Laurentide Ice Sheet extent over the last 130 thousand years traced by the Pb isotope signature of weathering inputs to the Labrador Sea

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A B S T R A C T

Understanding the history of continental ice-sheet growth on North America, and in particular that of the Laurentide Ice Sheet (LIS), is important for palaeoclimate and sea-level reconstructions. Information on ice-sheet extent pre-dating the Last Glacial Maximum (LGM) is heavily reliant, though, on the outputs of numerical models underpinned by scant geological data. Important aspects of LIS history that remain unresolved include the timing of its collapse during Termination 2, the first time that it expanded significantly during the Last Glacial Cycle, and whether its volume was significantly reduced during marine isotope stage (MIS) 3. To address these issues and more, we present authigenic iron-manganese (Fe–Mn) oxyhydroxide-derived high-resolution records of Pb isotope data and associated rare earth element profiles for samples spanning the past ~130 kyr from northwest North Atlantic Labrador Sea, IODP Site U1302/3. We use these new data to track chemical weathering intensity and solute flux to the Labrador Sea associated with LIS extent on the adjacent highly radiogenic (high Pb isotope composition) North American Superior Province (SP) craton since the Penultimate Glacial Maximum (PGM). Our new records show that relatively high (radiogenic) values characterise warm marine isotope stages (MIS) 5, 3 and 1 and the lowest (most unradiogenic) values occurred during cold stages MIS 6, 4 and 2. The radiogenic Pb isotope excursion associated with Termination 2 is short-lived relative to the one documented for Termination 1, suggesting that LIS retreat during the PGM was relatively fast compared to the LGM and that its collapse during the last interglacial occurred ~125 ka. Highly radiogenic inputs to the Labrador Sea during MIS 5d–a, ~116–71 ka, most likely reflect a spin-up in Labrador Current vigour, incipient glaciation and renewed glacial erosion of high grounds of the eastern SP craton by localised wet-based ice-caps. A large decrease in Pb isotope values towards unradiogenic LGM-like compositions between ~75–65 ka across the MIS 5/4 transition likely reflects a slow-down in Labrador Current vigour, an increase in subaerial deposition of aeolian dust and a significant advance of the LIS across Hudson Bay caused a strong reduction or even abandonment of Pb sourcing from the SP. The relatively radiogenic Pb isotope composition of bottom-waters bathing our study site during MIS 3, 57–29 ka, is unlikely to support a recently proposed major reduction in LIS extent for this time. Instead, we argue these values are better explained by southern Greenland Ice Sheet retreat, increased chemical weathering of the Ketelidian Mobile Belt and subsequent Pb runoff from Greenland.

1. Introduction

The growth and decay of continental ice sheets has resulted in major fluctuations in global sea-level during the Quaternary (Rohling et al., 2014). Accurate reconstructions of past ice-sheet
extent are necessary to understand the cause and timing of glacial terminations, rates of sea-level change and ice-sheet-ocean-atmosphere interactions. Knowledge of where ice-sheets grew during past glacials facilitates a firmer understanding of factors that drive their mass balance (e.g., astronomical cycles, atmospheric pCO2 and associated feedbacks) and is also required to correct interglacial sea-level reconstructions accurately for glacial isostatic adjustment (GIA; so crustal loading and unloading) when ice-sheets grow or melt (e.g., Dendy et al., 2017). We yet have only a rudimentary understanding of northern hemisphere continental ice-sheet histories prior to the Last Glacial Maximum (LGM), especially for the largest of these ice sheets, the North American Ice Sheet complex.

It has been shown that GIA-based corrections of Last Interglacial (MIS 5e, −129−116 ka) sea-level records are highly sensitive to uncertainties in the distribution of Northern Hemisphere ice-sheets during preceding glacial maxima to the order of up to ~5 m (Dendy et al., 2017; Dwyer et al., 2021), a figure equivalent to the lower-end estimate and range of sea-level rise reported for this interglacial (~6.6 ± 2 m; Kopp et al., 2013; Hibbert et al., 2016). Yet GIA models used to correct MIS 5e records for such factors mainly assume LGM ice-sheet distributions and volumes for the Northern Hemisphere during the Last Glacial Maximum (LGM) and older glacial maxima (Dendy et al., 2017). Multiple lines of indirect evidence suggest, however, that the North American Ice Sheet complex during the PGM may have been considerably smaller than its LGM counterpart (e.g., Potter and Lambeck, 2003; Svendsen et al., 2004; Rabineau et al., 2006; Colleoni et al., 2011, 2016; Wainer et al., 2017; Rohling et al., 2017). Uncertainty in our knowledge of the North American Ice Sheet complex history is not just restricted to the PGM and older times. Our understanding, for instance, of when a large North American Ice Sheet Complex first grew during the Last Glacial Cycle (LGC) is poorly constrained by empirical data (compare Batchelor et al. (2019) to Pico et al. (2017)).

Considerable debate also exists about the extent of North American Ice Sheet complex reduction during the LGC warm interstadial MIS 3, −29−57 ka (e.g., Pico et al., 2017; Dalton et al., 2019; Miller and Andrews, 2019; Gowen et al., 2021; Kerr et al., 2021). Uncertainty in MIS 3 ice-sheet extent is rooted in the sparsity of well dated terrestrial glaciomorphological evidence of pre-LGM age (e.g., Miller and Andrews, 2019). Much of our knowledge of the history of the North American Ice Sheet complex prior to the LGM is thus reliant on either numerical modelling experiments (e.g., Lambeck et al., 2006, 2010, 2017; Abe-Ouchi et al., 2013; de Boer et al., 2014; Colleoni et al., 2016) or indirect sedimentological evidence from glacially-derived terrace deposits sedimented in marine ice-proximal Quaternary settings (e.g., Hemming, 2004; Hodell et al., 2008; Bailey et al., 2013; Lang et al., 2014). One proxy that has great potential to reveal important new insights into these issues is the Pb isotope composition of oceanic waters recorded in the Fe–Mn oxyhydroxide fraction of marine sediments. Owing to the particle-reactive nature and short seawater residence time of Pb (20–30 years), the Pb isotope compositions of authigenic Fe–Mn oxyhydroxides have been proposed to track regional weathering intensity and solute flux associated with glacial extent on adjacent continental landmasses (e.g., Crocket et al., 2012; Foster and Vance, 2006; Gutjahr et al., 2009; Kurzweil et al., 2010; Süße et al., 2022). A short Pb isotope record from Integrated Ocean Drilling Program (IODP) Site U1302/3, recovered from North Atlantic Orphan Knoll proximal to northeast America, is most unradiogenic (so is characterised by its lowest 206Pb/204Pb, 207Pb/204Pb and 208Pb/204Pb) during the LGM and becomes more radiogenic (so increases) in response to increased continental chemical weathering of highly reactive glacial debris as the Laurentide Ice Sheet (LIS) retreated in the Hudson Bay region of northeastern North America during the last deglacial (Crocket et al., 2012). Yet, this record, and other available North America-proximal Pb isotope Fe–Mn oxyhydroxide datasets (e.g., Gutjahr et al., 2009; Kurzweil et al., 2010; Süße et al., 2022) are currently temporally limited to the past ~37 kyr.

