The biogeochemical impact of glacial meltwater from Southwest Greenland

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Abstract

Biogeochemical cycling in high-latitude regions has a disproportionate impact on global nutrient budgets. Here, we introduce a holistic, multi-disciplinary framework for elucidating the influence of glacial meltwaters, shelf currents, and biological production on biogeochemical cycling in high-latitude continental margins, with a focus on the silica cycle. Our findings highlight the impact of significant glacial discharge on nutrient supply to shelf and slope waters, as well as surface and benthic production in these regions, over a range of timescales from days to thousands of years. Whilst biological uptake in fjords and strong diatom activity in coastal waters maintains low dissolved silicon concentrations in surface waters, we find important but spatially heterogeneous additions of particulates into the system, which are transported rapidly away from the shore. We expect the glacially-derived particles – together with biogenic silica tests – to be cycled rapidly through shallow sediments, resulting in a strong benthic flux of dissolved silicon. Entrainment of this benthic silicon into boundary currents may supply an important source of this key nutrient into the Labrador Sea, and is also likely to recirculate back into the deep fjords inshore. This study illustrates how geochemical and oceanographic analyses can be used together to probe further into modern nutrient cycling in this region, as well as the palaeoclimatological
approaches to investigating changes in glacial meltwater discharge through time, especially during periods of rapid climatic change in the Late Quaternary.

Keywords
Biogeochemistry, nutrients, glaciers, primary production, silica cycling

Highlights
• Novel multi-disciplinary approach to tracing freshwater and particle transport into boundary currents;
• Significant glacial inputs reach coastal waters and are transported rapidly offshore;
• Low surface water dissolved silicon concentrations maintained by diatom activity despite strong glacial and benthic supplies.
1. Introduction

The high-latitude regions are experiencing some of the most rapid environmental changes observed globally in recent decades. This is particularly true for the Arctic. Here, temperatures are rising twice as fast as the global mean, the Nordic Seas are warming at an accelerated rate (Alexeev, Walsh, Ivanov, Semenov & Smirnov, 2017), Arctic sea-ice is thinning and moving faster (Lindsay & Schweiger, 2015), and multi-year Arctic sea-ice is declining (Maslanik, Fowler, Stroeve, Drobot, Zwally et al., 2007), with significant implications for the interaction between the atmosphere and the oceans (Provost, Sennéchael, Miguet, Itkin, Rösel et al., 2017). The Greenland Ice Sheet (GrIS) is experiencing significant mass loss largely through surface melting but also via ice discharge at glacier fronts (Enderlin, Howat, Jeong, Noh, Van Angelen et al., 2014; Felikson, Bartholomais, Catania, Korsgaard, Kjær et al., 2017; van den Broeke, Box, Fettweis, Hanna, Noël et al., 2017). This melting is likely to have a global impact: the North Atlantic receives freshwater from the Nordic Seas, GrIS, and the Canadian Arctic (Bamber, Tedstone, King, Howat, Enderlin et al., 2018), which influences the density structure, circulation, and stratification in regions where deep water-masses form; these represent a major component of ocean circulation that drives global fluxes of heat and freshwater (Carmack, Yamamoto-Kawai, Haine, Bacon, Bluhm et al., 2016; Proshutinsky, Dukhovskoy, Timmermans, Krishfield & Bamber, 2015; Yang, Dixon, Myers, Bonin, Chambers et al., 2016). In addition to freshwater budgets, there has been increasing focus on the role of glaciers and ice sheets in supplying organic material and inorganic nutrients to marine systems. There are significant fluxes of nutrients in GrIS runoff both in dissolved and particulate form, including nitrogen (Wadham, Hawkins, Telling, Chandler, Alcock et al., 2016), phosphate (Hawkins, Wadham, Tranter, Telling, Bagshaw et al., 2016), dissolved silicon (Hawkins, Wadham, Benning, Hendry, Tranter et al., 2017; Meire, Meire, Struuf, Krawczyk, Arendt et al., 2016), and iron (Bhatia, Kujawinski, Das, Breier, Henderson et al., 2013; Hawkins, Wadham, Tranter, Raiswell, Benning et al., 2014). The extent to which these nutrients reach the coastal oceans, and are subsequently advected or mixed from the continental shelves into the open oceans via boundary currents, is poorly constrained and a matter of debate (Hopwood, Bacon, Arendt, Connelly & Statham, 2015). Both dissolved nutrient and particulate dynamics are significantly impacted by circulation processes (Hopwood et al., 2015) and biological activity within glacially-influenced fjords. These regions could be a significant trap of dissolved
inorganic phases (Meire, Mortensen, Meire, Juul-Pedersen, Sejr et al., 2017), and have the potential to prevent the nutrient-rich glacial waters reaching the coastal seas. Despite this possibility, distal summer phytoplankton blooms have been detected off Southwest Greenland in association with glacial melt (Arrigo, van Dijken, Castelao, Luo, Rennermalm et al., 2017) and ecosystem models indicate sensitivity to meltwater input (Oliver, Luo, Castelao, van Dijken, Mattingly et al., 2018). An understanding of how natural resources— including fisheries, bird, and mammal stocks that are essential for food and encouraging tourism—will respond in the future to increasing anthropogenic stress on a regional and global scale relies on an understanding of foundational processes of these ecosystem services, including marine biogeochemistry and the sources and sinks of essential nutrients (Berthelsen, 2014; Meire et al., 2017; Weatherdon, Magnan, Rogers, Sumaila & Cheung, 2016).

The overarching goal of the Isotope Cycling in the Labrador Sea (ICY-LAB; icylab.wordpress.com) study is to understand the cycling of nutrients in the climatically critical but understudied regions of the Labrador Sea and Greenland fjords. The approach of ICY-LAB is to capture the whole biogeochemical system in these areas of marked environmental change using carefully planned field sampling strategies, with research expeditions to coastal Greenland and the open ocean Labrador Sea. The principal dataset was collected during an oceanic expedition on the RRS Discovery (DY081, July-August 2017), with the aim to investigate the influence of glacial meltwater on nutrient cycling in the shelf seas off SW Greenland. Particular focus was placed on the silica budget and how this is framed within the oceanographic and biological processes acting on the W Greenland margin. Uniquely, we combined a range of both traditional and novel methodologies to detect and trace meltwater and glacial material from the shelf across the slope, and to investigate the biogeochemical and biological impact of these inputs. Bringing these different approaches together is essential in these margin environments to obtain a full picture of biogeochemical cycling, providing a robust insight into the system over a range of spatial and temporal scales that are otherwise challenging to resolve (Figure 1).
2. Methods and materials

2.1. Fieldwork rationale

The data presented in this paper were collected during expedition DY081, centred on the coastal shelf and slope regions off Southwest Greenland. Model results indicate that this region is influenced by surface meltwater from the Western GrIS, in addition to a significant input of freshwater delivered from the Eastern GrIS via the strong East Greenland Current (EGC) (Luo, Castelao, Rennermalm, Tedesco, Bracco et al., 2016). The study locations were selected to represent this conduit to the open ocean for glacial runoff from the Western and Southwestern GrIS, which have been the focus of recent terrestrial studies carried out in collaboration with the Bristol Glaciology Centre (Hawkings, Hatton, Hendry, de Souza, Wadham et al., 2018; Hawkings et al., 2017). Full details of the oceanographic setting are given in Appendix A.

Figure 1: Summary figure showing the multi-discipline approach taken during expedition DY081 of project ICY-LAB.
We selected the main study location for the ICY-LAB project (“Gothâb (Nuuk) Trough”, Figure 2a) to be adjacent to Nuuk, which has experienced increasing glacial run-off in recent years (Van As, Andersen, Petersen, Fettweis, Van Angelen et al., 2014). During DY081, Southern Greenland was strongly influenced by both icebergs and sea ice, but two sites (“Narsaq” and “Cape Farewell”, Figure 2a) were still selected there for providing glacial troughs that could act as direct comparisons to the further north Nuuk site further north. Orphan Knoll, on the western margin of the Labrador Sea, was selected as a distal comparison site, and for complementary palaeoclimate, biological and habitat mapping studies (Figure 2a).

