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Abstract. Iceberg-hosted sediments and atmospheric dust transport potentially bioavailable iron to the Arctic and Southern oceans as ferrihydrite. Ferrihydrite is nanoparticulate and more soluble, as well as potentially more bioavailable, than other iron (oxyhydr)oxide minerals (lepidocrocite, goethite, and hematite). A suite of more than 50 iceberg-hosted sediments contain a mean content of 0.076 wt % Fe as ferrihydrite, which produces iceberg-hosted Fe fluxes ranging from 0.7 to 5.5 and 3.2 to 25 Gmoles yr$^{-1}$ to the Arctic and Southern oceans respectively. Atmospheric dust (with little or no combustion products) contains a mean ferrihydrite Fe content of 0.038 wt % (corresponding to a fractional solubility of $\sim$ 1%) and delivers much smaller Fe fluxes (0.02–0.07 Gmoles yr$^{-1}$ to the Arctic Ocean and 0.0–0.02 Gmoles yr$^{-1}$ to the Southern Ocean). New dust flux data show that most atmospheric dust is delivered to sea ice where exposure to melting/re-freezing cycles may enhance fractional solubility, and thus fluxes, by a factor of approximately 2.5. Improved estimates for these particulate sources require additional data for the iceberg losses during fjord transit, the sediment content of icebergs, and samples of atmospheric dust delivered to the polar regions.

1 Introduction

Iron (Fe) is an essential limiting nutrient for phytoplankton. Its supply exerts a significant impact on marine productivity with important implications for the carbon cycle and climate change (Mackenzie and Andersson, 2013). Quantifying Fe sources to the oceans, especially those that may be influenced by climate change, is therefore critical. Global Fe cycles commonly recognise important supplies of dissolved Fe (dFe, < 0.2 or 0.45 µm) from atmospheric dust, continental shelf sediments, and hydrothermal activity (e.g. Breitbarth et al., 2010). Iron isotopes are a promising novel approach (e.g. Conway and John, 2014) to quantifying these different sources but past contributions have commonly been based on estimates and/or measurements of dFe (see Tagliabue et al., 2010; Dale et al., 2015). However, quantifying dFe contributions from atmospheric dust requires an estimate of the solubility of iron. Estimating the solubility of Fe in particulates is particularly important to understanding the Fe cycle in the polar oceans where iceberg-hosted sediments are a source of bioavailable Fe (Smith et al., 2007; Raiswell et al., 2008; Hawkings et al., 2014; Duprat et al., 2016).

The Southern Ocean (SO) is the largest high nutrient–low chlorophyll area where productivity is limited by the delivery of Fe (e.g. Moore et al., 2013). Recent modelling
studies in the SO have focussed on understanding the factors which control spatial variations in productivity but reach different conclusions due to different representations of the Fe cycle and different assumptions regarding Fe solubility and scavenging. For example, Tagliabue et al. (2009) modelled measurements of dFe derived from atmospheric dust and shelf sediments. Atmospheric dust entering seawater was assumed to have a fractional solubility (soluble Fe expressed as a percentage of total Fe) of 0.5% with continued slower dissolution during sinking occurring at a rate of 0.0002% per day. Overall sediments were more important than atmospheric dust, although dust supplies dominated in some regions depending on the model assumptions used. Lancelot et al. (2009) modelled dFe supplies from atmospheric dust, iceberg melt, and shelf sediments. Sediments were the major source, iceberg melt was of lesser significance, and atmospheric dust (assumed to have fractional solubility of 2%) had little influence. The models gave good agreement with patterns of phytoplankton growth but large uncertainties were acknowledged in the magnitude of these sources. Boyd et al. (2012) compared biological utilisation patterns using four mechanisms of Fe supply (vertical diffusivity in areas free of sea ice, iceberg melt, atmospheric dust, and shelf sediments) that were found to have substantial areal extent. Phytoplankton Fe utilisation was highest in regions supplied by Patagonian dust (using fractional solubilities varying from 1 to 10%) and, to a lesser extent, shelf sediments. Wadley et al. (2014) compared the relative magnitudes and variations in supply of dFe from melting icebergs, shelf sediments and atmospheric dust. Sediments were again shown to be the most important source but considerable uncertainty was noted over the flux of Fe from iceberg-hosted sediments. Death et al. (2014) considered a range of sources that included iceberg-hosted sediments and atmospheric dust and found that modelled productivity was significantly enhanced in areas receiving iceberg-hosted sediments and subglacial melt compared to the productivity arising from atmospheric dust (assumed fractional solubility of 2%). However, the contribution from iceberg-hosted sediments was based on a suite of only six samples (Raiswell et al., 2008) that contained 0.15 wt% Fe as ferrihydrite.