To improve our understanding of LIS extent over the past ~130 kyr, we present new authigenic Fe–Mn oxyhydroxide-derived Pb isotope and rare earth element profiles of 87 samples from IODP Site U1302/3 reaching back to MIS 6. We examine the history of change in this new Pb isotope record and discuss the significance of its variability over this time in terms of LIS evolution.

2. Background

2.1. How are changes in Fe–Mn oxyhydroxide-derived Pb isotope ratios from site U1302/3 related to changes in ice-sheet extent on North America?

The Pb isotope composition of seawater bathing Orphan Knoll in the geological past was controlled by input from three main sources — the Mid Atlantic Ridge, the adjacent continents through chemical weathering and runoff and dust generation and its subaerial deposition in the Labrador Sea (Frank, 2002; Klemm et al., 2007). Given the high particle reactivity of Pb (Henderson and Maier-Reimer, 2002), an influence from chemical weathering and runoff of continental landmasses further afield than North America and Greenland seems unlikely for this region of the North Atlantic Ocean.

The Pb isotope composition of seawater can be determined by analysing the authigenic fraction of marine sediments. Lead has four naturally occurring isotopes. Three of these (206Pb, 207Pb, 208Pb) are the radiogenic daughter products of 235U (t1/2 = 7.04 Ma; 207Pb), 235U (t1/2 = 7.04 Ma; 207Pb), or 232Th (t1/2 = 14 Ga; 208Pb) respectively, while the other one (204Pb) is a stable isotope. In our study we focused on using 206Pb/204Pb, 207Pb/204Pb and 208Pb/204Pb ratios to track the Pb isotope composition of seawater bathing our study site. Due to the origin of the numerator isotope from U and Th decay, relatively high values for these ratios are described as being radiogenic, whereas low isotopic ratios are described as unradiogenic. Crocket et al. (2012) proposed that the relatively unradiogenic Pb isotope values bathing U1302/3 during LGM (~19.2 206Pb/204Pb) predominantly reflect a reduction in North American chemical weathering and runoff under a spatially extensive LIS and a dominance of a dust and Mid-Atlantic Ridge Pb isotope signal. They further argue that the subsequent trend towards more radiogenic values from ~19 ka records increased continental weathering intensity on these landmasses as well as increased runoff during Terminal 1 as these ice sheets retreated under a warming climate. The peak radiogenic values obtained during the early Holocene (~8 ka, −20.3 206Pb/204Pb) are attributed by these authors to incongruent weathering of highly reactive weakly chemically altered glacial debris exposed following ice-sheet retreat. In this model, the delay towards unradiogenic values of −19.6 206Pb/204Pb from −7 ka is suggested to reflect a reduction in weathering intensity and release of unradiogenic Pb as the exposure age of this glacial debris increased and its supply was diminished (see also Harlavan et al., 1998; Foster and Vance, 2006). Crocket et al. (2012) argued that incongruent weathering processes must have a major control on the Pb isotopic composition of seawater bathing U1302/3 over the past ~37 ka because the values recorded are much too radiogenic to be controlled by regional variations in the chemical weathering of continental crust. This assertion is based on two observations: (1) the Pb isotope composition of U1302/3 detrital sediments, which may provide a well-mixed measure of the bulk composition of the potential...
weathering source(s) from the adjacent continents, is much less radiogenic (~16–17 206Pb/204Pb) than the seawater signal (~19–21 206Pb/204Pb); and (2) a lack of co-variance exists between authigenic and detrital Pb isotope ratios at this site (Croquet et al., 2012).

Blaser et al. (2020) suggest, however, that the Pb isotope composition of detrital sediments at U1302/3 is much less radiogenic than the authigenic record because outside of Heinrich (H) events the authigenic radiogenic Pb signal recorded at Orphan Knoll must be dominated by dissolved and colloidal transport and subsequent scavenging into deep water and not terrigenous inputs. Recent laboratory experiments and observations from an enclosed Alpine lake challenge the notion that incongruent weathering can account for highly radiogenic Pb isotope ratios reported for U1302/3 during Termination (T) 1 and the early Holocene (see Daußmann et al., 2019; Sülke et al., 2019). Any incongruent weathering signal may take tens of thousands of years to decay (Harlawan et al., 1998; Sülke et al., 2019), which does not match the timescale of change observed during the Holocene at Orphan Knoll (~5 kyr) and elsewhere (e.g., Sülke et al., 2019). Instead, Blaser et al. (2020) argue that changing provenance may be a key factor. In other words, the Pb isotope signal at Orphan Knoll could also be generated by continental runoff transporting a congruent chemical weathering signal to the Labrador Sea. This is consistent with their reinterpretation of the whole-rock Pb isotope data. By the same token, it is not clear whether chemical weathering of the Abitibi Greenstone Belt alone could be solely responsible for the most radiogenic early Holocene Pb isotope values reported by Croquet et al. (2012, for our study site. This is because the drainage basins that sample this region of the SP (and host the Nottaway, Hurricana, Moose and Albany rivers of Quebec and Ontario) are responsible for only ~22% of modern annual freshwater river discharge to Hudson Bay (Déry et al., 2011). Yet over 60% of river discharge into Hudson Bay comes from catchment areas that exclusively drain the SP (Déry et al., 2011). It is possible that SP runoff took a different route to the Labrador Sea during the last deglaciation. SP runoff into Hudson Bay during T1 would have been restricted by the Hudson Bay Ice Saddle until ~8.1 ka (Dalton et al., 2020). Alternatively, isostatic rebound of North America during T1 could have altered geomorphologic gradients and re-routed SP runoff more directly eastwards into the Labrador Sea. A better appreciation of the magnitude of the role that changes in Pb provenance plays in setting the Pb isotope composition of bottom-waters bathing Orphan Knoll awaits the generation of whole-rock Pb isotope data from a wider region of the SP than is currently available, especially for its greenstone-granite belts with oceanic affinity and similar ore-deposit compositions to the Abitibi Subprovince (see those sectors highlighted green in Fig. S1). In the meanwhile, we discuss the potential role of both Pb provenance and incongruent chemical weathering in driving any radiogenic trends in our datasets.

