standard deviation (2 $\sigma$ ) on JNdi-1 for each measurement session is quoted as the
231	analytical uncertainty. $\epsilon Nd$ was calculated in parts per 10,000 relative to the
232	chondritic uniform reservoir (CHUR), <sup>143</sup> Nd/ <sup>144</sup> Nd <sub>CHUR</sub> =0.512638 (Jacobsen and
233	Wasserburg, 1980). For Nd isotopes, long-term reproducibility was monitored using
234	rock standards (Supplementary Text 1.4.2). Unless otherwise stated, where Nd
235	isotope replicates (measurement or full procedural) were carried out, the value
236	reported is the average of the replicates and the associated $2\sigma$ , if higher than the
237	measurement session uncertainty. For Sr isotopes, <sup>85</sup> Rb was monitored to correct for
238	Rb interferences on <sup>87</sup> Sr and Kr interferences were corrected for by measurement of
239	the baseline in a blank solution (on-peak zeros). Values were normalised to <sup>86</sup> Sr/ <sup>88</sup> Sr
240	= 0.1194 using an exponential correction. For Sr isotopes, samples were analysed in
241	duplicate, and the uncertainty quoted is the associated $2\sigma$ . Repeated measurements
242	of the standard NBS 987 yielded $^{87}$ Sr/ $^{86}$ Sr= 0.710275±46 ppm (2 $\sigma$ , n=20).
243	
244	3. Results

246 3.1 Dissolved Nd

248 Nd was partitioned between nano-particulate and colloidal phases and the 'truly' 249 dissolved (<1 nm; ≈1 kDa) phase (Table 1). Approximately 70% of Nd in the <0.2 µm 250 filtrate is hosted in colloids or nano-particulates (Table 1). In the Mackenzie River 251 (Middle Channel) ENd values between the <0.2 µm and <1 kDa fractions agreed within 0.43 epsilon units (Table 1). This demonstrates that nano-particulate and 252 253 colloidal phases and the 'truly' dissolved load have the same source and so the <0.2 254 μm εNd value is representative of the 'truly' dissolved composition, consistent with 255 previous observations in large rivers (Merschel et al., 2017).

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# 257 **3.2 Paired sediment and dissolved εNd**

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259 As the <0.2  $\mu$ m fraction is indicative of the 'truly' dissolved  $\epsilon$ Nd, it is possible to use 260 these measurements to compare with leachable phases and assess their reactivity. 261 We therefore refer to the <0.2 µm as the dissolved load. The dissolved load is 262 always more radiogenic than the sediment residue, with the offset ranging from 1.4 263 to 8.2 epsilon units (Fig. 3, average = 3.2). This is consistent with, and in some cases far exceeds, the offset observed by Goldstein and Jacobsen, 1987, for large 264 265 rivers with significant marine sedimentary source rocks in their catchments. In 266 contrast, ENd in the dissolved load is in close agreement with HH leachates targeting 267 reactive Fe-Mn oxides with a linear regression within error of the 1:1 line (Fig. 3). 268 The dissolved vs. residue ɛNd linear regression is systematically offset from the 1:1 line, with a weaker correlation (Fig. 3). 269

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271 Residue εNd compositions are in close agreement with previous measurements of
272 sediment already reported for the Mackenzie River (Vonk et al., 2015). Dissolved

273  $\epsilon$ Nd reported herein for the Mackenzie River at Tsiigehtchic (-13.4 ±0.2 ( $2\sigma$ , n=1,

274 2018) and -12.3 $\pm$ 0.3 (2 $\sigma$ , n=2, 2017)) is consistent with the only previous dissolved 275 measurement (2003, -12.9 $\pm$ 0.3 (Zimmermann et al., 2009)).

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SPM in the Middle Channel of the Mackenzie River is hydrodynamically sorted, due 277 278 to faster settling velocities of coarser sedimentary particles (e.g. Bouchez et al., 279 2011). This is illustrated by Al/Si, a proxy for grain size (Fig. 4), indicating the 280 dominance of finer sediments towards the surface and coarser at the bottom. 281 Despite this hydrodynamic sorting, there is no appreciable variation of  $\epsilon$ Nd with 282 depth in either the sediment residue or HH leachate (Fig. 4). Dissolved  $\epsilon$ Nd from the 283 surface and base of the water column demonstrate that the water is well mixed. 284 285 There are systematic differences in dissolved ɛNd between 2017 and 2018. The 286 furthest downstream site (Mackenzie, Middle Channel) dissolved composition 287 changes from  $\epsilon$ Nd=-10.71±0.41 (2 $\sigma$ , n=3, 2017) to -12.97±0.15 (2 $\sigma$ , n=1, 2018), 288 likely driven by changes in the Peel River which shows similar differences, changing 289 from εNd=-9.83±0.16 (2σ, n=1, 2017) to -11.47±0.15 (2σ, n=1, 2018). This could 290 represent inter-annual ɛNd variability, as is argued to be the case for changes in 291 organic carbon and dissolved nutrient sources on the same sample set (Schwab et

al., 2020), but additional data would be needed to determine if this is the case.

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The average HH leach  $\epsilon$ Nd from sediment cores was -11.3±1.9 (2 $\sigma$ , n=6), consistent with the same fraction in riverine sediments (Fig. 3) with the leachate always more radiogenic compared to the silicate residue (two cores, <10m deep, average offset = 2.7±1.2, 2 $\sigma$ , n=2). Shallow (<100 m) seawater in the Beaufort Sea is more radiogenic than these leachate compositions ( $\epsilon$ Nd = ~-6 to -9, Porcelli et al., 2009). Therefore, the offset between residue and leachate is preserved and exported in sediment to the shelf.  $\epsilon$ Nd in the HH leachate at the deepest and least proximal core site ( $\epsilon$ Nd = -9.55, 1054 m water depth) overlaps with  $\epsilon$ Nd measured in deep (> 1000 m) seawater in the Beaufort Sea, which is less radiogenic ( $\epsilon$ Nd between -9 and -11, Porcelli et al., 2009).

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### 305 **3.3 Sediment partitioning and characterisation**

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307 The partitioning of selected elements amongst all leachates relative to the bulk 308 sediment is shown in Fig. 5. In the NH<sub>4</sub>Cl leach, targeting exchangeable phases, Fe 309 and Nd were below detection. This is consistent with a previous study which reported 310 low concentrations of REEs in a river sediment exchangeable fraction (Adebayo et 311 al., 2018). Elemental concentrations in the NH<sub>4</sub>Cl leach which were not below 312 detection are reported in Tipper et al., 2021. The HH leach contains on average 3%, 8% and 30% of the total sediment Fe, Nd and Mn respectively (Fig. 5). Ca, Mg and 313 314 Sr are also high in the HH leach (Fig. 5) consistent with significant carbonate 315 lithologies present within the Mackenzie Basin (Millot et al., 2003). On average 60% 316 of the Fe and Nd are hosted within the silicate residue. 317 318 REE on the HH leach show a middle REE (MREE) enrichment (Fig. 6a). A MREE

enrichment of this magnitude is typical of an Fe-Mn oxyhydroxide signature (Fig. 6a)
and is consistent with that previously reported for similar sediment extractions (e.g.