These studies show that SO models produce significant differences in the relative magnitudes of the different Fe sources which complicate attempts to isolate overlapping contributions. For example, Tagliabue et al. (2016) show that global dust fluxes of dFe range from 1 to 30 Gmoles yr\(^{-1}\) between different models. Few studies also count for iceberg sources of Fe (see Tagliabue et al., 2016; Table 1), the importance of which may be particularly sensitive to climate change. Climate change is driving increased loss of ice from ice shelves in the Antarctic Peninsula (Vaughan, 2006; Rignot et al., 2011) and ice-shelf shrinkage has also been reported from other areas of Antarctica (Pritchard et al., 2012; Depoorter et al., 2013; Duprat et al., 2016). Ice-shelf losses increase the delivery of potentially bioavailable Fe by iceberg-hosted sediments. Unfortunately, iceberg-hosted sediment data are sparse but current estimates indicate Fe delivery appears to exceed meltwater delivery to the SO by at least an order of magnitude (Hawkings et al., 2014).

Increases in iceberg-hosted sediment delivery are also likely in the Arctic Ocean (AO). A relatively high proportion of primary production occurs on the AO shelves (Pabi et al., 2008) where ice-free areas experience intense phytoplankton blooms due to favourable light and nutrient conditions. Nitrate appears to be the primary limiting nutrient otherwise Fe and/or light become limiting (Popova et al., 2010). Hawkings et al. (2014) have estimated Fe delivery by meltwaters from the Greenland Ice Sheet but no data are available for Fe delivery from iceberg-hosted sediments, although marine-terminating glaciers in the AO are likely to respond to climate change, as in the SO, by producing more icebergs (Bamber et al., 2012) and thus increasing sediment Fe delivery.

Modelling the polar Fe cycles and assessing the impact of climate change requires an improved estimate of the Fe currently released from the particulates present as iceberg-hosted sediments and atmospheric dust. There is a substantial disagreement as to the strength of different sources and reducing their uncertainty is important (Tagliabue et al., 2016). This contribution presents new data for potentially bioavailable Fe from iceberg-hosted sediments and atmospheric dust and also shows how ice transport and storage may influence Fe delivery to the polar regions. The AO and the SO differ in several important respects. The AO receives a substantial riverine flux (~2400 km\(^2\) yr\(^{-1}\); Dyurgerov et al., 2010), more atmospheric combustion products (Luo et al., 2008), has a proportionately smaller area of winter ice (see later) and is also being disproportionately affected by global warming (IPCC, 2013). Changes in Fe delivery to the AO may influence productivity but this is unlikely in the AO where there is no evidence for Fe limitation (except perhaps in summer in the Irminger Basin; Nielsdottir et al., 2009).

The Fe budgets for the AO use the area >60° N (a larger area than that >66.56° N, which is conventionally used to define the Arctic Ocean; Pabi et al., 2008) and the SO budget is based on the area >60° S. The 60° S latitude lies close to the Antarctic Polar Front (the boundary between cold Antarctic waters and warmer sub-Antarctic waters), which runs clockwise from 140° E to 60° W, beyond which the front moves out to 48° S (Moore et al., 1999). Our new flux estimates are based on measurements of ferrihydrite Fe which are deter-

Table 1. Comparison of the FeA content of different size fractions of iceberg sediment.

<table>
<thead>
<tr>
<th>Sample</th>
<th>% FeA</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sieved &lt; 1 mm</td>
<td>0.175 ± 0.005</td>
</tr>
<tr>
<td>Sieved &lt; 250 µm</td>
<td>0.172 ± 0.003</td>
</tr>
<tr>
<td>Sieved &lt; 63 µm</td>
<td>0.162 ± 0.010</td>
</tr>
</tbody>
</table>
mined by the source and mode of delivery and have a fundamental influence on bioavailability. We are concerned only with glacial and atmospheric particulate sources that can be significantly influenced by terrestrial and/or transport processes prior to entry into seawater. The fate of these sources on entering seawater and their spatial variations are outside this focus although our data may inform these research areas.