2.3. Penultimate Glacial Maximum and Last Glacial Cycle ice sheet reconstructions of the Laurentide Ice Sheet

Global sea-level change is a primary indicator of global ice volume and is reasonably well constrained for the past ~130 kyr by data from fossil corals (e.g., Deschamps et al., 2012; Hibbert et al., 2016), submerged speleothems (e.g., Bard et al., 2002; Dorale et al., 2010; Antonioli et al., 2021), planktic foraminiferal 818O from restricted basins (e.g., Grant et al., 2014) and benthic foraminiferal 818O (e.g., Elderfield et al., 2012), albeit with well-known caveats (e.g., Skinner and Shackleton, 2005; Lisiecki and Raymo, 2009). The history of the individual continental ice-sheets that determines the changes in global sea-level that these records monitor is, however, much less well constrained.
The LIS was the largest continental ice-sheet in the northern hemisphere at the LGM. Its spatial extent at that time is widely documented by glaciomorphological evidence (e.g., Dyke et al., 2002) and inverse modelling (e.g., Lambeck et al., 2014). Our understanding of the evolution of its spatial extent prior to the LGM is, nevertheless, rudimentary (Ehlers et al., 2011). In this study, we focus on reconstructing LIS history prior to the LGM and as far back as MIS 6. Limited terrestrial glaciomorphological evidence exists for LIS extent for this time because any evidence would likely have been destroyed during its Last Glacial advance and what remains is challenging to date. Instead, most inferences of LIS evolution prior to the LGM are based on numerical models underpinned, where available, by scant empirical data (e.g., Colleoni et al., 2016; Stokes et al., 2012). This combined approach forms the basis of a recent synthesis by Batchelor et al. (2019) that arguably provides our current best constraints on pre-LGM ice sheet configurations in terms of a 'minimum', 'maximum' and 'best-estimate' for LIS extents for all isotage stages. Here we test these estimates for the past ~130 kyr, alongside insights provided by selected subsequently published research (e.g., Dalton et al., 2019; Kerr et al., 2021) using our new Pb isotope data (Figs. 3 and 4). We focus on reconstructing chemical weathering and runoff flux of North American bedrock adjacent to Hudson Bay because they underly a key sector of the LIS that is home to the biggest ice-stream of the Last Glacial (the Hudson Bay Ice Stream) and freshwater runoff from these landmasses represents the dominant source today of dissolved Pb that is ultimately routed to our study site via the Labrador Current.

### Table 1

<table>
<thead>
<tr>
<th>Regional source end members</th>
<th>Variable</th>
<th>$^{206}$Pb/$^{204}$Pb</th>
<th>$^{207}$Pb/$^{204}$Pb</th>
<th>$^{208}$Pb/$^{204}$Pb</th>
<th>Data Sources</th>
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<tr>
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<td>(Greenstone Belt)</td>
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<td>40.684</td>
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<tr>
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<td>Median</td>
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<td>15.572</td>
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<tr>
<td>(Greenstone Belt)</td>
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<td>40.357</td>
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<tr>
<td></td>
<td>Mean</td>
<td>18.464</td>
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<td>42.165</td>
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<td></td>
<td>Median</td>
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<td>15.334</td>
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<td>15.303</td>
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<td>35.517</td>
<td>12–14</td>
</tr>
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<td>15.078</td>
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<td>15.483</td>
<td>37.526</td>
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<td>14.475</td>
<td>35.584</td>
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<td>Nargssugtualiq Mobile Belt</td>
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<td>Mean</td>
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<td>38.295</td>
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<td>17.538</td>
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<td>North American aeolian dust</td>
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<td>18.865</td>
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<td>Median</td>
<td>18.880</td>
<td>15.655</td>
<td>39.053</td>
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</table>

Here, we follow Innocent et al. (1997) and Fagel et al. (1999; 2002, 2004, 2011) in using the mean and median of whole-rock data to define the Pb isotope composition of regional source end-members (see Fig. 2 and Figs. 54–16). This approach provides us with two estimates of the average weathering Pb isotope signal of each source based on the available whole-rock data. The degree of disparity between the mean and the median can be used as a first-order estimate of uncertainty in our understanding of each end-member’s average Pb isotope composition. The true average composition for each source likely falls somewhere between these two parameters. Data sources: (1) Gariepy and Allegre (1985); (2) Vervoort et al. (1993); (3) Smith (1988); (4) Thorpe (2008); (5) Wu et al. (2016); (6) Foland (1982); (7) Richardson et al. (2005); (8) Stevenson et al. (1999); (9) Doe and Delevaux (1980); (10) Peterson et al. (1994); (11) Thorpe (1982); (12) Baadsgaard et al. (1979); (13) Schiøtte et al. (1993); (14) Ashwal et al. (1986); (15) Scharer (1991); (16) Arcuri and Dickin (2018); (17) Sinha et al. (1996); (18) Kalsbeek and Taylor (1985); (19) Taylor and Upton (1993); (20) Moorabth et al. (1981); (21) Taylor et al. (1992); (22) Taylor et al. (1984); (23) Taylor et al. (1980); (24) Baadsgaard et al. (1986); (25) Kalsbeek et al. (1984); (26) Kalsbeek et al. (1988); (27) Kalsbeek et al. (1987); (28) Kalsbeek et al. (1993); (29) Hansen and Frederichsen (1989); (30) Thrane et al. (2004); (31) Jensen (1994); (32) Ellam and Stuart (2000); (33) Andreassen et al. (2004); (34) Barker et al. (2006); (35) Hansen and Nielsen (1999); (36) Farmer et al. (2003); (37) Saunders et al. (1999); (38) Holm (1988); (39) Jardine et al. (2021); (40) Aleinikoff et al. (2009).
3. Methods

3.1. Site description, sampling and age model

IODP Site U1302 (50°10’N, 45°38.3’W, water depth: 3560 m) and Site U1303 (50°12.4’N, 45°41.2’W, water depth: 3520 m) are located on the SE flank of Orphan Knoll within the Labrador Sea, proximal to the coast of Newfoundland (Channell et al., 2006). They are positioned on a rise between two canyons that help to protect these sites from debrite and turbidite deposition by funneling gravity flows down slope (Aksu and Hiscott, 1992). They are only ~3 nautical miles apart and have remarkably similar sedimentology (Expedition 303 Scientists et al., 2006). The similarity of strata between the two sites permitted the creation of a continuously spliced composite record using both sites based on correlation of physical property data down to 104 m composite depth (mcd; Expedition 303 Scientists et al., 2006).

To extend the previously published record of the Pb isotopic composition of the Fe–Mn oxyhydroxide fraction of sediments deposited at Site U1302/3 spanning the last ~37 kyr (Crocket et al., 2012) back to ~130 ka (MIS 3–6), 5-cc samples were taken every 10–50 cm (83 in total) between 0.45 and 23.38 mcd. The same samples were also used to determine the elemental concentrations of the seawater derived Fe–Mn oxyhydroxide fraction. A subset of these samples (n = 18) were used to determine the Pb isotope composition and elemental composition of the detrital fraction. A further four samples were taken between 0.45 and 4.71 mcd to evaluate the reproducibility of our new data with those reported by Crocket et al. (2012). Highly radiogenic Pb isotope excursions in authigenic Fe–Mn oxyhydroxide data derived from detrital carbonate (Heinrich) layers deposited at Site U1302/3 are suggested to represent exceptionally high inputs of detrital particles containing a “pre-formed” coating sourced from Hudson Bay during destabilisation of the LIS (Crocket et al., 2012). Not all past episodes of enhanced detrital inputs to the Labrador Sea are expressed in the Site U1302/3 stratigraphy by clear sedimentological layers. Because we are only interested in the evolution of seawater Pb isotope compositions in the Labrador Sea over the past ~130 kyr, we therefore used the XRF-derived detrital-layer stratigraphy for Site U1302/3 (Channell et al., 2012) to minimise sampling the pre-formed signal of the detrital carbonate layers.