The unique holistic observational approach we took is illustrated in Figure 1, and included (a) initial bathymetric mapping using acoustic methods (multibeam echosounders) to increase our understanding of the local terrain and to enable accurate planning of further
(b) characterisation of the water column structure using CTD casts and glider deployments; (c) sampling of surface and bottom waters for biogeochemical analyses using the CTD rosette and a trace metal clean towfish; (d) sampling of seafloor sediments using a megacorer (for geochemical studies of pore waters) and gravity corer (for palaeoceanographic investigations); (e) seabed observations and precision sampling for biological, palaeoceanographic and sedimentological studies using a work-class scientific ROV.

2.2. On-board methodologies and additional laboratory techniques

2.2.1. Mapping and acoustics

During DY081, multiple acoustic systems were deployed on the ship (e.g. EM122 multibeam echosounder (12 kHz); EM710 multibeam echosounder; SBP120 sub-bottom echosounder (2.5 – 6.5 kHz)). These were coordinated via a K-Sync system to avoid interference and crosstalk.

Bathymetry data were processed on-board with the Caris HIPS & SIPS software v.8, using standard settings and procedures (data import, navigation and attitude check, application of a “zero tide”, gridding into a 25 mx25 m pixel BASE surface). Backscatter data were processed with Fledermaus FMGT, again using default settings.

2.2.2. Physical oceanography

High-resolution water column studies surrounding the prominent glacial Gothâb (Nuuk) Trough were carried out utilising both a grid of Conductivity Temperature Depth (CTD) casts and the deployment of two 1000m-rated Slocum gliders (Figure 2). CTD casts were also used at the south Greenland and Orphan Knoll study sites. Hydrographic analysis enabled characterization of the water column structure in each study location, specifically to locate and quantify the freshwater inputs at the Greenland sites. Salinity was calibrated using bottle samples collected at discrete depths. After laboratory calibration of these samples, no drift corrections were required. Overall errors for temperature were 0.0006 °C (based on laboratory calibration) and 0.002 for salinity. Prior to analysis, data from these stations were gridded to a vertical and horizontal resolution of 10 m and 6 km respectively.

Vessel Mounted Acoustic Doppler Current Profilers (VMADP at 75 and 150kHz; Teledyne RD instruments) were secured onto the drop keel in surface waters near the centre-line and
beneath the RRS Discovery, and used to measure the horizontal current velocity profile.

Bottom tracking data were only collected from the 150 kHz instrument intermittently between 18th July and 24th July 2017 while close to Nuuk. In addition, downward and upward looking lowered 300 kHz ADCPs (LADCPs) were mounted on the CTD rosette. LADCP data were processed using LDEO LADCP processing software version IX_8, run on Matlab.

Full details of the other sensors attached to the CTD rosette and gliders can be found elsewhere (Hendry, 2017).

2.2.3. Biogeochemistry and chemical oceanography

2.2.3.1. Water column

Water column samples were collected using Niskin bottles attached to the CTD rosette (10L volume) and the Remotely Operated Vehicle (ROV) Isis (4 L volume), and via a trace-metal clean towfish. The towfish system comprised of a weighted titanium bodied fish lowered into the water at the stern and streamed as far from the ship as possible. When towed it was at approximately 2m depth, and water was pumped into the ship’s labs through an ultra-clean pump and tubing. Trace-metal sampling was only carried out when the ship was moving at speeds greater than 0.5 knots in order to avoid contamination from the hull of the vessel. Four stand-alone pumps (SAPs) were also deployed at key locations to collect water column particles.

Samples of seawater were collected for inorganic macronutrients, water oxygen isotope composition ($\delta^{18}O$) and carbonate chemistry parameters (pH, alkalinity), which are used for investigating freshwater input in high-latitude regions (Hendry, Pyle, Barney Butler, Cooper, Fransson et al., 2018; Meredith, Brandon, Wallace, Clarke, Leng et al., 2008; Thomas, Shadwick, Dehairs, Lansard, Mucci et al., 2011). Phytoplankton pigments were analysed on board, and compared to sensor-derived fluorescence data, to assess algal standing stocks in relation to meltwater input. Full details of sampling methods and laboratory techniques are available in Appendix B.

2.2.3.2. Diatom productivity

Biogenic silica (bSiO$_2$) production (i.e. diatom productivity) analyses were done using radioisotope $^{32}$Si as detailed in Krause, Brzezinski and Jones (2011). Briefly, samples were collected within the euphotic zone (sample depths based on light and relative to irradiance
just below the surface) and dispensed into acid-cleaned 125 mL polycarbonate bottles. 322 Bq of $^{32}\text{Si(OH)}_4$ was added to each sample, and bottles were incubated on deck in surface-seawater-cooled incubators covered with neutral density screening to mimic the depth of collection. After incubation, samples were filtered through 1.2 μm pore size polycarbonate membrane filters. Particulate $^{32}\text{Si}$ activity was quantified using a GM-25 Multicounter (Risø DTU National Laboratory, Denmark) after the samples had aged into secular equilibrium with the short-lived daughter isotope, $^{32}\text{P}$.

2.2.3.3. Radium isotopes

To investigate the fate of solutes sourced from benthic sediments or glacial meltwater, large-volume surface samples for radium (Ra) isotope analysis were collected from the trace-metal clean towfish system, both when the ship was underway (~2m water depth) and stationary (~5m water depth). A total of 200-300 L of seawater from a single sampling event were then passed through a plastic column holding MnO$_2$-coated acrylic fibre, which quantitatively binds Ra in the sample. The fibers and adsorbed Ra isotopes were then rinsed with deionized water (Milli-Q, Millipore), dried to an appropriate moisture content and loaded into a Ra Delayed Coincidence Counter (RaDeCC; Scientific Computer Instruments, USA) so as to quantify $^{223}\text{Ra}$ and $^{224}\text{Ra}$ content following the methods of Moore and Arnold (1996) and Moore (2008). Each sample was counted 4 times over ~4 months to determine the activities of excess $^{224}\text{Ra}$ and $^{223}\text{Ra}$, above the activities supported by their parent isotopes in the water column ($^{228}\text{Th}$ and $^{227}\text{Ac}$, respectively). Detector efficiencies were determined and monitored regularly at sea and in the laboratory with standards (Annett, Henley, Van Beek, Souhaut, Ganeshram et al., 2013). Final reported activities have been corrected for any decay that occurred between sample collection and analysis, activity supported by parent isotopes, detector background, and efficiency.

2.2.3.4. Sediment-water interface

High-latitude ocean margin sediments are increasingly being recognised as an important source of inorganic nutrients and key elements (Henley, Jones, Venables, Meredith, Firing et al., 2018; Kuzyk, Gobeil, Goñi & Macdonald, 2017; Sherrell, Annett, Fitzsimmons, Roccanova & Meredith, 2018). To investigate the role of sediments in these glacially-influenced shelf and slope environments, we collected pore-fluid samples at coastal
Greenland and the Labrador Sea. Short sediment cores (≤40 cm) were acquired from the study area with a mega corer. Using Rhizon filters (0.15 μm, Rhizosphere Research Products), pore water was extracted from the sediment cores and filtered into syringes, and the samples were stored under cool conditions prior to analysis. Pore water dissolved silicon concentrations were analysed on-board and post-expedition using a V-1200 Vis spectrophotometer, employing a standard molybdate-blue methodology (using Hach Lange reagents). The samples were corrected for blank and calibrated against a ten-point curve that was developed using Si standards of known concentrations. In addition to mega-cores, we also employed ROV push cores to obtain short sediment cores for pore-fluid sampling, in regions where complex bathymetry or the presence of ice-rafted material precluded the use of a megacorer.

2.2.4. Palaeoclimate

Two samples types were collected for palaeoclimate research – fossil deep-sea corals and sediment cores. ROV operations were the primary tool for benthic biological and fossil coral collections using grab or suction devices. In addition, where large fossil coral graveyards were observed, a net was used to sample fossil corals. Gravity cores were also collected to obtain long-term records of changes in meltwater flux and iceberg dynamics over the Late Quaternary, thereby providing a longer-term temporal context to the broader data set. Megacores, collected primarily for biogeochemical studies, were also subsampled to provide core-top material that could potentially replace any sections lost from the gravity cores during retrieval.