2 Methodology

2.1 Ice-hosted sediment sampling

Over 60 sediment samples have been collected from icebergs and glaciers at 15 different Arctic and Antarctic locations (Table S1 in Supplement). Data have previously been reported for only 15 of these samples (from 7 localities; see Table S1) and thus the new samples provide a significant expansion of the existing data that now represent a substantial database for Fe in ice-hosted sediments. A set of 41 new iceberg samples were collected from floating icebergs with sediment-bearing layers present in dense, clear blue ice, indicating compressed glacier ice rather than accreted frozen seawater. An additional suite of nine new glacial ice samples was collected from sediment-rich bands in the main body of glaciers (i.e. land-based ice, not icebergs). These samples represent basal ice which has been in contact with the ice-rock interface.

Samples were collected with a clean ice axe, geological hammer, or chisel. The outer layers of ice that might be contaminated were allowed to melt and drain away before the remaining ice was transferred into a new polyethylene bag and allowed to melt. Some loss of dissolved Fe by adsorption or the precipitation of (oxyhydr)oxides during melting is possible (Conway et al., 2015), but the presence of organic complexes (see later) may stabilise dissolved Fe. In any event, melt dFe concentrations are too low (Hawkins et al., 2014) to produce any significant increase in sediment Fe contents. Sediment samples were collected as soon as melting was complete by filtration through a Whatman 542 (2.7 µm pore diameter) filter paper or through a 0.40/0.45 µm membrane filter (Table S3). There is a significant difference in the size fractions produced by filtration through 2.7 µm and 0.40/0.45 µm; however, the filtered iceberg sediment is dominated by coarser material, and variations in the content and masses of the fraction passing through the different filters seem to be too small to produce significant differences in our extractable Fe contents, at least compared to the variations between different samples (see Tables 1, 3, and S3). Small pebbles and grit (> 1 mm diameter) were removed and the remaining material gently disaggregated but not crushed. Any further separations are as described below.

2.2 Atmospheric dust samples

A suite of 15 atmospheric dust samples (Table S2) has been analysed by the same extraction techniques used for the iceberg and glacial samples to ensure data comparability. Seven new samples were collected during a cruise through the eastern tropical Atlantic and into the Sea of Marmara (Baker et al., 2006). Aerosol samples (~ 100 mg) were collected using high-volume (1 m² min⁻¹) aerosol samplers onto single acid-washed Whatman 41 filters (pore size 20 µm; see Baker et al., 2006) and mainly represent mineral dust from the Sahara. Three new samples of dry deposition were collected from a clean window in Southern Patagonia and two new samples of dry deposition were collected from the Eastern Mediterranean: one from a dust collector located in Crete and the other from deposition on to a clean glass surface at Rosh Pina, Israel (Table S2). These bulk mineral dust samples were collected after dust storms and are unlikely to be significantly affected by contamination (see Shi et al., 2009). Relevant data from the literature (Table S2) are also included for three additional dry deposition samples from the Eastern Mediterranean and China (Table S2).

2.3 Analytical methodology

Each sample of air-dried sediment was treated for 24 h by an ascorbic acid solution buffered at pH 7.5. Air drying at room temperature does not achieve complete water loss but < 10 wt % more water is removed by oven drying. The extractant was a solution of 0.17 M sodium citrate and 0.6 M sodium bicarbonate to which ascorbic acid was added to produce a concentration of 0.057 M. This solution was deoxygenated (by bubbling with nitrogen; see Reyes and Torrent, 1997). Approximately 10–40 mg of sample were mixed with 10 mL of the ascorbate solution, shaken for 24 h at room temperature and then filtered through a 0.45 µm cellulose nitrate membrane filter (Kostka and Luther, 1994; Hyacinthe and Van Cappellen, 2004; Raiswell et al., 2010). The Fe removed by ascorbic acid is hereafter termed FeA and reported as dry wt %. Controlling these conditions produces a high degree of selectivity. Fe is quantitatively removed from fresh two-line ferrihydrite and partially dissolved from aged two-line and six-line ferrihydrite and schwertmannite with negligible effects on other Fe (oxyhydr)oxides or clay minerals (Raiswell et al., 2010). The presence of ferrihydrite in iceberg-hosted sediment and subglacial sediment has been confirmed by high-resolution photomicrographs and selected area electron diffraction by Raiswell et al. (2008) and Hawkins et al. (2014).