321 Leybourne and Johannesson, 2008).

323 REE in the SPM leachate phases were partitioned relative to the bulk (sum of 324 leachates and residue; Middle Channel, 2017, Fig 6b). 15% of REEs relative to the 325 total sediment are mobilised during the HH leaching step. A smaller fraction (<1%) of 326 REEs are released during the subsequent acetic acid leach step (Fig. 6b), despite 327 the large amount of Ca and Mg (22% of total sediment Ca, 11% of total sediment 328 Mg). Less Fe and Mn are mobilised in the acetic acid leach (1% of total Fe, 5% of 329 total Mn), relative to the HH leach (4% of total Fe, 42% of total Mn). This 330 demonstrates that the REE are not associated with carbonate phases. 331 332 <sup>87</sup>Sr/<sup>86</sup>Sr on the dissolved load and sediment residues (Fig. 6c) are in agreement 333 with previously published data for this river (e.g. Millot et al., 2003). <sup>87</sup>Sr/<sup>86</sup>Sr values

indicate carbonate and silicate sources in the HH leach and residue respectively. Sr isotopes do not indicate extensive leaching into the silicate detrital fraction. Although there is scatter in the leachable  $\varepsilon$ Nd compositions (Fig. 6c), on stronger leaching  $\varepsilon$ Nd is not substantially more radiogenic than the HH leach.

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#### 339 4. Discussion

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341 £Nd values from paired dissolved load and SPM, alongside the characterisation of 342 elemental concentrations in sequential extractions, are used to trace both the source 343 and the reactivity of different phases within the SPM. Firstly, we seek to assess the 344 reactivity of the phases in the HH leach and then characterise sources. We then 345 estimate the flux of labile Fe from the Mackenzie to the Arctic Ocean and address 346 the implications of this study in terms of the oceanic Nd budget.

#### 348 **4.1 Reactivity of the labile phases in SPM**

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350 The reactivity of phases hosted in the HH leachate are traced with ɛNd and 351 elemental partitioning. Significant fractions of Fe, Mn, Ca, Mg, and REEs relative to 352 the bulk sediment are present in what is an extremely weak and buffered leach (5 353 mM HH, 1.5% acetic acid, Fig. 5, Fig. 6). Although subsequent leaches use stronger 354 reactants and are not buffered, they release smaller amounts of Nd compared to the 355 HH leach (Fig. 6b). This suggests that some of these reactive phases are 356 susceptible to acid-reductive dissolution with small changes in pH or redox 357 conditions (which may occur in the delta or offshore). Nd and REEs are depleted in 358 carbonate rocks and are unlikely to be directly incorporated into biogenic calcite (e.g. 359 Tachikawa et al., 2014). Less Nd is released in the 1.7 M acetic acid leach, 360 compared to the HH leach, confirming that during the HH leaching step the Nd 361 released is not associated with carbonates. 362 363 Although an MREE enrichment in the HH leach is typical of an Fe-Mn oxyhydroxide 364 phase, this could also be consistent with some phosphate phases (such as biogenic

apatite, Fig. 6a., Leybourne and Johannesson, 2008, and references therein).

366 Secondary precipitation of phosphates could also result in an MREE enrichment

367 (Leybourne and Johannesson, 2008), but the lack of an LREE enrichment (low

368 HREE/LREE, Fig. 6a) in HH leaches makes this unlikely (Köhler et al., 2005).

369

The dissolved εNd composition is always similar to the HH leachate composition
(Fig. 3). This similarity is prevalent in different sampling localities and tributaries,
which differ in sediment and water characteristics. This points to a fundamental

process that is shared between these catchments and suggests that this observation
is not unique to the time of year sampled (freshet). In rivers, the SPM is supplied by
bank erosion and hillslope processes (Hilton et al., 2015), which are likely to be
decoupled in space and time from the hydrological pathways that deliver river water
(Vonk et al., 2019). Therefore, the consistency in εNd between dissolved and HH
leach εNd is most likely explained by rapid and recent cycling within riverine implying
equilibrium between the dissolved load and Fe-oxide phases in the HH leach.

381 Dissolved and HH leachate ɛNd are always similar at each site in the Mackenzie 382 River, both upstream and downstream of major confluences with the Peel and Arctic 383 Red Rivers (Tsiigehtchic to Middle Channel, Fig. 1a). Previous discharge and 384 sediment flux estimates demonstrate that water and suspended sediment do not mix 385 in the same proportions downstream of the Peel confluence, with the Peel supplying 386 17% of the sediment load (Carson et al., 1998), but only 7% of the total water 387 discharge (Schwab et al., 2020 and references therein). This implies the potential for 388 rapid cycling between the dissolved load and HH leach phases during transit 389 between the upstream and downstream sites, despite only limited changes in pH and 390 Nd concentration (supplementary Table S4 and S10).

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Although the mechanism for equilibration of Nd between HH leach phases and the dissolved load is unclear, since the weakly adsorbed exchangeable phases are removed in the leaching step prior to the HH leach, this rapid cycling is likely driven by dissolution of Nd from phases such as Fe-Mn oxyhydroxides. The concentration of Nd in the HH leach (ppb to ppm, Supplementary Table S9 and S11) is several orders of magnitude higher than the dissolved concentration (Supplementary Table S10), suggesting that phases within the HH leach may drive the dissolvedcomposition.

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Our findings demonstrate dynamic cycling between the dissolved load and phases
soluble in the HH leach, such as Fe-Mn oxyhydroxides. The similarity of the
dissolved load and HH leach phases and decoupling from the silicate residue
suggests that the HH leach phases are labile and likely bioavailable. These results
imply not only rapid cycling of Nd but potentially other immobile elements contained
within the HH leach.

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# 408 **4.2 Marine authigenic phases as a source of radiogenic Nd in river waters**409

410 The mass balance of sources of radiogenic Nd that contribute to higher ɛNd in the 411 dissolved and HH fractions relative to silicates provides information on the relative 412 supply of Nd from different phases. Here we discuss possible sources of radiogenic 413 Nd that could explain both the HH leaches and dissolved load. Average global 414 seawater ɛNd (~-8.8, Lacan et al., 2012) and modern core top authigenic fractions 415 (Haley et al., 2017) are more radiogenic than the upper continental crust (-11.4±2.5, 416 Goldstein et al., 1984). Therefore, when marine sediments are emplaced on the 417 continents, they can provide a distinct source of radiogenic Nd to the weathering 418 reactor. Measurements of ɛNd on Fe-Mn oxides and other inherited authigenic 419 phases of marine origin hosted in sedimentary rocks show that they are on average 420 more radiogenic than associated silicates (Jang et al., 2020, Hindshaw et al., 2018). 421 They can be reactive and so result in a more radiogenic dissolved and leachate

422 composition when compared to the bulk source rock in rivers (Bayon et al., 2020;
423 Goldstein and Jacobsen, 1987; Hindshaw et al., 2018; Jang et al., 2020).