The age model for our record was generated by Channell et al. (2012), based on matching relative palaeointensity (RPI) and δ18O data from Neogloboquadrina pachyderma (sin.) to the PSIO-1500 RPI stack (Channell et al., 2009) and the LR04 benthic δ18O stack (Lisiecki and Raymo, 2005). The detrital layer stratigraphy for our study site (Channell et al., 2012) also permits confident correlation of our datasets to those of other North Atlantic marine cores where Heinrich event iceberg rafting events have been identified.

3.2. Pb isotope analysis

All sample processing and laboratory analyses were conducted at the School of Ocean and Earth Science, University of...
Based on a whole rock data compilation by Fagel et al. (2002), supplemented by isotope composition. (For interpretation of the references to colour in this figure legend, the reader is referred to the Web version of this article.)

**Fig. 2.** The Pb isotope composition of cratonic bedrock adjacent to Orphan Knoll Site U1302/3: (A) $^{206}\text{Pb}/^{204}\text{Pb}$ versus $^{207}\text{Pb}/^{204}\text{Pb}$ and (B) $^{206}\text{Pb}/^{204}\text{Pb}$ versus $^{208}\text{Pb}/^{204}\text{Pb}$.

Based on a whole rock data compilation by Fagel et al. (2002), supplemented by isotope composition. (For interpretation of the references to colour in this figure legend, the reader is referred to the Web version of this article.)

Southampton. We extracted Pb from the authigenic Fe–Mn oxyhydroxide fraction in sediments of each sample closely following Blaser et al. (2016). Briefly, between 25 and 30 mg of dried, homogenised sediment was washed with ultra-pure water (18.2 MΩ) for 20 min on a vertical tube rotator, centrifuged at 3200 rpm for 7 min and the solution pipetted into vials and prepared for analysis.

To ensure 200 ng of Pb was obtained for column chemistry, an aliquot of each leached solution was screened on a Thermo Scientific X-Series 2 ICP-MS. Based on the Pb concentration obtained through the above screening, an aliquot of leached solution was evaporated to dryness in Teflon pots, followed by the addition of 0.5 ml concentrated HNO$_3$ and left overnight on a hot plate at 130 °C to react. The samples were left to evaporate to dryness followed by the addition of 0.5 ml 0.5 M HBr and left on a hot plate at 130 °C for at least 1 h. The Pb fraction was separated from the matrix using Eichrom AG1-X8 200–400 mesh anion exchange resin (Strelow, 1978). The resultant aliquots were screened on a Thermo Scientific X-Series 2 ICP-MS to measure the Pb concentration to allow accurate spiking for isotope analysis.

To confirm that the above procedure was successful at extracting a seawater Pb signal, 30 of the same samples spanning the record were analysed for detrital (residual fraction) Pb isotopes. Samples were exposed to a second HH-leach with 10 x solution strength for 24 h on a vertical tube rotator to ensure complete removal of Fe–Mn oxyhydroxides following Gutjahr et al. (2007). The samples were then centrifuged, decanted, washed in Milli-Q water and decanted again. After drying, 100 mg of homogenised sediment was weighed into Teflon pots and subject to 3 ml aqua regia overnight to remove organic compounds, evaporated to dryness and redissolved in concentrated HNO$_3$. The remaining silicates were digested in 2 ml of HF for 48 h at 130 °C. Finally, HClO$_4$ was added and left overnight at 130 °C. Once dried, a few drops of 6 M HCl were added for 24 h at 130 °C dried and repeated. Stock solutions were comprised of the sample, ~10 ml 6 M HCl and Milli-Q. These detrital fraction solutions were prepared for column chemistry following the same procedure as outlined above for the Fe–Mn oxyhydroxide fraction solutions.

Pb isotope ratios were measured using a Thermo Neptune MC-ICP-MS, after the samples were diluted with 3% HNO$_3$ to obtain ~20 ng/ml of Pb. A double spike run for each sample was used to correct for instrumental mass fractionation where the $^{206}\text{Pb}/^{204}\text{Pb}_{\text{sample}}$/$^{206}\text{Pb}/^{204}\text{Pb}_{\text{spike}}$ was 0.09 ± 0.03. Procedural blanks for Fe–Mn oxyhydroxide and detrital fraction samples averaged 0.028 ng (n = 14) and 0.001 ng Pb (n = 1), respectively with samples being blank-corrected. Blanks were treated in the same way as sediment samples to determine possible contamination from reagents and general handling. Authigenic Pb masses range from ~36 to 198 ng. Reproducibility of Pb isotope ratios was estimated based on 17 measurements of the standard SRM NBS 981 run from 2018 to 2020. This analytical period yielded average ratios and uncertainties at 2σ (relative to the values of Baker et al. (2004) as shown in parentheses): $^{206}\text{Pb}/^{204}\text{Pb} = 16.9424 ± 0.00044$ (16.942 ± 0.00006), $^{207}\text{Pb}/^{204}\text{Pb} = 15.4982 ± 0.00097$ (15.501 ± 0.00006) and $^{208}\text{Pb}/^{204}\text{Pb} = 36.7182 ± 0.00276$ (36.730 ± 0.00019). All accuracies are <1%. To monitor reproducibility of Pb isotope ratios further, an internal replicate of an authigenic Fe–Mn oxyhydroxide sample from IODP Site U1302/3 was measured in every run (n = 5), producing average ratios of $^{206}\text{Pb}/^{204}\text{Pb} = 19.1946 ± 0.000416$ (2σ; variance = 0.0057 ± 0.0008), $^{207}\text{Pb}/^{204}\text{Pb} = 15.6341 ± 0.000859$ (2σ; variance = 0.0042 ± 0.0001) and $^{208}\text{Pb}/^{204}\text{Pb} = 35.2377 ± 0.00205$ (2σ; variance = 0.000071 ± 0.00008).