Ferrihydrite only exists as a fine-grained and highly defective nanomaterial. The more disordered form (Hiemstra, 2013) contains two diffraction lines (two-line ferrihydrite, often called hydrous ferric oxide) and exists as smaller crystals than the form with six diffraction lines (six-line ferrihydrite). The measurement of ferrihydrite is important be-
cause this mineral phase is directly or indirectly bioavailable (Wells et al., 1983; Rich and Morel, 1990; Kuma and Matsunga, 1995; Nodwell and Price, 2001). The delivery of fresh ferricydrate to the open ocean thus has the potential to stimulate productivity in Fe-limited areas (Raiswell et al., 2008; Raiswell, 2011).

The residual sediment was treated for 2h with a solution of 0.29 M sodium dithionite in 0.35 M acetic acid and 0.2 M sodium citrate, buffered at pH 4.8 (Raiswell et al., 1994). Following the ascorbic acid extraction step, the dithionite extracts the remaining (oxyhydr)oxide Fe (aged ferricydrate, goethite, lepidocrocite, and hematite; Raiswell et al., 1994). Dithionite-soluble Fe is hereafter termed FeD and is reported as dry wt %.

Both the FeA and FeD extractant solutions were analysed for Fe either by an atomic absorption spectrometer with an air-acetylene flame or by spectrophotometry using ferrozine (Stookey, 1970). Replicate analysis of a river sediment internal laboratory standard gave analytical precisions of 3% for FeA and 10% for FeD using this sequential extraction. Errors associated with sampling glacial sediments are examined below. Blank corrections were negligible.

2.4 Approach

Estimates of the solubility of Fe in atmospheric dust have utilised a variety of extraction techniques which have produced estimates of fractional solubility ranging from 0.2 to 80% (Jickells and Spokes, 2001), depending on time, pH, and the extractant (Baker and Croot, 2010). Recent studies have attempted to recognise a soluble Fe fraction (extracted with ultra-pure distilled water or seawater) and/or a labile fraction (using a low pH chemical extraction). Distilled water leaches (Sedwick et al., 2007; Berger et al., 2008) have attempted to recognise a soluble Fe fraction (extracted with ultra-pure distilled water or seawater) and/or a labile fraction (using a low pH chemical extraction). Distilled water leaches (Sedwick et al., 2007; Berger et al., 2008; Conway et al., 2015) provide a consistent and reproducible result but losses of Fe can occur due to precipitation of Fe(OH)₃. Rapid filtration or flow through techniques can be used to minimise such Fe losses. Seawater extractions are thought to be less reproducible due to variations in the concentrations of natural binding ligands (Sedwick et al., 2007).

Few of the extractions used to determine labile or leachable Fe have been fully calibrated against different Fe minerals. Baker et al. (2006) extracted Fe using ammonium acetate at pH 4.7, which dissolves negligible concentrations of Fe (oxyhydr)oxides but significant concentrations of Fe as carbonate (Poulton and Canfield, 2005). Chen and Siefert (2003) extracted Fe with a 0.5 mM formate–acetate buffer at pH 4.5, which was stated to dissolve Fe (oxyhydr)oxides (mineralogy unspecified). Berger et al. (2008) use a pH 2 leach with acetic acid and hydroxylamine hydrochloride followed by a 10-min heating step at 90°C. This method (Winton et al., 2015) extracts metals associated with biogenic material, Fe, and Mn (oxyhydr)oxides and adsorbed to clay minerals. Our ascorbic acid extraction is stronger than that by Baker et al. (2006) but weaker than the extractions used by Chen and Siefert (2003) and Berger et al. (2008). The ascorbic acid extraction is, however, selective for fresh ferricydrate, which is the most soluble, and thus potentially bioavailable, Fe (oxyhydr)oxide mineral.

We recognise two particulate fractions (Raiswell and Canfield, 2012) that contain Fe (oxyhydr)oxide minerals (ferricydrate, lepidocrocite, goethite, and hematite), as described below.

1. FeA reported as wt % Fe that is extractable by ascorbic acid and consists mainly of fresh ferricydrate (Raiswell et al., 2011).
2. FeD reported as wt % Fe that is extractable by dithionite. Extraction of FeD following removal of FeA mainly dissolves residual, aged ferricydrate plus lepidocrocite, goethite, and hematite (Raiswell et al., 1994).

An important issue concerns the bioavailability of FeA and FeD. Experimental work suggests that some part of sediment Fe can support plankton growth (Smith et al., 2007; Sugie et al., 2013). Sediment Fe present as fresh ferricydrate (the most soluble Fe (oxyhydr)oxide) is directly or indirectly bioavailable (see above) and is extracted as FeA. FeA mainly comprises nanoparticulate ferricydrate that probably encompasses a range in bioavailabilities (Shaked and Lis, 2012) due to variations in the extent of aggregation and associations with organic matter (which may partially or wholly envelope Fe (oxyhydr)oxide minerals; Raiswell and Canfield, 2012).