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425 Preferential weathering of inherited marine authigenic phases is the most likely 426 reason for decoupled ɛNd between dissolved and silicate residue sediment in the 427 Mackenzie. Radiogenic Nd in Fe-Mn oxyhydroxides of Mackenzie River SPM is likely 428 derived from the weathering of these phases through physical transportation, and 429 equilibration with waters. These radiogenic phases become mixed with Fe-oxides 430 resulting from the weathering of silicate minerals, with *c*Nd inherited from the source 431 silicate. Nonetheless, it is possible that alternate sources contribute labile and 432 radiogenic Nd, and these are addressed below.

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#### 434 **4.3 Alternative sources of radiogenic Nd to river waters**

Accessory minerals enriched in REEs and garnet can have more radiogenic εNd
than their bulk source rocks (e.g. Rickli et al., 2017). However, radiogenic release of
Nd during incongruent weathering of crystalline rocks is minimal (Dausmann et al.,
2019). Another possible source of radiogenic Nd is the preferential weathering of
volcanic rocks, but they are rare in the Mackenzie basin (Fig 1b, Millot et al., 2003).

Several studies indicate the dominance of shale and carbonate weathering on the Mackenzie dissolved load, with no evidence of large volcanic or crystalline rock contributions (Horan et al., 2019; Huh et al., 2004; Millot et al., 2003). cNd decoupling between dissolved and silicate residue is at a maximum in a small Peel River tributary that, to the best of our knowledge, does not have crystalline or volcanic rocks in its catchment and drains almost exclusively shales. Moreover, preferential weathering of a specific rock type would not explain the coupling of the
dissolved load with the HH leach, and their decoupling from the silicate residue, in all
sampling sites, with differing source lithologies.

450 Phosphates in sedimentary rocks are enriched in REEs and may be partially 451 mobilised in the HH leach. Phosphates in the main source rock type (shales) are 452 likely to have formed during marine diagenesis, so can be considered as inherited 453 authigenic phases, with a comparable  $\varepsilon$ Nd to concurrent Fe-Mn oxides. Fe-oxides 454 are ubiguitous in all sediments and sedimentary rocks. Given the range in source 455 rock age (Proterozoic to Cenozoic, Wheeler, 1996) and depositional environment, 456 Fe-Mn oxides are the most probable universal constituent consistent with the high 457 fraction of Fe and Mn in the HH leach, whereas phosphates may not always be 458 present in sufficient abundance. Equilibration between the dissolved load and 459 phosphates hosted in the HH leach would not likely result in the observed REE 460 patterns, as secondary phosphates are typically LREE enriched (Fig. 6a, Köhler et 461 al., 2005). Therefore, it is unlikely that phosphate minerals play a significant role, and 462 the more positive Nd is derived from the preferential weathering of inherited Fe-Mn 463 oxides.

464

While Fe-Mn oxides are insoluble in oxygenated and neutral range pH which typify large rivers including the Mackenzie (Tank et al., 2016) mobilization of amorphous and nano-particulate Fe can occur, and in acidic environments dissolution of Feoxides may occur. Extensive pyrite oxidation provides the majority of dissolved sulfate in the Mackenzie (Calmels et al., 2007). A product of the oxidative weathering of pyrite is Fe-oxide, and if instantaneous buffering of this reaction by dissolution of carbonate minerals does not occur, it will result in an acidic pH alongside high sulfate 472 concentrations (Horan et al., 2019). This is seen in one tributary of the Peel River 473 where high sulfate concentrations (4660 μM) and relatively low pH of ~6 are coupled 474 to elevated Nd (950 ppt), high Fe (1 μM) and Mn (7.8 μM) concentrations in the 475 dissolved load (<0.2 μm, Supplementary Tables S9 and S12) with the highest 476 observed offset between the residue and HH leach εNd (8 epsilon units, 477 Supplementary Table S10).

478

479 Nd is not likely to be incorporated directly into pyrite but will be present in

480 'contaminant' phases in pyrite or in other phases within a sedimentary rock

481 succession, namely Fe-Mn oxyhydroxides or phosphates (Raiswell and Plant, 1980).

High Nd and Mn dissolved concentrations indicate the dissolution of reactive Fe-Mn
oxides within the source rock (shale), with differing εNd to silicate sources, coupled

484 to the oxidative weathering of pyrite.

485

486 The HH leach of a shale fragment from this catchment has comparable  $\varepsilon$ Nd to the 487 bedload HH leachate and dissolved load (Fig. 3, Supplementary Table S10). Although the mineralogical source of Nd and Fe is different (Fe from pyrite and Nd 488 489 from other phases) these observations imply that the mobilization of Fe and Nd are 490 coupled during the oxidative weathering of pyrite. This suggests that oxidative 491 weathering of pyrite may provide an important source of reactive Nd and Fe bearing 492 phases, with radiogenic  $\varepsilon$ Nd. Once the REEs are in solution, they can be 493 incorporated into secondary Fe-oxides upon increasing pH, when Fe-oxides bearing 494 REEs will precipitate out of solution. Close coupling between riverine Nd and Fe has 495 been noted in previous studies (e.g. Elderfield et al., 1990; Ingri et al., 2000).

#### 497 **4.4 Constraints on the contribution of different sources of labile Fe and Nd**

498

Using a simple mass balance model it is possible to estimate the fraction of Nd and
Fe sourced from inherited authigenic phases contained within the sedimentary
source rocks.

502

503 Dissolved  $\epsilon$ Nd is set by exchange with labile HH leach phases in the SPM. The labile 504  $\epsilon$ Nd is considered to be a mixture of Fe-oxides derived from modern-day weathering 505 of silicate minerals ( $\epsilon$ Nd<sub>sil</sub>) and Nd derived from inherited authigenic phases ( $\epsilon$ Nd<sub>auth</sub>, 506 equation 1, Figure 7).

508 where  $\epsilon Nd_{labile}$  is the average of the leachate and dissolved composition, and f is the 509 fraction of Nd in either the inherited authigenic or silicate phases where  $f_{Nd,auth} + f_{Nd,sil}$ 510 = 1.