3.3. Rare and trace elements analysis

Trace element concentrations of the Fe–Mn oxyhydroxide
Fraction stock solutions were also used to evaluate the fidelity of our seawater Pb isotope data. To generate these data, 100 μl aliquots of the above-mentioned stock solutions were dried in Teflon pots, then concentrated HNO₃ was added to remove HH-leach chemicals. The samples were dried and dissolved in 3% HNO₃ spiked with In (5 ppb), Re (5 ppb) and Be (20 ppb) to achieve a 100 x dilution. Analyses were conducted on the Thermo Scientific X-Series 2 ICP-MS. Measurements were calibrated using a suite of international rock standards: JB-1a, JB-1b, BIR-1, BHVO-2, JB-3, JB-2, BCR-2, AGV-2 that were diluted with 3% HNO₃ spiked with In (5 ppb), Re (5 ppb) and Be (20 ppb) to achieve an ~4000 x dilution for the trace elements. An additional set of standards were run with

![Fig. 3. Reconstructions of Laurentide Ice Sheet and Greenland Ice Sheet extents following Batchelor et al. (2019). Maximum (black line), minimum (red line) and best-estimate (blue) ice-sheet extents for: (A) marine isotope stage (MIS) 6 (132–140 ka); (B) MIS 5d/b (108–117; 86–92 ka, respectively); (C) MIS 5c/a (92–108; 72–86 ka); (D) MIS 4 (58–72 ka); (E) MIS 3 (29–57 ka) where minimum estimate is at peak warmth (40–45 ka; red line) and maximum estimate achieved following a period of LIS growth (30 ka; black line); (F) Last Glacial Maximum (26–19 ka). Black cross-hatched area denotes geographical extent of Superior Province craton (Montion et al., 2018). Red stars highlight location of IODP Site U1302/3 on Orphan Knoll. (For interpretation of the references to colour in this figure legend, the reader is referred to the Web version of this article.)](image-url)
Hudson Bay Lowlands used by Dalton et al. (2016, 2017; 2019) to infer a marine incursion into this area (and its deglaciation) between 52 ka. R.L. Parker, G.L. Foster, M. Gutjahr et al. Quaternary Science Reviews 287 (2022) 107564

REE bulge: MREE

REE slope: MREE

Middle REE:

Heavy REE:

~80,000 × dilution for Al and Fe. In-house reference materials BAS206 and BRR-1 and international reference material JA-2 were also run after the standards. Long term accuracy of JA-2 relative to the reference value (Jochum et al., 2016) is better than 5% except for Cu and Pb which are 6% and 7%, respectively.

Rare earth element (REE) concentrations of the samples were expressed relative to the REE profile of the Post-Archean Australian Shale (PAAS), which is a good approximation of the upper continental crust (Taylor and McLennan, 1985). To characterise REE patterns, ratios of PAAS-normalised concentrations were used following Martin et al. (2010) and Blaser et al. (2016, 2019) and shown in Eqs. (1)–(6):

Light REE: \[ LREE = \frac{La_n + Pr_n + Nd_n}{Yb_n + Lu_n} \]

Heavy REE: \[ HREE = \frac{Er_n + Yb_n + Lu_n}{Yb_n + Lu_n} \]

Middle REE: \[ MREE = \frac{Gd_n + Tb_n + Dy_n}{Yb_n + Lu_n} \]

REE slope: \[ HREE/LREE \]

REE bulge: \[ MREE/MREE^* = \frac{MREE}{0.5(HREE + LREE)} \]

where \( REE_n = \frac{REE_{sample}}{REE_{PAAS}} \)

4. Results and discussion

4.1. New LGC Pb isotope data from site U1302/3

In Fig. 5 we integrate our new and published data to present the first high-resolution LGC records of authigenic Fe–Mn oxyhydroxide-derived Pb isotopes from the subpolar North Atlantic that spans both the last (T1) and penultimate (T2) deglaciations. Comparable values, where overlap exists (for the LGM to Holocene), between our new data from Orphan Knoll Site U1302/3 and those previously published by Crocket et al. (2012) for this site, confirm the validity of splicing these two records together (see overlap of filled and white circles between 37 and 0 ka in Fig. 5c–e).

The new longer record from Site U1302/3 confirms the suggestion of Crocket et al. (2012) that the U1302/3 authigenic signal of sediments deposited at our study site is characterised by short-lived highly radiogenic excursions during the deposition of Heinrich-like layers in the Labrador Sea (see Pb data within green and light yellow vertical bars in Fig. 5c–e). However, on orbital timescales, Pb isotope ratios typically vary between ~19 and 21.5 \( 206Pb/204Pb \) ratios across T2 is comparable to its evolution across T1 (compare Pb isotope zone \( \delta \) in Fig. 5c–e) but return to ratios closest to late Holocene compositions \( \sim 19.6 \) \( 206Pb/204Pb \); Pb isotope zone \( \delta \) in Fig. 5c–e; also see Fig. S2). LGM-like values are first recorded during the LGC in MIS 4 (Pb isotope zone \( \epsilon \) in Fig. 5c–e) but return to ratios closest to late Holocene compositions \( \sim 19.6 \) \( 206Pb/204Pb \) during MIS 3 (Pb isotope zone \( \zeta \) in Fig. 5c–e). By contrast, the detrital fraction Pb isotope record shows different absolute ratios (e.g., ~16–18 for \( 206Pb/204Pb \) and contrasting temporal trends to that recorded by the authigenic records (compare Fig. 5c–e to 5h; also see Fig. S3).

4.2. Fidelity of the authigenic Fe–Mn signal from Site U1302/3 as a record of Labrador Sea seawater Pb isotope composition

To assess the integrity of the seawater Pb isotope record extracted from the authigenic Fe–Mn oxyhydroxide sediment fraction, we employ a series of tests to check for the possible release of non-seawater-derived Pb from the detrital sediment fraction. These tests are: (1) examination of detrital and authigenic Pb isotope trends for evidence of covariance, (2) evaluation of REE data to identify what material was leached (Haley et al., 2004; Martin...
Fig. 5. Pb isotope and δ18O records from IODP Site U1302/3 spanning marine isotope stage (MIS) 6–1: (A) Relative sea-level (RSL) record from the Red Sea with data points (blue crosses), 95% probability interval for the RSL data (dark grey envelope) and 95% probability interval for the probability maximum (light grey envelope; Grant et al., 2012). Also shown is a benthic foraminiferal-derived RSL record from Ocean Drilling Program Site 1123 (red line; Elderfield et al. (2012)) and the PGM-LGM difference in global sea-level (horizontal dashed lines); (B) U1302/3 planktic δ18O (black line; Hillaire-Marcel et al., 2011) and the LR04 benthic δ18O stack (blue line; Lisecki and Raymo, 2005); Pb isotope ratios of authigenic Fe–Mn oxyhydroxides: (C) 206Pb/204Pb; (D) 207Pb/204Pb; (E) 208Pb/204Pb; Pb isotope ratios of detrital sediment fraction: (F) 206Pb/204Pb; (G) 207Pb/204Pb; (H) 208Pb/204Pb ratios of Site U1302/3. “Detrital” refers to the full digestion of the authigenic-free fraction of marine sediment. Solid circles/stars = this study; open circle/stars = Crocket et al. (2012). MIS cold (warm) stages are shown by vertical light grey (white) bars. Heinrich (detrital) layers preserved in both the Labrador Sea (identified using core-scanning-derived XRF Ca/Sr ratios by Channell et al. (2012)) and subpolar North Atlantic sediments (Hemming, 2004; Hodell et al., 2008) are shown by labelled vertical light green bars. Labelled vertical light-yellow bars highlight Heinrich-like detrital layers where deposition was restricted to the Labrador Sea (Channell et al., 2012; Hodell et al., 2008). Star-shaped symbols denote Pb isotope data from U1302/3 detrital layers. Horizontal dashed lines in (C) (D) and (E) highlight Pb isotope values at U1302/3 during the late Holocene and Last Glacial Maximum. Horizontal black bars and arrows in (C) highlight intervals over which we infer enhanced supply of dissolved radiogenic weathered material from the Superior Province craton exposed following Laurentide Ice Sheet retreat in the Hudson Bay region during Terminations 2 and 1 (data symbols in CE with black outlines). Greek symbols α–ι labelled in (C) correspond to Pb isotope data zones referred to in main text. See Fig. 1 for site locations. (For interpretation of the references to colour in this figure legend, the reader is referred to the Web version of this article.)
et al., 2010) and, (3) comparison of Al/Nd, Al/Pb and Al/Th ratios of the authigenic and detrital fractions of our samples and abyssal ocean Fe–Mn crusts (Gutjahr et al., 2007).