2.3. Mass balance calculations

To calculate the freshwater mass balance in the study area, the seawater samples are presumed to comprise a mixture of three source water end-members: ocean, sea ice melt and meteoric water, which is assumed to be dominated by glacial discharge. The three end-member assumption enables quantification of the freshwater fractions via the following mass balance equations (Meredith, Heywood, Dennis, Goldson, White et al., 2001):

\[
F_{ir} + F_{me} + F_{si} = 1
\]

\[
F_{ir}S_{ir} + F_{me}S_{me} + F_{si}S_{si} = S_{ms}
\]
\[ F_{ir} \delta_{ir} + F_{me} \delta_{me} + F_{si} \delta_{si} = \delta_{ms} \]

Where \( F_{ir}, F_{me}, F_{si} \) are the calculated fractions of Irminger Water, meteoric and sea ice melt respectively (Irminger Water being the chosen ocean endmember), which sum to 1 by definition. The result is clearly dependent on the exact choice of endmembers for salinity \((S_{ir}, S_{me}, S_{si})\) and \(\delta^{18}O (\delta_{ir}, \delta_{me}, \delta_{si})\) for the Irminger Water, meteoric and sea ice melt respectively. \(S_{ms}\) and \(\delta_{ms}\) are the measured salinity and \(\delta^{18}O\) of each sample.

Properties for the sea ice melt and meteoric endmembers (Table 1) were based on values reported in Dodd, Heywood, Meredith, Naveira-Garabato, Marca et al. (2009); Melling and Moore (1995); Meredith et al. (2001); and the CTD observations from the DY081 research cruise. Note that negative sea ice melt percentages reflect a net sea ice formation from the water parcel sampled.

3. Results

EM-122 multibeam swath bathymetry datasets (e.g. Figure 2) are now published on PANGAEA (doi: 10.1594/PANGAEA.892825). The full water column data from CTD profiles and bottles (hydrography, oxygen isotopes, carbonate chemistry, macronutrients e.g. Figure 3) are published on PANGAEA (doi.pangaea.de/10.1594/PANGAEA.896544). Here we present a subset of the results, focusing on characterisation of the silica cycle in the water column and sediments.

3.1. Silica cycling parameters in the water column and sediments

3.1.1. Water column macronutrients and pigments

The nearshore macronutrient concentrations were typically low (< 11 \(\mu\)M nitrate, < 0.2 \(\mu\)M nitrite, < 0.75 \(\mu\)M phosphate and < 5 \(\mu\)M DSI in the upper 50 m of the water column) reaching minima in the surface waters with lowest salinity and lowest \(\delta^{18}O\) (an example of which is given in Figure 3a, see also Appendix C). Sections of CTD macronutrient and pigment bottle data from the Nuuk grid, integrated over the top 50m, reveal consistent onshore-offshore trends (an example of which is given in Figure 3b, see also Appendix B). Integrated macronutrient concentrations decreased towards shore concurrent with an
Figure 3: A) Example cross section of macronutrient concentrations from the Nuuk grid (from CTD 8 to CTD 16), showing (from top to bottom): nitrite, nitrate, silicic acid, and phosphate (all in μM). This section crosses the shelf break and occupied the prominent glacial trough, as shown in Figure 2.

B) Example cross section of integrated (top 50 m) macronutrient and algal pigment concentrations from the Nuuk grid (from CTD 8 to CTD 10, see Figure 2). i) Integrated macronutrients; ii) Integrated pigment concentrations. Error bars show propagated errors on integration calculation (±1SD).
increase in Si:N. Integrated Chl $a$ increased towards shore, indicative that at least some of the macronutrient decrease is a result of biological uptake into biomass. However, the ratio of Chl $a$:Chl $c$ peaked at the shelf break, and then decreased again towards shore, indicating a lower diatom proportional contribution to biomass in the same locations as the lowest integrated water column DSi (Figure 3b, see also Appendix C).

### 3.1.2. Diatom productivity

Surface bSiO$_2$ production among the three sampling regions ranged from 0.05 – 0.31 µmol Si L$^{-1}$ d$^{-1}$ (Figure 4). For the most part, rates declined with depth; however, the Orphan Knoll site had subsurface maxima at the 20% isolume (~10-20 m) and base of the euphotic zone (i.e. 1% isolume, 20-50 m). The production rates of bSiO$_2$ among the Nuuk profiles were typically higher than Orphan Knoll and Southern Greenland (Figure 4); however, samples incubated in the dark (collected from below the euphotic zone) in Nuuk still had measurable production.

![Figure 4: Biogenic silica (bSiO$_2$) production during ICYLAB. A) Averaged (±1SD) gross rates of bSiO$_2$ production (µmol Si L$^{-1}$ d$^{-1}$) versus relative light depth (i.e. 100% is surface, 1% is base of the euphotic zone) among profiles within Orphan Knoll (filled circles), Nuuk (open circles), and Narsaq (filled triangles). B) Box plots of euphotic-zone integrated bSiO$_2$ production (mmol Si m$^{-2}$ d$^{-1}$) for profiles in Orphan Knoll (n = 2), Nuuk (n = 9), and Narsaq (n = 6).](image)

### 3.1.3. Sediment dissolved silicon profiles

Pore water DSi collected from the mega cores and the ROV push cores from the Labrador Sea and coastal Greenland during DY081 expedition range from 49–616 μM (Figure 5). All cores generally show an initial increase in pore water DSi with core depth, indicating...
Figure 5: Pore water dissolved silicon concentration (DSi) measurements versus mega core (MGA, solid lines) or ROV push core (dash-dot lines) depths. Results from (a) mega cores, (b) Orphan Knoll cores, (c) cores off Nuuk, and (d) cores off Narsaq and Cape Farewell. Empty circles are results from replicate push cores retrieved at certain sites. Bracketed numbers in plot legends indicate water depths of the sediment cores.
Differences in pore water dissolved silicon concentration between replicate push cores that were sampled at different time periods ($\Delta$DSiPW). $\Delta$DSiPW was calculated by subtracting push core pore water DSi sampled at the latter time period from the corresponding replicate push core pore water DSi sampled at the earlier time period. The calculation was carried out between measurements from the same or nearby ($\leq \pm 1\text{ cm}$) sediment core depths. (e) $\Delta$DSiPW versus the time difference between the sampling periods of the replicate cores ($\Delta t$), (f) $\Delta$DSiPW versus core depth.

Supply of Si to bottom waters from the sediment via dissolution. At greater depths, the rates of increase of pore water DSi with depth slow down or even reverse (Figure 5).

There is good agreement between pore water DSi results (Figure 5) observed in ROV push cores and mega cores from Orphan Knoll and South Greenland (off Narsaq and Cape Farewell). To the best of our knowledge, this is the first formal comparison of pore water DSi profiling using ROV push cores compared to megacorers, and our results provide confidence to the application of this sampling methodology in future studies, especially in settings where the use of a megacorer proves to be challenging (e.g. complex bathymetry, presence of coarse ice-rafted material). In contrast, there is larger variation in pore water DSi (Figure 5) among the sites off Nuuk (Appendix C), which could in part be related to the greater variability in sediment composition (e.g. ice-rafted debris, fossil fragments) and characteristics (e.g. grain size) observed at these sites. More importantly, the highest pore water DSi concentrations are observed (Figure 5) at the shallowest coastal site closest to Nuuk (Appendix C), which is under the influence of meltwater from glacial fjords.