511

Equation 1 was solved for  $f_{Nd,auth}$ , with  $\epsilon Nd_{sil}$  given by the residue silicate composition 512 513 measured for sample set. ENdlabile is the average of the HH leachate composition and 514 (if measured) the dissolved composition for each sample set. ENd<sub>auth</sub> is unknown and 515 given the range of ages and types of sedimentary rocks in the Mackenzie basin, will 516 be variable.  $\epsilon$ Nd<sub>auth</sub> was considered over the range  $\epsilon$ Nd<sub>auth</sub>=0 to  $\epsilon$ Nd<sub>auth</sub>=  $\epsilon$ Nd<sub>labile</sub> 517 sampled from a synthetic uniform distribution to account for the uncertainty of this 518 end-member. ENdiabile and ENdsil were measured at each sampling location and 519 considered as normal distributions. We used Monte-Carlo methods to estimate the 520 error when solving for f<sub>Nd,auth</sub> (calculating f<sub>Nd,auth</sub> for 100,000 values of each variable). 521 f<sub>Nd,auth</sub> was calculated for 4 individual sample sets (2017 and 2018 Middle Channel,

522 Arctic Ocean and in shelf sediment cores, Fig. 8, Supplementary Table S13). Taking 523 the interquartile range from the sum of all four sample sets it estimated that, at the 524 mouth of the Mackenzie River a minimum of 23-50% of labile Nd is derived from 525 inherited authigenic phases (Fig. 8).

526

527 Given this significant estimate for f<sub>Nd,auth</sub>, it is likely that a similarly high fraction of 528 other immobile elements such are Fe are also sourced from inherited authigenic 529 phases. The fraction of labile Fe sourced from inherited authigenic phases (including 530 from pyrite and inherited Fe-oxides) was estimated using mass balance:

531 
$$\left(\frac{Nd}{Fe}\right)_{labile} = f_{Fe,sil} \left(\frac{Nd}{Fe}\right)_{sil} + f_{Fe,auth} \left(\frac{Nd}{Fe}\right)_{auth}$$
 (2)

532 where  $f_{Fe,auth} + f_{Fe,sil} = 1$ , and  $\left(\frac{Nd}{Fe}\right)_i$  is the molar ratio of Nd to Fe in each phase.

This equation was solved for  $f_{Fe,auth}$ , the fraction of labile Fe sourced from inherited 533 authigenic phases.  $\left(\frac{Nd}{Fe}\right)_{auth}$  is unknown, and the largest source of error and was 534 535 based on the range from HH leaches of a shale fragment and coarse bank sediment  $\left(\left(\frac{Nd}{Fe}\right) \times 1000 = 0.63 \text{ to } 2.4\right)$ .  $\left(\frac{\{Nd\}}{\{Fe\}}\right)_{cil}$  was taken as the average of the measured 536 537 silicate residue values which are within error of average upper continental crustal estimates  $\left(\binom{Nd}{Fe} \times 1000 = 0.33 \pm 0.1$ , Supplementary Text 3).  $\left(\frac{Nd}{Fe}\right)_{labile}$  is the measured 538 HH leachate composition at each sampling site. Using Monte Carlo methods to 539 540 estimate errors, f<sub>Fe,auth</sub> is estimated to be between 11-29% (interguartile range, from 541 sum of Middle Channel samples sets, 2017 and 2018, Supplementary Table S14). 542

543 Despite the uncertainty it is clear that the SPM in the Mackenzie must host a 544 significant fraction of labile Nd and Fe which is derived from inherited authigenic 545 phases in sedimentary rocks. Accordingly, the weathering of sedimentary rocks,

546 especially marine shales, must supply a significant amount of reactive immobile

547 elements (Nd, Fe) in the Mackenzie basin.

548

# 549 **4.5 The flux of labile, potentially bioavailable, Fe**

550

551 We have demonstrated that the SPM in the Mackenzie River has a labile Fe phase.

552 To estimate its supply to the Arctic Ocean, we use the suspended sediment flux,

553 which is well documented for the Mackenzie in the years 1974-1994. There is

significant inter-annual variability ranging from 81 Tg/yr to 224 Tg/yr (Carson et al.,

555 1998) with a consistent seasonal peak in the summer months following freshet, the

556 time period over which our samples were collected (Fig. 2).

557

558 The labile, potentially bioavailable, Fe concentration was considered to be equal to 559 the HH leach Fe concentration measured on SPM from the Mackenzie (Middle 560 Channel, 2017 and 2018), which approximated to a normal distribution (1466±350 561  $\mu g/g$  (±1 $\sigma$ , n=16)). The uncertainty was estimated using Monte-Carlo methods 562 (n=100000), similar to that used by Hilton et al., 2015. The sediment flux was 563 considered as a uniform distribution, taking into account all potential values at equal 564 probability (81 Tg/yr to 224 Tg/yr, Carson et al., 1998), a maximum approximation of 565 error. The median (±interguartile range) flux of labile Fe associated with SPM in the Mackenzie River was calculated to be 0.21(+0.06,-0.05) Tg/yr. We do not include 566 567 dissolved (<0.2 µm) Fe concentrations in this estimate as they are negligible in 568 comparison to Fe hosted in the reactive sediment phase (estimated dissolved Fe flux 569 represents <0.03% of the labile sediment Fe flux, Supplementary Text 4).

570

571 The calculated flux of potentially bioavailable Fe in the SPM of the Mackenzie is 572 substantial; comparable to that from the Greenland Ice Sheet (~0.3 Tg/yr, Bhatia et 573 al., 2013). Greenland Ice Sheet fluxes are estimated using the sum of strong SPM 574 leaching and dissolved Fe followed by the application of an estuarine loss factor 575 (90%, Bhatia et al., 2013).

576

577 Much of the SPM from the Mackenzie is rapidly buried offshore or trapped in the 578 delta, and the loss due to estuarine processes is unknown. Even if ~90% of the 579 bioavailable Fe in the SPM is removed or trapped, the resulting flux to the oceans is 580 still significant (~0.02 Tg/yr). Moreover, benthic release of this particulate associated 581 Fe, alongside Nd, on the Mackenzie shelf may play an important role in delivering a 582 higher proportion of this labile phase to the Arctic Ocean.

583

584 For every two moles of sulfate released from the oxidative weathering of pyrite in the Mackenzie basin, 1 mole of Fe will be incorporated into an Fe-oxide. Using the 585 586 pyrite-derived sulfate flux of Calmels et al., 2007 it is estimated that the oxidative 587 weathering of pyrite will result in the production of ~3.6 Tg/yr of Fe, as Fe-oxides. 588 This is around an order of magnitude greater than the labile Fe flux associated with 589 SPM that we estimate herein (0.2 Tg/yr). This implies a large accumulation of Fe-590 oxides derived from the oxidative weathering of pyrite within the critical zone in the 591 Mackenzie basin, which are not exported to labile phases in the SPM. 592

4.6 Nd isotopic end-member compositions and inputs to the Arctic ocean

595 The Mackenzie River is a large point source of sediment and water to the Arctic 596 Ocean: therefore, it is an important finding that labile Nd has an isotopic composition 597 that is distinct from the bulk SPM. This finding is potentially applicable to all riverine 598 catchments with marine sedimentary source rocks. It is equally noteworthy that there 599 is variability in dissolved ɛNd between sampling years. Although our data is similar to 600 the only previous dissolved data point on the Mackenzie (Zimmermann et al., 2009), 601 it extends the range with implications for studies involving  $\epsilon$ Nd in the Arctic Ocean 602 (e.g. Deschamps et al., 2019). Despite variability the HH leachates from shallow 603 marine core sites reflect the average dissolved composition of the Mackenzie over 604 decadal timescales. Therefore, such leachates from shelf sediment may provide a 605 more useful integrated ɛNd end-member. This isotopically distinct, labile, phase is 606 maintained offshore, meaning that any labelling of seawater via 'boundary exchange' 607 processes on the shelf will have a significantly more radiogenic composition than the 608 bulk sediment. This suggests that bulk shelf sediment and continental margin ɛNd 609 may be systematically offset from the likely oceanic source (Jeandel et al., 2007) as 610 it is the reactive (leachable) compositions which will characterise any shelf inputs.