We are concerned with Fe mineral reactivity at the point of delivery to seawater where ferricydrate measured as FeA is more labile than FeD (the dithionite-soluble (oxyhydr)oxides which are relatively stable and poorly bioavailable). However, Fe present as FeD may become partially bioavailable after delivery to seawater (for example by dissolution and grazing; Barbeau et al., 1996; Shaked and Lis, 2012), but these complex interactions are outside the scope of the present contribution.

3 Results and interpretation

3.1 Reproducibility of iceberg sediment sampling

The collection of small samples from heterogeneous sediment with a range of grain sizes (clay up to sand size and beyond) is difficult to do reproducibly. Our approach has been to examine the variability both within and between different size fractions. Our previous practice (Raiswell et al., 2008) has been to remove only coarse material > 1 mm diameter, which might severely affect our ability to analyse sub-samples of 10–40 mg reproducibly. Table 1 compares the composition of different size fractions produced by sieving iceberg sediment (from Wallenbergfjorden, Svalbard) first to < 1 mm and then by taking two further replicate subsamples: one sieved to < 250 µm and the other to < 63 µm. Five replicates were analysed from each size fraction to give the means and standard deviations in Table 1.
Reproducibility of the <1 mm fraction of iceberg sediments.

<table>
<thead>
<tr>
<th>Sample</th>
<th>% FeA &lt; 1 mm</th>
<th>% FeA &lt; 63 µm</th>
</tr>
</thead>
<tbody>
<tr>
<td>K1</td>
<td>0.374 ± 0.019</td>
<td>0.377</td>
</tr>
<tr>
<td>K2</td>
<td>0.094 ± 0.019</td>
<td>0.056</td>
</tr>
<tr>
<td>K3</td>
<td>0.044 ± 0.017</td>
<td>0.058</td>
</tr>
<tr>
<td>K4</td>
<td>0.129 ± 0.021</td>
<td>0.102</td>
</tr>
<tr>
<td>K5</td>
<td>0.089 ± 0.007</td>
<td>0.134</td>
</tr>
</tbody>
</table>

A Student’s t test showed no significant differences between mean analyses of wt % FeA in the three different size fractions. In general the wt % FeA would be expected to be larger in the finer fractions, but the enrichment need not be large. A comparison of the FeA contents of the glacial flours studied by Hopwood et al. (2014) showed that <500 µm fractions contained 40–130% of the FeA content of the <63 µm fraction. Shaw et al. (2011) also found a rather similar wt % of FeA in the 63–125 µm (0.038%) and 125–500 µm (0.053%) fractions of iceberg sediment. Thus the finest fractions are not always large enough in mass, or have a high enough wt % FeA, to produce substantial differences between the different size fractions. We next examined the sampling reproducibility using five different iceberg samples (K1-5) from Kongsfjorden, Svalbard (see Table S3), that were sieved through 1 mm with a replicate subsample then produced by sieving to <63 µm. Table 2 shows the mean and standard deviation for five replicate analyses of these iceberg samples sieved through <1 mm and compared to a single analysis of the <63 µm fraction.

No consistent pattern emerged from the data presented in Table 2. Samples with low wt % FeA values (K2 and K3) tended to show the most variation. However, a z test showed a high probability of there being no significant difference between the <1 mm and <63 µm samples for K1, K3, and K5 (p > 5%) but a low probability (p < 0.2%) that samples K2 and K4 were not significantly different. We conclude that our practice of removing only very coarse material by sieving through <1 mm provides a reasonable compromise that achieves good reproducibility (unless the wt % FeA is less than 0.05%) in samples that are coarse enough to be representative of the sediments delivered by icebergs.

### 3.2 Ice-hosted sediment composition

Table 3 summarises the wt % FeA and FeD contents of the iceberg and glacier sediments and the mean and standard deviations of FeA and FeD. Wide variations mainly result from source area geology but there are no significant differences between the compositions of the Arctic and Antarctic icebergs (when the outlying data for Weddell Sea IRD4 are ignored; see Table S3) and hence we are justified in presenting all the iceberg samples as a single group (Table 3).