Replicate ROV push cores were collected at certain sites to evaluate any consistent change in pore water DSi with time when the sediment cores were left standing onboard at ambient temperature. Results show only minor discrepancies in pore water DSi: up to $\pm 60 \mu$M when the sampling was carried out on paired replicate cores within 3 hours of each other (Figure 5); this is likely due to spatial heterogeneity in sediments and pore waters. In contrast, when pore water sampling was carried out on paired replicate cores more than 10 hours apart from each other, there are greater discrepancies with higher DSi values measured in the pore water sampled later in time (Figure 5). Our results suggest that pore water in the upper core depths might not reflect original DSi values if the sampling is carried out more than 10 hours after retrieval of the sediment core.
4. Discussion

Our multi-disciplinary framework allows a nuanced understanding of the whole silica cycle in this climatically critical region, and specifically, the impact of glacial meltwater in shelf seas and into the open ocean. We are able to address questions surrounding the amount of meltwater reaching the ocean, the mechanism by which it does so, and identify mechanisms for how the meltwater is entrained in coastal and boundary currents. For the main context of this study, we can then use this information to understand better the implications of meltwater inputs on macronutrient distributions and biological production, in particular the supply and uptake of DSi. Lastly, we use our palaeoclimate archives to interrogate past changes in meltwater supplies, which are likely to have had a major impact on nutrient cycling.

4.1. Bathymetric features and their role in shelf-water dynamics

The shipboard bathymetric and backscatter data provided a rapid insight into the geomorphology of the study regions. The bathymetry features, resulting from their glacial history, are likely to be an important influence on modern shelf-water dynamics and biogeochemical cycling. The bathymetric grids obtained off the W Greenland coast include the dedicated surveys offshore Nuuk, Narsaq and Cape Farewell, in addition to data collected along the transits (Figure 6). They illustrate a wealth of geomorphological features typical of glaciated margins, such as cross-shelf troughs, iceberg ploughmarks, gully systems, submarine canyons and submarine landslides (Dowdeswell, Canals, Jakobsson, Todd, Dowdeswell et al., 2016). Most notably off Nuuk, the inshore-deepening Gothåb (Nuuk) Trough, previously described by Ryan, Dowdeswell and Hogan (2016), is likely to be important in driving instabilities in localised circulation, influencing the mixing of melt and glacially derived material into the shelf waters. The trough harbours a number of drumlins, elongated features typical of glacial weathering, in addition to intricate patterns of iceberg ploughmarks at the shallow trough mouth (Figure 6). Systems of gullies and submarine canyons can be found at both the Gothåb and Narsaq trough-mouth fans, and along the shelf edge (Figure 6). Some of the canyons are cut up to 350 m into the continental slope, and feature steep to near-vertical walls along their flanks together with scoured channels at their floors. Offshore Narsaq, some of the submarine canyons appear to have evolved as a result of retrogressive failures cutting upslope along gullies (Figure 6).
4.2. Physical tracers of glacial meltwater inputs, mixing and advection

We can use the gridded hydrographic and geostrophic velocity fields to constrain the flow of water across the shelf break, focusing on the Gothåb (Nuuk) Trough CTD section (Figure 2b). The subsurface temperature minimum indicates the core of Polar Surface Water, whilst the subsurface temperature maximum is Irminger Water. The Irminger Water is present between 200 m and 700 m over the continental slope, with a core temperature of 4.9°C. There are clear traces of this water mass in the trough also, where warm (θ > 4°C) and saline (S > 34.7) water is observed near the bottom at CTD16 and CTD10. The Polar Surface Water core is spread across the whole section at 50-150 m, with minimum temperatures recorded at the station furthest onshore (CTD16).
Geostrophic velocities were calculated by referencing the geostrophic shear to the velocity field measured directly with a Lowered Acoustic Doppler Current Profiler (LADCP). The LADCP velocity field was de-tided, by subtracting the barotropic tide solution obtained from the Oregon State University (OSU) model (Egbert & Erofeeva, 2002). The resultant geostrophic velocity field reveals two surface intensified current cores, one on each side of the trough mouth (Figure 7). The offshore core is associated with a hydrographic front around 15 km west of the shelf break and extends down beyond 400m depth. Its offshore location is consistent with the long-term average position of the West Greenland Current (Myers, Donnelly & Ribergaard, 2009). The inshore velocity core has a less distinct hydrographic signature and therefore its origin is more uncertain at present. One possibility is that the velocity maximum is the result of topographic steering of the inshore portion of the West Greenland Current by the bathymetric trough. Alternatively, it could represent an eddy of offshore water that has detached from the boundary current. A velocity maximum was also detected in a bathymetric trough inshore of Fylla Bank (near Nuuk) in a numerical model simulation (Myers et al., 2009), but there has been no further study into the nature of this feature to date. Such strong inshore current anomalies have significant consequences for the transport of terrestrially-derived freshwater and nutrients.

In addition to the mean flows, eddies formed from the West Greenland Current transport water from the boundary current to the interior Labrador Sea, where they contribute to the process of Labrador Sea Water formation (Katsman, Spall & Pickart, 2004). Baroclinic instability is thought to be a key formation mechanism, and years of enhanced baroclinicity tend to coincide with high eddy activity (Rykova, Straneo & Bower, 2015), which transfer hydrographic anomalies into the interior. There is significant baroclinicity in both branches of the WGC in our section, implying that eddy generation may be significant. In addition, wind driven Ekman transport is likely to be a key driver in the export of shelf water across the WGC and into the Labrador Sea interior (Schulze & Frajka-Williams, 2018).

Cuny, Rhines, Niiler and Bacon (2002) have suggested that, around this location, the West Greenland Current splits into westward and northward flowing components. However, the location of the splitting, and the partitioning of the water masses involved is not well understood. There are two westward components that flow around the northern perimeter of the Labrador Sea and a northward branch that extends close to the Greenland shelf break.
Hydrographic signatures of the West Greenland Current have been reported near Greenland to the north, in the vicinity of Davis Strait and Baffin Bay (Cuny, Rhines & Kwok, 2005; Myers et al., 2009).

Figure 7: Gridded vertical sections, from top to bottom, of potential temperature (°C), salinity, and geostrophic velocity (m s⁻¹) along Gothåb (Nuuk) Trough. The direction of geostrophic flow is perpendicular to the CTD section (positive northward and negative southward). The black lines with the white boxed labels are isopycnals, which are lines of constant potential density (in kg m⁻³). The dashed vertical lines indicate the CTD stations, labelled at the top of each panel, from which these vertical sections were derived.
4.3. Chemical tracers of glacial inputs

4.3.1. Chemical tracers of meltwater

Vertical sections of sea ice melt and meteoric percentages based on seawater salinity and $\delta^{18}O$ for the Gothåb (Nuuk) Trough CTD section (Figure 8), reveals a negative offshore gradient in both sea ice melt and meteoric water percentages. Higher freshwater concentrations are found in the trough at all depths, highlighting the potential for glacially-sourced waters to reach the outer shelf and the strong boundary currents.

Freshwater mass balance calculations have, alternatively, been carried out in the High Arctic using salinity and alkalinity endmembers resulting in robust meteoric water percentage reconstructions (e.g. Hendry et al., 2018; Jones, Anderson, Jutterström & Swift, 2008). We compared the two mass balance methods using DY081 data, using both a high meteoric water alkalinity endmember typical of the riverine input to the Arctic Ocean (Jones et al., 2008), and a lower meteoric water alkalinity endmember typical of glacial meltwater (Meire, Søgaard, Mortensen, Meysman, Soetaert et al., 2015). Our comparison indicates that – irrespective of the endmember values chosen - alkalinity-derived values of meteoric water percentages are impacted by subsurface processes that show correlations with nitrite concentrations and temperature (Appendix C). Such non-conservative behaviour likely arises as a result of enhanced alkalinity flux due to water column nitrification, and/or sedimentary denitrification (Fennel, Wilkin, Previdi & Najjar, 2008; Wolf-Gladrow, Zeebe, Klaas, Körtzinger & Dickson, 2007).