The absence of any significant covariance between trends in the authigenic and detrital Pb isotope records (compare panels c–e to fh in Fig. 5; Fig. S3) strongly suggests that detrital sediment leaching during sample processing does not control temporal variability in our leachate record. This is confirmed by the presence of a distinct PAAS-normalised mid-REE (MREE) enrichment in our leachate samples that is not characteristic of the distribution of REE in our detrital samples (Fig. 6; Piper, 1974). This MREE enrichment is somewhat different from water column REE concentration patterns (e.g. Patton et al., 2021) and considered characteristic of marine pore waters because particulate Fe$^{3+}$ oxides are reduced and dissolved in pore waters to release REE scavenged in the water column (Haley et al., 2004). It has been shown that the REE signature of HH-extracted leachate seawater plots separately from that of the residual detrital fraction in cross plots of the REE slope (HREE/LREE) and MREE bulge (MREE/MREE$^*$) (Haley et al., 2004; Martin et al., 2010). Most of the samples analysed in this study plot in this region (Fig. 7). Yet some of our MIS 5 authigenic data (n = 18 out of 83) encroach on the detrital field (red and yellow triangles in Fig. 7a–b) consistent with incomplete removal of oxides, partial leaching of the detrital component during sediment processing (Martin et al., 2010) or potentially by pore waters in the sediment pile (i.e., in situ chemical weathering at the seafloor after deposition; Blaser et al., 2020). The presence of positive Ce anomalies in some MIS 5 authigenic data may record contributions of detritally-sourced Ce to the leachate signal (Fig. 6). It may also reflect highly elevated early-deglacial trace metal input into the Labrador Sea due to generally highly elevated chemical weathering rates following the LGM (compare with Vance et al., 2009).

These MIS 5 data are characterised by some of the most radiogenic Pb isotope values in our records (of ~21.5 206Pb/204Pb; Fig. 5c–e). The general absence of detrital H-layer deposition at U1302/3 during this interval (green/light yellow bars in Fig. 5) rules out leaching of preformed radiogenic coatings from sediments sourced from Hudson Bay for the origin of this signal (Crocket et al.,

![Fig. 6. Post-Archean Australian shale (PAAS) normalised rare earth element (REE) multi-element plots for paired authigenic Fe–Mn oxyhydroxide (solid lines) and detrital (dotted lines) samples deposited at IODP Site U1302/03 during (A) marine isotope stage (MIS) 3, (B) MIS 4, (C) MIS and (D) MIS 6.](image-url)
We cannot rule out the possibility that the detrital fraction itself was partially leached during the processing of our MIS 5 samples but consider it unlikely (see below). The relatively low MREE/MREE* ratios for many of these samples also suggest in-situ chemical weathering of sediments (i.e., they encroach on the pristine seawater signal and are significantly lower than the Al/Nd, Al/Th and Al/Pb ratios of their detrital counterparts (Fig. 9). We therefore conclude that the major trends in U1302/3 authigenic Pb isotope data are not dominated by release of Pb from sediments deposited in the Labrador Sea during the LGM.

4.3. What does the evolution of authigenic Pb isotopes at U1302/3 reveal about Laurentide Ice Sheet extent over the past ~130 kyr?

An increase in (i) physicochemical weathering intensity, (ii) incongruent weathering of highly reactive weakly-chemically altered glacial debris on North America (Crocket et al., 2012) and (iii) enhanced sourcing of weakly-weathered SP radiogenic material (Blaser et al., 2020) have been proposed to explain how the Pb isotope composition of bottom waters bathing U1302/3 can become more radiogenic following LIS retreat. Regardless of which of these mechanism prevail, our new Pb isotope records from U1302/3 provide a way to shed new light on the evolution of the spatial extent of the LIS back to T2.

4.3.1. MIS 6 and Termination 2 LIS history

The Batchelor et al. (2019) reconstruction for MIS 6 suggests Hudson Bay was glaciated throughout the PGM, but no accurately dated empirical data exist that can be used to ground-truth this suggestion. While only the oldest datapoints in our Pb isotope records were recovered from the end of MIS 6 according to our RPI-based age model, they suggest that the Pb isotope composition of seawater bathing U1302/3 just prior to Termination 2 were just as unradiogenic as during the LGM at this site (compare Pb isotope zones $z_1$ and $z_2$ in Fig. 5c–e; Fig. 10). Our new data therefore suggest that the spatial footprint of the LIS in Hudson Bay and the degree to
which it suppressed the delivery of weathered Pb from the interior of North America to Site U1302/3, at least immediately prior to the end of the PGM, were comparable to that of its LGM counterpart. The radiogenic peak in the U1302/3 record during the last deglaciation is argued either to reflect incongruent weathering of highly reactive weakly-chemically altered glacial debris exposed following ice-sheet retreat (Crocket et al., 2012) or enhanced sourcing of weakly-weathered SP radiogenic material exposed following LIS retreat (Blaser et al., 2020). The overprint of H-layer deposition during T2 (the ‘so-called’ H11) in our Pb isotope data partially masks the equivalent weathering and runoff signal for the MIS 6/5 transition (Pb isotope zone b in Fig. 5c–e; Fig. 10). There is, however, no evidence in our records that the radiogenic supply of Pb from the interior of North America continued beyond ~122 ka. This is because the Pb isotope values in all our authigenic records return to compositions observed during the late Holocene by this time (as recorded at the transition between Pb isotope zones b and γ in Fig. 5c). The restriction of H-layer deposition to the Labrador Sea during MIS 6 (Naafs et al., 2013; Obrochta et al., 2014) points towards large differences in ice-mass distribution and ice-stream dynamics between the PGM and LGM LIS in the Hudson Bay area.