611

# 612 5 Conclusions

613

This study highlights the potential for sedimentary rock weathering in the Arctic to provide a critical source of labile Fe. As approximately half of the rocks weathered on land draining into the Arctic Ocean are shales (Amiotte Suchet et al., 2003), similar observations to this study would be expected in other Arctic rivers. The on-going state of permafrost thaw in the Mackenzie basin has resulted in increased slumping in the Peel catchment which has exposed fresh pyrite minerals and resulted in increased sulfuric acid weathering (Zolkos et al., 2018). Increased exposure of pyrite
may also lead to increased mobility and dissolution of REEs (and Fe) within this
catchment. However, it is also important to note that delivery of bioavailable Fe from
the Mackenzie basin to the Arctic Ocean will have changed in the past, including
during periods of glaciation and glacial retreat in this region. Melt water and icerafting could have allowed for more direct delivery of un-weathered labile Fe phases
to the Arctic Ocean.

627

628 Recycled sedimentary components within source rocks may provide labile 629 micronutrients and other trace elements, a process which is likely applicable globally, 630 and not just in the Arctic. Fe, and the supply of other micronutrients that limit primary 631 productivity, are thought to be capable of having long-term and far-reaching effects 632 on global climate, by way of their potential for increasing productivity and organic 633 carbon burial in the oceans. Therefore, when considering increases in nutrient 634 availability in the past, the source lithology and its reactivity must be taken into 635 consideration. We highlight the impact of source lithology on the budget of labile 636 trace elements (Nd) and micronutrients (Fe) in rivers, which has implications for both 637 present and past fluxes to the oceans and paves the way for restructuring how 638 riverine SPM is viewed in terms of its reactivity when sedimentary rocks are present 639 in the source catchment.

640

#### 641 Acknowledgements

642

M. Greaves, G. Hughes, H. Chapman, M. Bickle and K. Relph are thanked for their
assistance in the lab. E. Amos and M. Schwab are thanked for assisting with

645 fieldwork. E. Stevenson is thanked for her help and advice on Arctic fieldwork and the co-precipitation method. M. Murphy and C. Arendt are thanked for sharing their 646 647 methods for ultra-filtration and FeCl<sub>3</sub> cleaning respectively. Y. Plancherel and E. 648 Stevenson are thanked for supplying filtered Severn River water used as an internal standard. K. Jarrett and the Geological Survey of Canada, Marine Geoscience 649 650 Collection are thanked for supplying sediment core subsamples. The authors both 651 thank and acknowledge the assistance of the Aurora College Research Institute, 652 Inuvik. Samples were collected under research licenses 15288 and 16106. C.S.L 653 was funded by a NERC studentship (NE/L002507/1) with support for travel to 654 Canada in 2018 from the University of Cambridge Department of Earth Sciences 655 Leave to Work Away Research Fund. R.G.H acknowledges funding from the 656 European Research Council (Starting Grant ROC-CO<sub>2</sub>, 678779) and a NERC UK-657 Canada Arctic Partnership Bursaries Program. E.T.T acknowledges funding from 658 NERC Standard Grant 703 NE/P011659/1. The manuscript was greatly improved by 659 comments from two helpful anonymous reviewers and the editor L. Derry. 660

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882	Figure Captions
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884	Figure 1. Area of study A. Sample locations and elevation (GDEM, resolution 30 Arc
885	second). B. Bedrock geology of the Mackenzie Basin (shaded black line),
886	categorized by rock type (Wheeler et al., 1996).White circles indicate the location of
887	sampling sites identified in A.

888

Figure 2. Water discharge from the Mackenzie River (2017 and 2018). Daily
discharge (lines) from Environment Canada gauging station at Tsiigehtchic, retrieved
from the ArcticGro database (https://www.arcticrivers.org/data, Supplementary text
S2). ADCP Discharge at the Middle Channel and Tsiigehtchic (this study, light blue,
2017, dark blue 2018).

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895 **Figure 3.** ENd on coupled HH leachates (circles), dissolved (<0.2µm only shown) 896 and silicate residue sediment (triangles) at all sampling localities (2017 and 2018). 897 Probability density of sediment residue (green) and leachate (pink) shown, difference 898 in modes (dashed grey line) is indicated (black arrow). Open symbols are bank 899 sediment and coloured symbols, SPM, coloured diamonds are sediment core data. 900 Coloured squares are a shale fragment HH leachate and residue (pink and green 901 respectively), which was not used in the linear regression. Error bars are  $2\sigma$ 902 analytical uncertainty (where visible or smaller than the symbol size). 1:1 line 903 (dashed line) is shown.

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Figure 4. Two sediment depth profiles from the Mackenzie River (Middle Channel,
2017). Water depth is normalised to the deepest sample. Error bars are 2σ analytical
uncertainty. Dissolved compositions are from waters obtained at the surface and at
depth.

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Figure 5. Probability density of the fraction of a suite of elements in leachates
relative to the total (sum of leachates and residue). Data is from all sampling sites
across both years. The fraction of the HH leachate, acetic acid leachate, and the

913 sum of all leaching steps are shown. Symbols are data points that form the914 probability distribution.

915

## 916 Figure 6. Characterisation of sequential extractions A. MREE/MREE\*

917 HREE/LREE cross-plot, following Martin et al., 2010, alongside a non-exhaustive 918 literature compilation (plotted for reference, Supplementary Text S2). HREE=  $\Sigma(Tm, Text S2)$ 919 Yb, Lu) LREE=  $\Sigma$ (La, Pr, Nd), MREE=  $\Sigma$ (Gd, Tb, Dy), MREE\*=(HREE+LREE)/2. All 920 concentrations are normalised to PAAS (after Pourmand et al., 2012). Foraminiferal 921 coatings and marine leachates are extractions thought to represent Fe-Mn 922 oxyhydroxides. **B.** A REE partitioning in a Mackenzie (Middle Channel, 2017) SPM 923 sequential extraction. Left hand side axis is the REE concentration, right hand side 924 axis fraction of total (sum of leachates and residue) REE concentration (bars) in 925 each phase. **C.**<sup>87</sup>Sr/<sup>86</sup>Sr against εNd on Mackenzie (Middle Channel, 2017) SPM sequential extractions and water (87Sr/86Sr data point from Tipper et al., 2021). 20 926 927 analytical uncertainty is shown, unless smaller than the symbol size.