Figure 8: Vertical sections of meteoric water and sea ice melt percentages along Gothåb (Nuuk) Trough. The black area is the bathymetry of the section, as measured by the ship multibeam. Red pluses indicate the bottle sampling locations, with the CTD stations annotated above.
4.3.2. Geochemical tracers of particulate flux

Radium (Ra) is produced continuously from lithogenic material by the decay of thorium (Th) and thus displays elevated concentrations near any sediment-water interface. Short-lived Ra activities did not show a clear relationship with salinity (Figure 9), but did exhibit informative regional variability. The relatively low activities around Cape Farewell indicate a lack of recent lithogenic input upstream of this region (along the eastern coast of Greenland). From Cape Farewell $^{224}$Ra$_{xs}$ increases westwards to $\sim$51$^\circ$ W, consistent with increasing cumulative sedimentary inputs into the Greenland coastal current (Figure 9C). As

Figure 9: Spatial distribution of salinity (A), $^{223}$Ra$_{xs}$ (B) $^{224}$Ra$_{xs}$ (C) and $^{228}$Th (D) from towfish samples taken at 2-5 m depth. Elongated symbols show underway samples which took ~1h to collect. For B-D, inset plots show the same isotope versus salinity in surface waters. Vertical error bars denote ±1 standard error for activities, and horizontal error bars show the standard deviation of salinity measurements taken each minute during the ~1 h sampling interval.
CTD profiles show that the shallow surface mixed layer sampled by our towfish extended to only 10-20 m, the most likely sources for $^{224}$Ra$_{XS}$ are glacial meltwater, very shallow sediments within fjords, or resuspended shelf sediments by storm-driven mixing. This signal then decreases towards Nuuk, which will reflect both dilution due to mixing and decay of the short-lived isotope (half-life 3.66 d), and also suggests minimal further lithogenic input into the surface mixed layer. The difference in activity from $\sim 9$ dpm m$^{-3}$ at $\sim 51^\circ$ W to $< 4.5$ dpm m$^{-3}$ at the Nuuk sampling stations suggests a decay time of one half-life between these sampling regions, a distance of 300-400 km, although dilution or additional inputs would also affect the measured activity and cannot be fully quantified at present. However, the maximum $^{224}$Ra$_{XS}$ activities of 8-9 dpm m$^{-3}$ above deep water off the shelf break demonstrate that this lithogenic signal is persistent, with the potential to rapidly transport other glacial and sedimentary-derived compounds far offshore.

Although this spatial pattern is not clear in the longer-lived $^{223}$Ra$_{XS}$ (Figure 9B), the lower overall activity of this isotope - as well as lower detection efficiency - lead to lower signal to noise ratios. It is also likely that regional lithologies may lead to different input patterns due to differing distribution of the $^{227}$Ac and $^{228}$Th parents.

Our analysis also quantifies $^{228}$Th, the parent of $^{224}$Ra. Due to its lithogenic origin and higher particle reactivity, trends in $^{228}$Th may be more closely associated with particulate phases than Ra, especially in our unfiltered samples. Samples collected on fibre are generally assumed to retain the majority of both particulate and dissolved Th, although in coastal waters this approach may underestimate total $^{228}$Th where particles can be flushed through the sample, and the activities presented here also include supported activity from $^{228}$Ra (the parent of $^{228}$Th) within the water column. Nevertheless, our $^{228}$Th data (Figure 9D) shows a statistically significant correlation with salinity. We therefore suggest that contrasting patterns in the $^{224}$Ra$_{XS}$ daughter may be attributed to additional inputs from fjord or shallow shelf sediments, as these inputs are more significant for dissolved species. Additional samples from inner fjord locations and depths below the mixed layer will enable us to differentiate between inputs from marine and glacial sediments.

### 4.4. Diatom productivity in glacially-influenced shelf seas

Despite the low nutrient concentrations, bSiO$_2$ production rates were significant in the meltwater influenced waters off the shelf and slope. Similar surface DSi concentrations are
observed in the open-ocean gyres (e.g. North Pacific, Brzezinski, Krause, Church, Karl, Li et al. (2011); North Atlantic, Krause, Nelson and Lomas (2010)) and typically do not support high rates of diatom productivity. Spring-season bSiO$_2$ production rates in Svalbard and the Barents Sea (Krause, Duarte, Marquez, Assmy, Fernández-Méndez et al., 2018) and at the MarineBasis Nuuk station in Godthaabsfjord (Krause, Schulz, Rowe, Dobbins, Winding et al., In review) were routinely lower than those quantified at Orphan Knoll or Nuuk. The active production below the euphotic zone suggests either recently exported diatoms (e.g. from surface waters within 1-2 days) or the presence of siliceous and active Rhizaria (Biard, Krause, Stukel & Ohman, 2018; Biard, Stemmann, Picheral, Mayot, Vandromme et al., 2016). Integrated bSiO$_2$ production rates in the euphotic zone ranged two orders of magnitude, 0.13 – 14.4 mmol Si m$^{-2}$ d$^{-1}$. Four of the six stations off Southern Greenland had integrated rates <1 mmol Si m$^{-2}$ d$^{-1}$, similar to mid-ocean gyres (Brzezinski et al., 2011), whereas all other profiles during the cruise exceeded 2 mmol Si m$^{-2}$ d$^{-1}$. These rates are on the lower end for the Southern Ocean (1-93 mmol Si m$^{-2}$ d$^{-1}$, Nelson, Treguer, Brzezinski, Leynaert and Queguiner (1995)) but are similar to Svalbard and the Barents Sea during spring (0.3–1.5 mmol Si m$^{-2}$ d$^{-1}$, Krause et al. 2018). Overall, diatom bSiO$_2$ production consumed 4% (median) – 10% (average) of the euphotic zone DSi inventory daily. These data are the first such reports for this region of the Labrador Sea and Greenland, and demonstrate a surprisingly active diatom assemblage despite low nutrients, temperature, and biomass.

### 4.5. Influence of glacial meltwaters in the open ocean: Si cycling as a case study

Although glacial meltwaters exhibit elevated concentrations of some dissolved nutrients and reactive phases (e.g. Hawkings et al., 2017), glacial fjords – the conduits between the source of these nutrients and the open ocean – are characterised by complex physical, chemical and biological processes and are highly variable in space and time (Hopwood, Carroll, Browning, Meire, Mortensen et al., 2018). Despite the highly heterogeneous nature of these environments, we can use the findings from DY081 to shed more light on the key common processes that characterise the transfer of nutrients across the land-ocean interface, which will likely vary in rates and importance between different glaciated regions. In many high-latitude regions, upwelled waters are thought to dominate the supply of nutrients to the euphotic zone supporting most of the primary production. For example, off the West Antarctic Peninsula (WAP), biological “hotspots” were thought to be fed by
upwelling Circumpolar Deep Water (CDW) being channelled onto the shelf via glacially carved canyons (Schofield, Ducklow, Bernard, Doney, Patterson-Fraser et al., 2013).

However, there is increasing evidence that CDW is heavily modified as it transits onto the shelf likely due to a flux of silica and iron from shallow marine sediments (Henley et al., 2018; Sherrell et al., 2018). However, even in these relatively nutrient-rich environments, there is still some important direct input from glacial meltwaters (e.g. Annett, Skiba, Henley, Venables, Meredith et al., 2015) due to the release of reactive phases and promotion of biological mediation of nutrient cycling through the formation of organic matter and biogenic minerals.

Despite different boundary conditions compared to the WAP, similar processes are likely to be happening in glaciated regions of SW Greenland. For example, elevated DSi (>20 μM) in Greenlandic fjords measured in surface waters (Hawkings et al., 2017) cannot simply be explained by mixing between the freshwater and marine end-members: these concentrations are higher than the freshwater endmember, and there are no seawater masses with sufficiently high DSi concentrations in the top 100m to supply this flux (Figure 3, see also Appendix C). The fjord water must be modified – in an analogous way to the WAP – likely by particle-water interactions, including the release of DSi from reactive phases derived from glacial weathering products, or biogenic silica (Hawkings et al., 2018; Hawkings et al., 2017; Meire et al., 2016). This modification may be active in the water column, as well as at the sediment-water interface, in the fjords as well as in the shallow water shelf-sediments (Figure 5).