The absence of detrital limestone-bearing H-layers in the central subpolar North Atlantic during MIS 6 has been attributed to a relatively thin (and therefore less voluminous) LIS that was cold-based and unable to generate H-events through surging of its Hudson Bay ice-stream (Obrochta et al., 2014). In this scenario, physical erosion of the SP terrane would have been reduced during the PGM relative to the LGM, leaving behind comparatively less-weakly weathered radiogenic material available to be chemically weathered following LIS retreat during T2. This may therefore explain the shorter time over which radiogenic values are a feature of our record during the MIS 6/5 transition (over no more than ~7 kyr) compared to the last deglaciation (~17 kyr) (compare width of horizontal black bars in Fig. 5c; Fig. 10).

The relatively short time over which radiogenic values are a feature of authigenic Pb isotope records during T2 and T1 support the idea that incongruent weathering did not play a leading role in changes in the Pb isotope composition of seawater bathing Orphan Knoll during LIS retreat. This is because it has been shown that any incongruent weathering signal should take tens of thousands of years to decay (Harlavan et al., 1998; Süfke et al., 2019), not ~3–5 kyr as we observe at U1302/3 (Fig. 5c–e). Furthermore, leaching...
experiments on granitic rock from the Swiss Alps show that increases in the $^{208}\text{Pb}^{204}\text{Pb}$ ratios of the solute products of incongruent weathering should also lag those in $^{207}\text{Pb}^{204}\text{Pb}$ because the main source of radiogenic $^{208}\text{Pb}$ (Th-bearing minerals) is more resistant to dissolution than (U-bearing) mineral sources of $^{209}\text{Pb}$ and $^{207}\text{Pb}$ (Dausmann et al., 2019; Süfke et al., 2019). Yet, all three authigenic Pb isotope ratios from U1302/3 increase at the same time during both T2 and T1 (Fig. 5c–e).

Independent reconstructions of LIS extent since the LGM highlight that while it began to retreat from ~20 ka (and seemingly first in Long Island Sound, Connecticut), a significant proportion of the SP was likely to have remained glaciated during T1 until ~12 ka (e.g., Dalton et al., 2020). If enhanced chemical weathering of SP craton was responsible for the radiogenic trends in U1302/3, the peak of radiogenic Pb supply to U1302/3 during the early Holocene is also broadly coincident with the final outburst of glacial Lake Agassiz-Ojibway south of Hudson Bay that most recent estimates suggest to have occurred between 8.7 (Dalton et al., 2020; Süfke et al., 2022) and 8.15 ka (Brouard et al., 2021). While these radiogenic spikes in Pb runoff to our study site cannot provide us with a precise understanding of the regional history of LIS retreat during T1 and T2, their respective durations may actually therefore provide us with a stratigraphic marker for the timing of collapse of the Hudson Bay Ice Saddle (or at least a likely latest possible date for it). This is argued to have occurred faster following the PGM than it did following the LGM, and in response to greater boreal summer insolation forcing (Carlson et al., 2008).

**4.3.2. MIS 5 LIS history**

The sudden return to elevated radiogenic Pb isotope values between ~115 and 75 ka during MIS 5d-a following the start of the LGC is arguably the most notable feature of our new dataset (Pb isotope zone δ in Fig. 5c–e). If a change in Pb provenance is involved, it cannot be attributed to, e.g., an increase in chemical weathering of exposed Greenland bedrock (e.g., the Ketilidian Mobile Belt) due to a diminished Greenland Ice Sheet. This is because the only circum-North Atlantic bedrock sufficiently radiogenic to explain these values is the greenstone belts of the SP (Fig. 2). Numerical ice-sheet modelling also shows that Greenland Ice Sheet regrowth following MIS 5e would have been relatively rapid (Kleman et al., 2013; Colleoni et al., 2014). Furthermore, it is challenging to explain the origin of this highly radiogenic interval by invoking continued chemical weathering (incongruent or otherwise) of weakly-chemically altered glacial debris on North America exposed following PGM LIS retreat because Pb isotope values recover to late Holocene-like values during the latter half of MIS 5e (by ~122 ka; compare Pb isotope zones γ & ι in Fig. 5c–e). This observation shows that the radiogenic weathering signal sourced from North America during T2 had already been exhausted. Instead, we consider that two mechanisms acting in concert are likely responsible for the observed radiogenic increase: (1) MIS 5 Labrador Current invigoration and, (2) incipient LIS glaciation.

The Labrador Current contributes to the transport of dissolved Pb to U1302/3 and its flow was likely reinvigorated following MIS 5e (Mao et al., 2018). The elevated radiogenic Pb isotope signal that we observe between ~115 and ~75 ka may therefore reflect a long-term increase in Labrador Current flow speed. An increase in Labrador Current vigor would supply more Pb from inner parts of the Labrador Sea that under less vigorous flow rates would not make it to U1302/3 given the particle-reactive nature and short seawater residence time of Pb. While our knowledge of Pb isotope compositions of the SP remain somewhat limited, the highly radiogenic signal preserved by our records for the MIS 5d-a may itself provide indirect evidence that SP rocks are on average highly radiogenic.

Numerical modelling by Batchelor et al. (2019) propose that the high grounds of the SP terrane (the Quebec/Labrador region) were glaciated during MIS 5. The highly radiogenic values in our records may therefore also reflect renewed glacial erosion of the SP in high
4.3.3. MIS 4 LIS history

The pronounced decline in Pb isotope values across the MIS 5a/4 transition from ~20.6 to ~19.3 $^{206}\text{Pb}/^{204}\text{Pb}$ ($\text{Pb}$ isotope zone $e$ in Fig. 5c) most likely reflects a decrease in Labrador Current vigour (Mao et al., 2018). This decline also likely reflects an increase in deposition of North American-sourced aeolian dust (Fig. 8f), which has a relatively unradiogenic source signature (Lang et al., 2014; Jardine et al., 2021, Fig. 2) and a significant reduction in the supply of weathered Pb from the SP craton as the spatial footprint of the LIS increased notably for the first time during the LGC. This decline in Pb isotope ratios is also associated with the first sea-level lowstand of the LGC (compare panels a and c-e in Fig. 5) so likely documents a significant advance of the LIS across the SP and Hudson Bay. The 75 cm (~5.2 kyr) gap in our record spanning 10.05–10.80 mcd prevents us from establishing the exact timing of LIS advance in Hudson Bay during MIS 4. Regardless, the unradiogenic trend in our Pb isotope records at this time most likely reflects the westward advance of the Quebec-Labrador dome over Hudson Bay inferred for this time from till fabrics and lineation swarms in the Hudson Bay Lowlands (Kleman et al., 2010; Gauthier et al., 2019). Using GIA modelling, Pico et al. (2017) conclude that relative high MIS 3 sea-level along the US mid-Atlantic is most consistent with a scenario where the eastern Labrador (Quebec) sector of the LIS is ice-free between ~80 and 44 ka (MIS 5a to MIS 3). Our data strongly imply, however, that the eastern LIS expanded significantly over the SP craton across the MIS 5/4 transition and that their choice of ice-sheet configuration for MIS 4 in northeast North America is likely to be incorrect.