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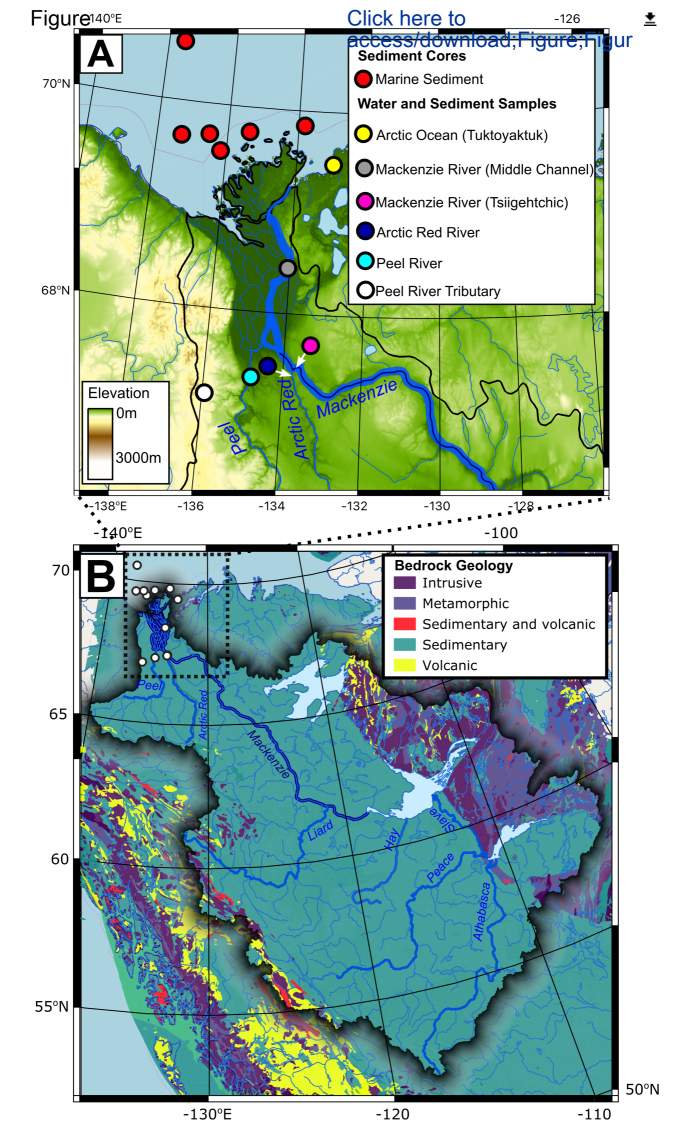
Figure 7. Schematic indicating the mechanisms presented herein that may lead to
more radiogenic εNd in the HH leachates of SPM and the dissolved load of the
Mackenzie river, when compared to the SPM silicate residue.

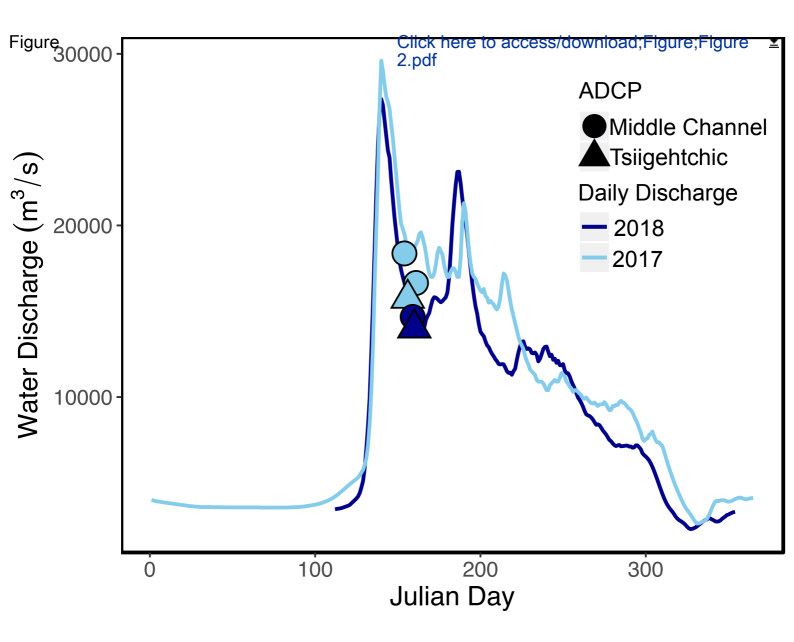
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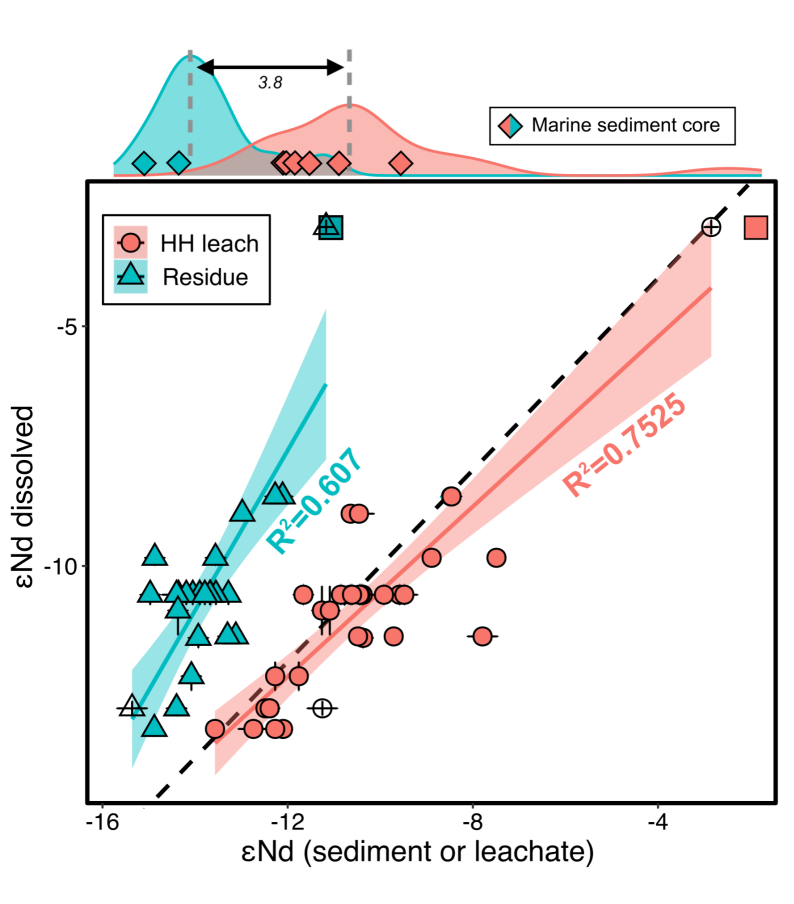
Figure 8. Probability density of calculated f<sub>Nd,auth</sub> (equation 1) for 4 sampling sites
generated using Monte Carlo methods. Median value (white line) and interquartile
range (dashed lines) are shown. Input compositions for this calculation are listed in
Table S13.