However, despite this enrichment within the fjord, the low-salinity waters reaching the coastal ocean are low in DSi as a result of uptake mechanisms that are active as the fjordic waters reach the shelf; our forthcoming studies of uptake kinetics and algal physiology from within the fjords themselves will elucidate whether biological uptake is playing a key role in Si cycling in these regions. Although there is apparently a limited supply of DSi exported from fjords, there appears to be active cycling of silica by diatoms in coastal waters. Our findings show the potential for meteoric waters, and glacially-derived particles, to be exported as far as the coastal and boundary currents, and into the open ocean, where further processing could act to release bioavailable elements. Whilst some of these exported particles may dissolve within the water column during sinking, some will reach the sediment-water interface. Pore water DSi can be used to evaluate the chemical changes of
the sediments post-deposition such as the dissolution of this reactive glacial material in addition to dissolution of biogenic silica remains of diatoms and sponges, secondary or ‘reverse’ weathering, and the recycling of DSi back to the bottom waters (Rahman, Aller & Cochran, 2017). The high, but variable, DSi concentrations found in the pore waters at our coastal study sites point towards high rates of benthic regeneration fluxes. Calculated sedimentary diffusive fluxes off SW Greenland, using the approach of Ragueneau, Gallinari, Corrin, Grandel, Hall et al. (2001), range from 0.1-0.3 mmol Si m\(^{-2}\) d\(^{-1}\), and are at least 10% of the diatom production rates. Our findings suggest that the total DSi flux across the sediment-water interface, including from advective processes, could rival the magnitude of water column biogenic silica production rates. The high uptake rates of diatoms, together with this rapid recirculation of DSi across the sediment-water interface, points towards a silica cycle maintained by strong pelagic-benthic coupling.

4.5. Approaches to reconstructing glacial meltwater inputs through time

Glacial meltwaters are enriched in both dissolved and particulate nutrients, including silicon, and our new data highlight that these meltwaters extend across the shelf into boundary currents. In the context of the marine silicon cycle, our data show that, whilst DSi reaching the shelf waters from the glacial fjords may be low, diatom activity is surprisingly high. DSi must be reaching the surface, potentially by mixing with modified shelf waters. Our Ra isotopic data (section 4.3.2.) reveal that there is input of glacial particles into these shelf waters, potentially via sediment reworking, which may contribute bioavailable silicon via dissolution both in the water column and at the sediment-water interface. This system is likely to be sensitive to glacial inputs, and so quantifying changes in meltwater fluxes through time - using a variety of climate archives - is going to be key to understanding shelf and slope productivity during past episodes of climatic change.

4.5.1. Fossil deep-sea corals

The geochemistry of fossil skeletons of deep-sea corals has the potential to record aspect of past environmental conditions (Chen, Robinson, Beasley, Claxton, Andersen et al., 2016; Robinson, Adkins, Frank, Gagnon, Prouty et al., 2014). In particular, water masses distribution and food supply are thought to be important for deep-sea coral populations off the West Greenland margin. Given their environmental sensitivity, cold-water coral
distributions are likely to susceptible to changes in water mass properties and primary production caused by meltwater inputs. In 2017, the first living samples of the cold-water scleractinian coral, *Lophelia pertusa*, were collected from approximately 60° 22'N off the Greenland within a layer of relatively warm, modified Atlantic Water (Kenchington, Yashayaev, Tendal & Jørgensbye, 2017).

We have now been able to make in situ observations of cold water corals off Greenland, as well as showing that corals have been present on the Greenland Margin for at least 10,000 years (Figure 10). These first populations likely appeared with melting of the large ice fields of the last glacial period. Supporting prior research, we also found that scleractinian corals have been present further south on Orphan Knoll for at least 130,000 years (Figure 10; Cao, Fairbanks, Mortlock & Risk, 2007; Hillaire-Marcel, Maccali, Ménabréaz, Ghaleb, Blénet et al., 2017). In both locations our suite of dates show that the populations have not been stable. This observation implies shifts in environmental pressures over these timescales, likely driven by a shift in balance between warmer Atlantic waters and cold meltwater-rich polar waters.

![Figure 10](image_url)  
*Figure 10: Uranium-thorium age data for fossil corals collected during DY081.*
During the last ice age, catastrophic iceberg discharge events episodically flooded the subpolar North Atlantic with meltwater (Heinrich, 1988). Evidence suggests that these “Heinrich events” (Broecker, Bond, Klas, Clark & McManus, 1992), were related to, and may have acted as the trigger for (Broecker, 2003; Clark, Pisias, Stocker & Weaver, 2002), dramatic changes in ocean circulation (McManus, Francois, Gherardi, Keigwin & Brown-Leger, 2004) and heat distribution that were felt globally (e.g. Wang, Cheng, Edwards, An, Wu et al., 2001). In addition to global impacts, changes in ice sheet dynamics and meltwater inputs would also have had considerable impacts on more regional biogeochemical cycling.

While previous work has identified the existence of eight such events over the last 70 ka (Andrews, Jennings, Kerwin, Kirby, Manley et al., 1995; Bond, Heinrich, Broecker, Labeyrie, McManus et al., 1992; Bond & Lotti, 1995; Rashid, Hesse & Piper, 2003; Stoner, Channell & Hillaire-Marcel, 1996), questions remain about the origin(s) of these events, their trigger(s) and the nature of their primary signatures across the North Atlantic (Andrews & Voelker, 2018). Gravity cores collected at Orphan Knoll (DY081-GVY002 & GVY002), Southwest Greenland (DY081-GVY003) and Cape Farewell (DY081-GVY004 & GVY 005) represent new opportunities to constrain the timing, geometry and character of iceberg discharge and glacial meltwater release in the paleo record.

Existing work on marine sediment cores from the high latitude North Atlantic has employed a variety of proxies to identify pulses of meltwater delivery and associated ice rafted debris delivery. Previously applied proxies include $^{230}$Th$_{xs,0}$ (to assess changes in sedimentary fluxes (McManus, Anderson, Broecker, Fleisher & Higgins, 1998)), counts of IRD and foraminifera (to identify the relative abundance of terrestrially-derived debris and foraminifera (Heinrich, 1988)), foraminifera census (to determine the relative abundance of cold-dwelling planktonic species such as *N. pachyderma* s. (Ruddiman, Sancetta & McIntryre, 1977)), $\delta^{18}$O of *N. pachyderma* s. (to quantify the cooling and/or freshening of surface waters (Bond, Broecker, Johnsen, McManus, Labeyrie et al., 1993)), magnetic susceptibility (to identify detrital sediment (Grousset, Labeyrie, Sinko, Cremer, Bond et al., 1993)), x-ray diffraction (XRD) (to assess changes in the abundance of quartz and plagioclase feldspar (Moros, Kuijpers, Snowball, Lassen, Bäckström et al., 2002; Moros, McManus, Rasmussen, Kuijpers, Dokken et al., 2004)), and X-ray fluorescence (XRF) (to identify changes in sedimentary elemental ratios (Hodell, Channell, Curtis, Romero & Röhl, 2008)).
Unfortunately, the comparison of sediment core records has been limited by difficulties in establishing tightly constrained chronologies. This has impeded analysis of the triggers of events and the range of their influence (Andrews & Voelker, 2018). While previous work has utilized one or more of the proxy approaches identified above, complete assessment of the proxies described above, in a single core, would greatly assist in interpreting the paleoceanographic record. In addition, despite the widespread inference that icebergs originated from the Laurentide ice sheet during these events (Bond & Lotti, 1995; Broecker, 2003; Hodell et al., 2008; McManus et al., 1998), relatively fewer detailed studies have examined their imprint and consequences in the proximal Labrador Sea (Andrews & Voelker, 2018).

Figure 11: Proxies for North Atlantic Heinrich Events. a) N. Pachyderma (s.) $\delta^{18}$O from DY081-GVY001 (purple). b) Ca/Sr XRF count ratio from DY081-GVY001 (yellow). c) N. Pachyderma (s.) $\delta^{18}$O (Hillaire-Marcel & Bilodeau, 2000) on the age model of Lynch-Stieglitz, Schmidt, Henry, Curry, Skinner et al. (2014) (navy blue). Vertical grey dashes suggest age-depth assignments for DY081-GVY001.
Our goal in analysing the new DY081 cores is to use a wide range of proxies to address temporal relationships and the spatial signatures of Heinrich events. Our findings show that GVY001 (Orphan Knoll) represents a record of continuous sedimentation from 40-50 ka to present and likely captures the Younger Dryas and Heinrich events 1-4 (Figure 11). In GVY001 Heinrich event signatures have been identified in records of IRD counts, the $\delta^{18}$O of planktonic foraminifera, XRD and XRF scans. These results confirm the utility of using the suite of proxies in DY081 cores to tie together previously incomparable records from sediment cores in which disparate proxies have been applied, to reconstruct meltwater inputs in the region back through time.