4.3.4. MIS 3 LIS history and implications for $H$-event mechanisms

The subsequent increase in U1302/3 Pb isotope ratios across the MIS 4/3 transition (e.g., of ~0.3 in $^{206}\text{Pb}/^{204}\text{Pb}$; Pb isotope zone $e$ in Fig. 5c) shows that the onset of MIS 3 is associated with increased delivery of radiogenic Pb to the Labrador Sea relative to MIS 4, perhaps enhanced by a strengthened Labrador Current (Mao et al., 2018). Excluding H-events, the values in our $^{206}\text{Pb}/^{204}\text{Pb}$ and $^{208}\text{Pb}/^{204}\text{Pb}$ records are persistently late Holocene-like (e.g., ~19.5 in $^{206}\text{Pb}/^{204}\text{Pb}$ and ~40 in $^{208}\text{Pb}/^{204}\text{Pb}$) throughout MIS 3 (see Figs. 2 and 5) before decreasing towards an LGM unradiogenic minima (e.g., of ~19.2 $^{206}\text{Pb}/^{204}\text{Pb}$ and ~39.5 $^{208}\text{Pb}/^{204}\text{Pb}$) over ~5 kyr from ~25 ka. These relatively radiogenic MIS 3 values were also recorded in the Crocket et al. (2012) Pb isotope records for Site U1302/3 between 25 and 37 ka, and have previously been interpreted by values. (For interpretation of the references to colour in this figure legend, the reader is referred to the Web version of this article.)
Blaser et al. (2020) to mean that Hudson Bay was ice-free during this time.

Widespread deglaciation of Hudson Bay, the Hudson Bay Lowlands and the eastern Quebec-Labrador-sector of the LIS has also been hypothesized for MIS 3 by Dalton et al. (2019) based on a new stratigraphic framework for the Hudson Bay Lowlands and their synthesis and re-interpretation of late Pleistocene geochronological data largely assembled by Dyke (2002). If the spatial extent of the LIS was significantly reduced at this time, we might expect to see clear evidence of a large spike in radiogenic Pb runoff during the MIS 4/3 transition (i.e., as is recorded for T2 and T1 at U1302/3, Fig. 10) in response to an increase in chemical weathering rates inland on North America. A short-lived radiogenic spike is potentially evident in our Pb isotope records at ~61 ka (see black stars in Fig. 5c–e), but it is much smaller than those associated with our study site for the last two deglaciations. This observation may indicate LIS retreat across MIS 4/3 reflects a failed termination of sorts (see also Schaefer et al., 2015) but the hydrological cycle remained largely sluggish resulting in a diminished chemical weathering response in North America, whether incongruent or otherwise. Yet, the corresponding 207Pb/204Pb isotope ratios of deep water bathing our study site during MIS 3, which are more unradiogenic than their late Holocene counterparts (by ~0.1, Fig. 5d), may highlight that a different explanation is required.

If the LIS did not retreat markedly during the MIS 4/3 transition (as argued by, e.g., Miller and Andrews, 2019), the only non-North American-based regional source that could potentially explain the Pb isotope composition of the deep–waters bathing our study site during MIS 3 is the Ketilidian Mobile Belt and Pb runoff from Greenland. A subsequent large decrease in Pb isotope values towards LIS retreat northwards across the Superior Province during Termination 2 was relatively rapid compared to Termination 1, and in response to greater boreal summer insolation forcing. The renewed existence and runoff of radiogenic Pb to the Labrador Sea during the MIS 6/5e transition are characterised by highly unradiogenic values most likely derived from chemical weathering of the adjacent landmasses and its runoff to the Labrador Sea over the past ~130 kyr, which we argue predominantly varied as a function of Laurentide Ice Sheet extent across Hudson Bay.

We present new high-resolution records of authigenic Fe–Mn oxyhydroxide-derived Pb isotopes from Orphan Knoll IODP Site U1302/3 for the past ~130 kyr. We combine these new records with previously published Pb isotope data from this site spanning MIS 3–1 (Crockett et al., 2012) to track temporal variability in the input of Pb sourced from chemical weathering of the adjacent landmasses and its runoff to the Labrador Sea over the past ~130 kyr, which we argue predominantly varied as a function of Laurentide Ice Sheet extent in Hudson Bay.

We propose that the relatively short duration over which Fe–Mn authigenic Pb isotope compositions of bottom waters at U1302/3 during the MIS 6/5 transition are characterised by highly radiogenic values most likely reflects that Laurentide Ice Sheet retreat northwards across the Superior Province during Termination 2 was relatively rapid compared to Termination 1, and in response to greater boreal summer insolation forcing. The renewed existence and runoff of radiogenic Pb to the Labrador Sea during MIS 5d–a is recorded by the highest Pb isotope values in our data, which we argue most likely reflects invigoration of the Labrador Current and incipient glaciation and renewed glacial erosion of high grounds of the eastern Superior Province craton by wet-based localised ice-caps, most probably in the Quebec/Labrador region.

A subsequent large decrease in Pb isotope values towards unradiogenic LGM-like compositions between ~75 and 65 ka across the MIS 5/4 transition reflects an increase in North American glacial aeolian dust deposition, Labrador Current slow-down and a reduction in chemical weathering and runoff of Pb from North America due to westward advance of the Laurentide Ice Sheet across Hudson Bay. The relatively radiogenic Pb isotope composition of bottom–waters bathing our study site reported by Crocket et al. (2012) for MIS 3 has previously been argued to reflect that Laurentide Ice Sheet extent in Hudson Bay was significantly reduced at this time compared to the Last Glacial Maximum (Blaser et al., 2019). The relatively unradiogenic composition of authigenic 207Pb/204Pb ratios for our study site during MIS 3 (~0.1 lower) relative to the late Holocene highlights, however, that this interpretation may be incorrect. Instead, we propose that Laurentide Ice Sheet extent in Hudson Bay, and the Pb composition of runoff from this region of North America into the Labrador Sea, may not have changed significantly during MIS 3 compared to MIS 4 (cf. Miller and Andrews, 2019). Under this scenario, only southern Greenland Ice Sheet retreat, as recently inferred for this interval (Siedenkrantz et al., 2019), could explain the Pb isotope signature of seawater bathing U1302/3 during this time through increased chemical weathering of the Ketilidian Mobile Belt and Pb runoff from Greenland.

Data availability

The new IODP Site U1302/3 data presented and compiled Superior Province Pb isotope data are available in the attached Excel spreadsheets. Supplement 1 and 2 is also deposited in PANGAEA.

Author contributions

All authors contributed to the discussion of ideas presented in the manuscript. RLP acquired funding, generated the U1302/3 data and wrote the manuscript; IB designed the study, supervised RLP and co-wrote the manuscript; GLF, PAW and MG contributed to manuscript writing; MJC, AM, JAM helped to generate Pb isotopes and rare earth element data, oversaw instrumentation and helped with analysis. PAW co-supervised RLP.

Declaration of competing interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.quascirev.2022.107564.