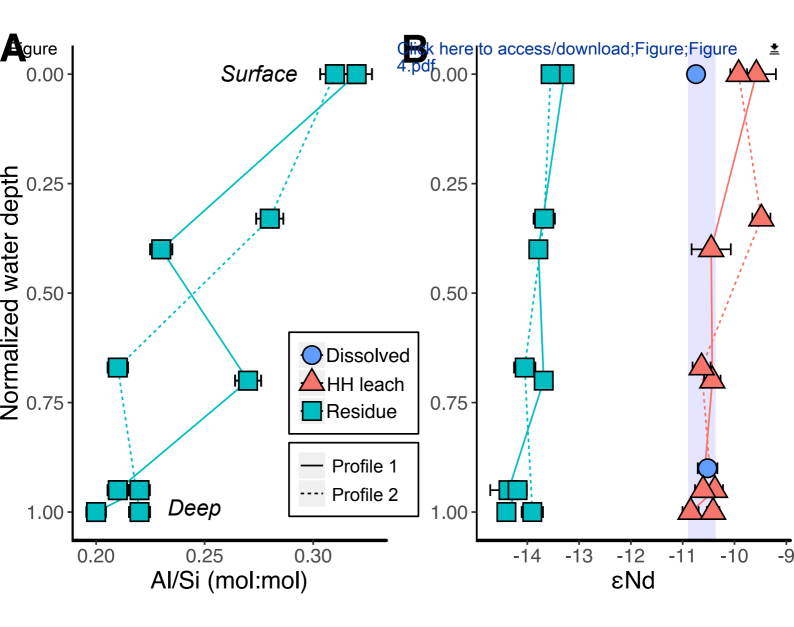
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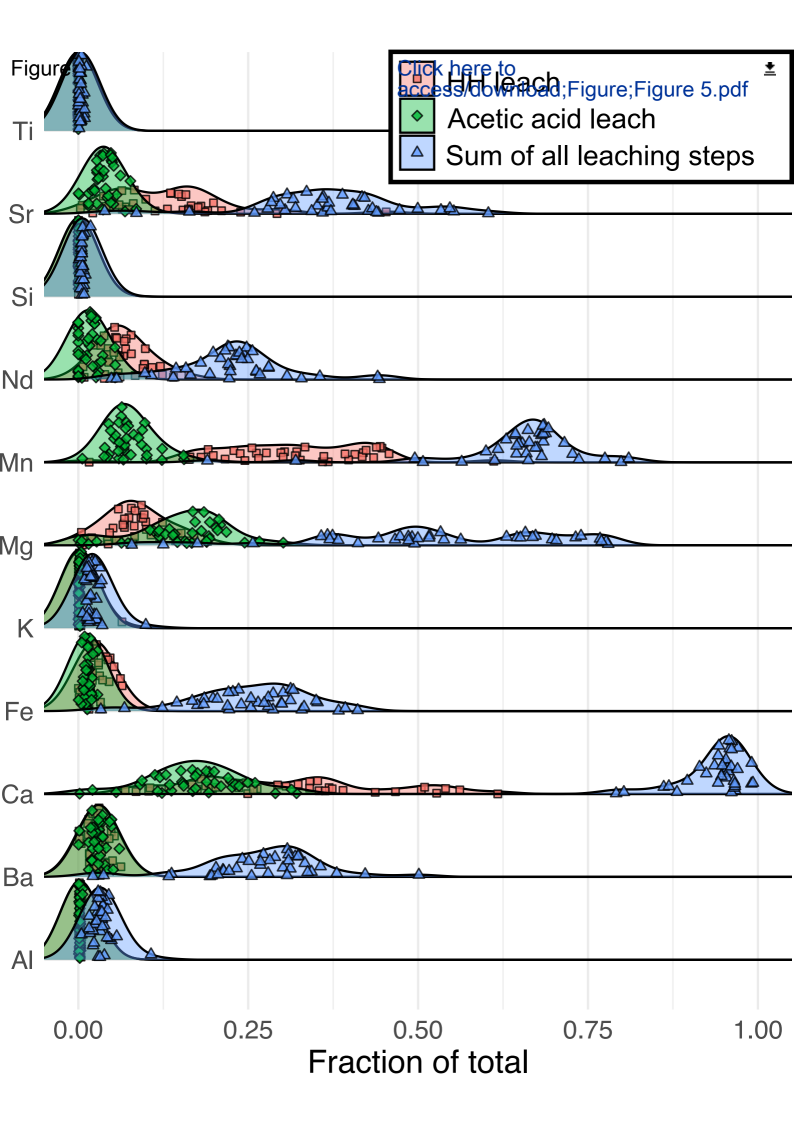
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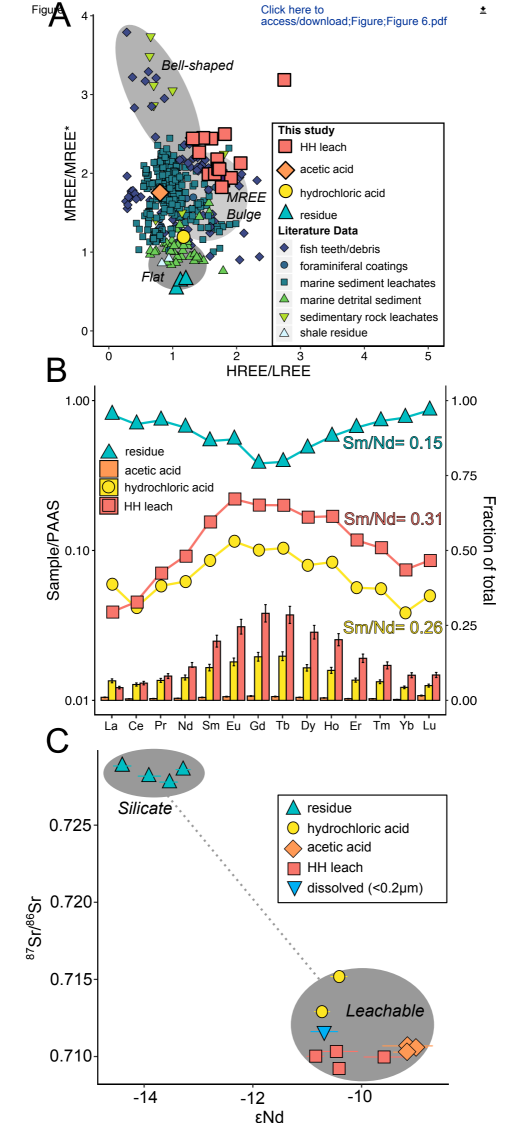