5. Synthesis and Outlook

Coastal and shelf sea biogeochemical research in the polar regions requires high-resolution spatial and temporal datasets, due to the inherent heterogeneous nature of the high-latitude margin settings. Whilst obtaining the necessary temporal resolution is challenging, especially in the absence of expensive long-term monitoring programs, it is possible to combine traditional physical and biogeochemical measurements with novel isotopic and observational methods that integrate signals over a variety of timescales (days to thousands of years). Stable and radiometric isotope geochemistry also allows the identification of common processes within this highly-variable system, which are active over given timescales. All of these approaches can be combined with palaeoceanographic techniques to obtain a reliable baseline for pre-industrial conditions, and also look for analogues of future change from the past.

In the context of the marine silica cycle, our findings from DY081 are able to show that:

- There are strong mean flows that are conducive to eddy generation, which will likely transport freshwater from the margin into the boundary currents that supply the Labrador Sea;
- Oxygen isotopes robustly trace meteoric water composition adjacent to Greenland, with near-surface concentrations of 5% over the shelf reflecting significant glacial discharge;
- Ra isotopes indicate additions of glacial meltwater and sedimentary particulates are spatially heterogeneous and rapidly transported;
• Low macronutrient concentrations (e.g. DSi) are found in coastal waters influenced by glacial melt;

• Diatoms are surprisingly active given the low nutrient availability and low temperature (i.e. consuming 5-10% of euphotic zone Si daily);

• There is a strong benthic flux of DSi from sediments;

• Fossil corals can be used to track changes in benthic ecosystems through time, likely influenced by water mass distribution;

• Sediment cores recovered of sufficient length and resolution to reconstruct meltwater inputs at least back to 45 ka (i.e. Heinrich events 0-4).

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Table 1: End-member values used in the mass balance calculations.

<table>
<thead>
<tr>
<th></th>
<th>Irminger</th>
<th>Meteoric</th>
<th>Sea ice</th>
</tr>
</thead>
<tbody>
<tr>
<td>Salinity</td>
<td>34.88</td>
<td>0</td>
<td>3</td>
</tr>
<tr>
<td>$\delta^{18}O$ (‰)</td>
<td>+0.34</td>
<td>c. –21</td>
<td>Surface values +2.1</td>
</tr>
</tbody>
</table>
APPENDIX A: Oceanographic setting

The strong West Greenland Current (WGC) brings cold Arctic Water (AW) around the southern tip of Greenland and northwards into the Labrador Sea (Yang et al., 2016). The Irminger Current (IC) brings in warmer North Atlantic Water (NAW) from the North Atlantic into the Labrador Sea – this water is ultimately derived from the Gulf Stream. The typical water column structure near coastal Western Greenland is stratified by salinity, comprising cold, surface water (found shallower than approximately 100 m) that consists of AW and additional meltwater, overlying a strong thermocline; NAW (temperature > 3°C, salinity < 34.5) is found below the thermocline, with the water temperature peaking at a depth of approximately 400m, most likely representing the core of Irminger Water inflow (McCartney and Talley, 1982).

At approximately 64 °N, the latitude of Nuuk, the WGC bifurcates into the Labrador Current (LC), and a proportion is diverted into the interior of the Labrador Sea, where deep water formation occurs by winter-time convection driven by oceanic heat loss (McCartney, 1992). The newly formed Labrador Sea Water (LSW) enters the North Atlantic as the Deep Western Boundary Current, to form the upper layers of North Atlantic Deep Water (NADW), which subsequently become the intermediate layers of the North Atlantic meridional overturning circulation. By the time the boundary current reaches Orphan Knoll (Figure 2a), sub-thermocline warm, saline waters are underlain by the LSW and overflow waters (likely Gibbs Fracture Zone Water and Demark Strait Overflow Water) (Yang et al., 2016).

APPENDIX B: Details of methodologies

Full details of on-board methodology and approach can be found in the DY081 cruise report (Hendry, 2017), but are summarised briefly here.

B.1. Water column sampling and on-board processing: glacial meltwater inputs, nutrients, and phytoplankton

Water column samples were collected using Niskin bottles attached to the CTD rosette (10L volume) and the Remotely Operated Vehicle (ROV) Isis (4 L volume), and via a trace-metal clean towfish.
Duplicate samples for inorganic macronutrients were filtered using either an Acropak or in-line polycarbonate filter (0.2 μm) into acid-cleaned and rinsed high density polyethylene (HDPE) bottles and frozen immediately at -20°C. Clean handling techniques were adopted and only Semperguard vinyl non-powdered gloves were used for the sample handling. Samples for biogenic/amorphous silica were filtered through 25mm 0.6 μm polycarbonate filters, dried and stored for analysis back on land.

Samples were also collected for water oxygen isotope composition (δ¹⁸O) and carbonate chemistry parameters (pH, alkalinity), which are used for investigating freshwater input in high-latitude regions (Hendry et al., 2018; Meredith et al., 2008; Thomas et al., 2011). For δ¹⁸O, an unfiltered water sample was sealed tightly in a (rinsed) 60 mL HDPE plastic bottle and stored in a cool, dark storage location. For carbonate chemistry, a 250 mL borosilicate glass bottle was rinsed twice with seawater from the Niskin before being filled using a PVC tube and allowed to overflow one volume; a glass stopper was placed in the bottle neck to displace excess seawater then 2.5 mL of seawater was pipetted off to allow a 1% headspace. The sample was poisoned with 50 μL saturated mercuric chloride solution and sealed, homogenised and stored in a cool, dark storage location.

Phytoplankton pigments were analysed on board, and compared to sensor-derived fluorescence data, to assess algal standing stocks in relation to meltwater input. Seawater was filtered using GF/F filters and then frozen at -20 °C until extraction. A trichromatic method (Mackereth et al., 1978) was used to determine chlorophyll (Chl) a, b, and c spectrophotometrically in the near surface seawater samples. Chlorophyll extraction was carried out using aqueous acetone buffered with magnesium carbonate, for 24 hours in the dark at 4°C. The samples were then centrifuged and analysed at 750 nm (to correct for turbidity), 664 nm, 647 nm, and 630 nm on a V-1200 Vis spectrophotometer. Absorbance values were then used in equations 1 to 3 to calculate the concentration of Chl a, b, and c per volume of filtered sample (Eq. 4).

\[
\begin{align*}
\text{Chl } a \text{ (mg/L)} &= 11.85 \times (\text{OD664}) - 1.54 \times (\text{OD647}) - 0.08 \times (\text{OD630}) \\
\text{Chl } b \text{ (mg/L)} &= 21.03 \times (\text{OD647}) - 5.43 \times (\text{OD664}) - 2.66 \times (\text{OD630}) \\
\text{Chl } c \text{ (mg/L)} &= 24.52 \times (\text{OD630}) - 7.60 \times (\text{OD647}) - 1.67 \times (\text{OD664}) \\
\text{Chl } x, \text{ mg/m}^3 &= (\text{Chl } x) \times \text{ extract volume, L} / \text{ Volume of sample, m}^3
\end{align*}
\]
B.2. Additional laboratory techniques

Seawater oxygen isotope (δ¹⁸O) measurements were made using the CO₂ equilibration method with an Isoprime 100 mass spectrometer plus Aquaprep device at the British Geological Survey (Keyworth). 200µl samples of water were loaded into exetainers (3.7ml Labco Ltd.) and placed in the heated sample tray at 40°C. The exetainers were then evacuated to remove atmosphere, then flushed with CO₂ and left to equilibrate for between 12 (first sample) - 37 (last sample) hours. Each individual gas sample was then admitted to the cryogenic water trap where any water vapour was removed. The dry sample gas was then expanded into the dual inlet where it was measured on the transducer before being expanded in the dual inlet bellows. Ionvantage software balanced the reference bellows relative to its volume. The sample and reference CO₂ gases entered alternatively into the Isoprime100 through the dual changeover valve for isotope ratio measurement. In each run two laboratory standards (CA-HI and CA-LO) plus up to two secondary standards were analysed in triplicate. The value of these laboratory standards has been accurately determined by comparison with international calibration and reference materials (VSMOW2, SLAP2 and GISP) and so the ¹⁸O/¹⁶O ratios (versus VSMOW2) of the unknown samples can be calculated and are expressed in delta units, δ¹⁸O (‰, parts per mille).