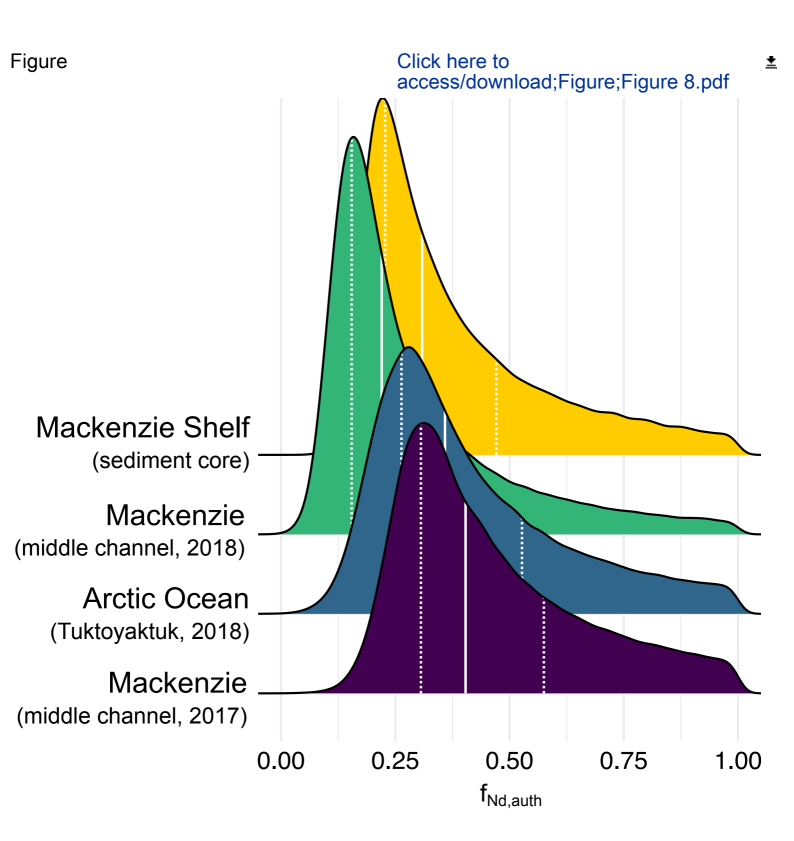


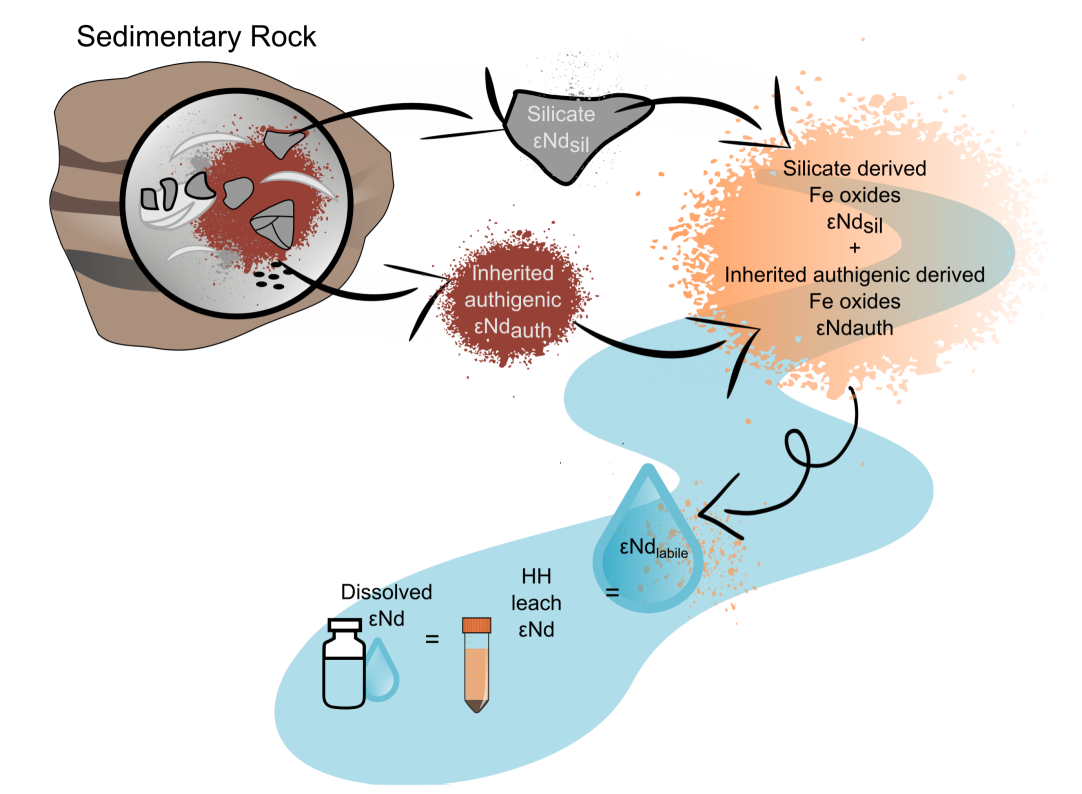












**Table 1.** εNd and Nd concentrations on different filtrates from the Mackenzie River and a major tributary (2018).

River	Filter	εNd	Error	[Nd]	Error	% of
NIVEI	Size/MWCO	cinu	(2σ) <sup>1</sup>	ppt	(2σ)	<0.2µm
Mackenzie	<0.2µm	-12.97	0.15	20.7 <sup>2</sup>	0.7 (n=2)	
River	10kda	-12.41	0.51	10.4 <sup>3</sup>	1.4 (n=2)	50%
(Middle	1kda	-13.40	0.34	5.9 <sup>3</sup>	0.4 (n=2)	29%
Channel)						
	<0.2µm	-11.47	0.15	12.1	n/a (n=1)	
Peel River	10kda	-11.61	0.34	7.1 <sup>2</sup>	0.1 (n=2)	59%
	1kda	n.d	n.d	4.1 <sup>3</sup>	0.8 (n=2)	34%

<sup>1</sup>Analytical uncertainty, <sup>2</sup>Average and 2σ of two replicate measurements during different analytical sessions, <sup>3</sup>Average and 2σ of two full procedural replicates (including filtration)

## Author contributions

C.S.L, E.T.T., A.M.P, R.G.H and R.S.H designed the study. C.S.L, R.G.H, M.D, J.J.B carried out the sample collection and fieldwork. R.G.H and E.T.T and C.S.L obtained funding for fieldwork and analysis. C.S.L and R.W carried out method development to analyse the samples. C.S.L analysed the samples and wrote the manuscript under the supervision of E.T.T and A.M.P with input and guidance from all co-authors.