Isotope measurements used internal standards calibrated against the international standards VSMOW2 and VSLAP2. Errors are typically +/− 0.05‰ for δ¹⁸O.

Carbonate analysis was carried out at GEOMAR, Germany. For Total Dissolved Inorganic Carbon (TDIC), carbonate species were converted to CO₂ by addition of phosphoric acid (10% in 0.7 M NaCl), this generated CO₂ is then carried into the measurement cell using N₂ and analysed by coulometric titration using a VINDTA 3C (Marianda, Germany) connected to a 5011 coulometer (UIC, USA). For Total Alkalinity (TA) samples are titrated with 0.1 M HCl (prepared in 0.7 M NaCl) in 150 µL increments until the carbonic acid equivalence point is reached. The titration is monitored with the VINDTA 3C in a closed cell titration (Dickson et al., 2007). Measurements were calibrated using certified reference material from Prof. Dickson, Scripps. The temperature, salinity, and nutrient concentrations of the samples at time of sampling are then combined with the TDIC and TA measurements to calculate CO₂ system parameters (pH and pCO₂).

Samples for inorganic nutrients were all analysed at the Plymouth Marine Laboratory using the latest GO-SHIP (Hydes et al., 2010) recommended defrosting technique of heating
in a warm water bath for 45 minutes from frozen and then equilibrating to room
temperature for another 30 minutes before analysis. The analysis was carried out using a
SEAL analytical AAIII segmented flow colorimetric auto-analyser using classical analytical
techniques for nitrate, nitrite, silicic acid (or DSi) and phosphate, as described in Woodward
and Rees (2001). Clean sampling and handling techniques were employed during the
defrosting, sampling and manipulations within the laboratory, and where possible carried
out according to the International GO-SHIP nutrient manual recommendations of Hydes et
al. (2010). Seawater nutrient reference materials (KANSO Ltd. Japan) were also analysed to
check analyser performance and to guarantee the quality control of the final reported data.
The typical uncertainty of the analytical results were between 2-3%, and the limits of
detection for nitrate and phosphate were 0.02 µM, 0.01 µM for nitrite, and silicic acid did
not ever approach the limits of detection.

We dated 274 fossil scleractinian corals from the three main target sites (Orphan
Knoll, Nuuk, South Greenland) using a reconnaissance dating technique based on the decay
of uranium to thorium (Spooner et al. 2015, Chen et al. 2015).

Foraminifera samples for stable isotope analysis were processed and analysed at the
Lamont-Doherty Earth Observatory of Columbia University. Samples were freeze-dried and
then washed through a 63 µm sieve and the >63 µm fraction dried at 45°C overnight. Five to
ten specimens of *Neogloboquadrina pachyderma* (sinistral) were picked from the >250 µm
size fraction, weighed to ensure consistent sample sizes and then analysed on a Thermo
Delta V Plus with Kiel IV individual acid bath device. Values were calibrated to the VPDB
isotope scale with NBS-19 and NBS-18. Reproducibility of the in-house standard (1SD) is
±0.06‰ for δ18O and ±0.04‰ δ13C.

Ca and Sr intensities (count rates) were measured with an x-ray fluorescence (XRF)
core scanner (ITRAX, Cox Ltd., Sweden) at the British Ocean Sediment Core Research Facility
(BOSCORF). Split core surfaces were smoothed and covered with polypropylene film to
minimize desiccation during analysis. Elemental counts were collected at 1 mm resolution,
using an integration time of 2s and a molybdenum x-ray source set to 30kV and 30mA.
Figure A1. Maps showing DY081 mega core (MGA) and ROV push core sites where pore-fluid samples were taken, and cruise track (black line). Bathymetry resolution of the sampling regions were improved with shipboard multibeam survey. Circles marking the core locations have the same colours corresponding to those denoting the core identities in Figure 5.
Figure A2: A) Example cross section of macronutrient concentrations from the northern section of the Nuuk grid (from CTD 4,5,6,7,14, see Figure 2), showing (from top to bottom): nitrite, nitrate+nitrite, silicic acid, and phosphate (all in µM). This section encompasses the margin from the slope onto the shelf, and did not occupy the bathymetric trough (c.f. Figure 3, main text).
B) Example cross section of integrated (top 50m) macronutrient and algal pigment concentrations from northern section of the Nuuk grid (from CTD 4 to CTD 7, 14, see Figure 2). i) Integrated macronutrients; ii) Integrated pigment concentrations. Error bars show propagated errors on integration calculation (1SD).
A. Calculation one

Salinities set as per $d^{18}O$ mass balance (see main text)
Meteoric water alkalinity = 1000
Seawater alkalinity = 2312
Sea-ice alkalinity = 175
(Alkalinity in $\mu$mol/kg; Data from Jones et al., 2008)

PC1 = 59%
PC2 = 17%
PC3 = 12%
Anomaly PC3 eigenvector < 0.1

Anomaly = $\%$Met$_{318O}$ - $\%$Met$_{ALK}$
Figure A3: Comparison of different freshwater mass balance calculations. Mass balance calculations were carried out using i) salinity and seawater $\delta^{18}O$ (see main text, Table 1), and ii) salinity and total alkalinity (TA) measurements:
\[ F_{ir} + F_{me} + F_{si} = 1 \]
\[ F_{ir}S_{ir} + F_{me}S_{me} + F_{si}S_{si} = S_{ms} \]
\[ F_{ir}A_{ir} + F_{me}A_{me} + F_{si}A_{si} = A_{ms} \]

Where \( F_{ir} \), \( F_{me} \), \( F_{si} \) are the calculated fractions of Irminger Water, meteoric and sea ice melt respectively (Irminger Water being the chosen ocean endmember), which sum to 1 by definition. They result is clearly dependent on the exact choice of endmembers for salinity \((S_{ir}, S_{me}, S_{si})\) and TA \((A_{ir}, A_{me}, A_{si})\) for the Irminger Water, meteoric and sea ice melt respectively. \( S_{ms} \) and \( A_{ms} \) are the measured salinity and TA of each sample. Two different calculations were carried out using different end-members for meteoric water alkalinity: A) 1000 \( \mu \)mol/kg (typical value for the Arctic Ocean, from Jones et al., 2008), and B) 159 \( \mu \)mol/kg (glacial meltwater value from Meire et al., 2015). This latter value is more likely to be a robust representation of the end-member in this region compared to the High Arctic as glacial meltwater is likely to dominated meteoric inputs.

The anomaly between the percentage meteoric water was then calculated using:

\[ \text{Anomaly} = \%\text{Met}_{18O} - \%\text{Met}_{\text{alk}} \]

Principal Component Analysis was then carried out comparing this anomaly with other environmental parameters. Component loadings (lines) and scores (dots) for the main axes are shown (top) together with depth profiles of scores (bottom). Note that the \%Met anomalies show strong loading with axes that also show high scores at subsurface depths, and correlate well with nitrite concentrations and temperature. This suggests that the anomalies arise as a result of non-conservative behaviour perhaps associated with water column nitrification or sedimentary denitrification.
Additional references


Hydes, D. et al., 2010. Recommendations for the determination of nutrients in seawater to high levels of precision and inter-comparability using continuous flow analysers. GO-SHIP (Unesco/IOC).


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