The wt % FeA and FeD data approach a log-normal distribution and hence logarithmic means are used to calculate the mean values and the logarithmic standard deviations are used to derive the low and high values in Table 3. This approach produces a logarithmic mean FeA content of 0.076 wt % for the iceberg sediments and a range of 0.030 to 0.194%. These new values are based on more than 50 iceberg samples; thus this mean is more reliable than the earlier mean value of 0.15 wt % FeA (based on only six samples from Raiswell et al., 2008) and the large number of samples also permit an estimate of the variation. A Student’s t test on the logarithmic data showed that the iceberg sediments are significantly higher (p < 0.1%) than the logarithmic mean and standard deviation of the wt % FeA contents of the sediments from glacial ice (mean 0.03%; range 0.015 to 0.060%). The logarithmic mean and standard deviation of the values for wt % FeD in Table 3 are also significantly higher (p < 0.1%) in the icebergs (mean 0.377%; range 0.20 to 0.715%) than in the sediments from glacial ice (mean 0.091%; range 0.042 to 0.196%).

### Ice processing effects

The wt % FeA and FeD contents of the iceberg sediments are significantly higher than the glacier-hosted sediments. The icebergs were not all derived from the land-based glaciers we sampled, and part of the differences in FeA and FeD may result from mineralogical/geochemical variations in the glacial bedrock. An alternative explanation for the high wt % FeA and FeD values is that iceberg sediments have undergone alteration during post-calving transport as temperature fluctuations induced melting/freezing cycles that caused dissolution and precipitation. The slightly acidic pH (5.5–6.0) of glacial ice melt (Meguro et al., 2004; Tranter and Jones, 2001) accompanied by the presence of extracellular polymeric substances (EPS) (Lannuzel et al., 2014; Lutz et al., 2014; Hassler et al., 2011, 2015) is able to accelerate the dissolution of Fe (oxyhydr)oxides.

Experimental work by Jeong et al. (2012) showed enhanced dissolution rates of goethite and hematite trapped in ice compared to dissolution rates in water. The degree of enhancement depended on the presence of organic ligands and the surface area of the iron (oxyhydr)oxides; thus the high surface area of ferrihydrite (compared to goethite and hematite) should produce large enhancements. Jeong et al. (2012) found that dissolution was ligand-enhanced and not reductive. However, Kim et al. (2010) have also observed that UV radiation causes the photoreductive dissolution of Fe (oxyhydr)oxides (goethite, hematite) encased in ice to ferrous Fe. Photoreductive dissolution was significantly faster in ice than in aqueous solutions at pH 3.5 (and was 7–8 times faster than the dissolution rates observed by Jeong et al., 2012) and was not influenced by the presence of electron donors. Acids are concentrated by several orders of magnitude at the ice-grain boundary due to freeze concentration ef-
ferts, and the resulting low pH (≈1.5) further enhances both ligand and reductive dissolution (Kim et al., 2010; Jeong et al., 2015). Lin and Twinning (2012) have found elevated concentrations of ferrous Fe within 1 km of a melting iceberg in the Southern Ocean, which they suggest could be derived by the photoreduction of FeA in melt pools. However, most ferrous Fe is likely to be rapidly re-oxidised and precipitated as (oxyhydr)oxide minerals once exposed to the atmosphere by melting, which dilutes the acids and increases pH. The dissolution/precipitation effects of repeated melting/freezing events are explored below, as they might apply to any sediments (including atmospheric dust, see later) encased in ice.

Figure 1 shows an idealised melting/freezing reaction scheme for any sediment in which Fe (oxyhydr)oxides are initially absent and that only contains silicate Fe. Dissolution is initiated in acidic snow melt where Fe is leached slowly by silicate dissolution (Step 1). Subsequent freezing initially concentrates the acids and accelerates dissolution until complete freezing (or consumption of the acids) halts dissolution and induces the precipitation and aggregation of Fe (oxyhydr)oxides as FeA and FeD (Step 2). The transformation of ferrihydrite (FeA) to goethite/hematite (FeD) has a half-life of several years at −3°C (Schwertmann et al., 2004; Brinza, 2010) and hence a proportion of FeA can be preserved over the life time of an iceberg. A new phase of melting (Step 3) causes the dissolution or disaggregation of the newly formed FeA and FeD and also restarts the slow dissolution of silicate Fe. Renewed freezing again accelerates dissolution but finally precipitates FeA and FeD in amounts (Step 4) that have now been increased by the Step 3 dissolution of silicate Fe. Provided there is insufficient time for the transformation of FeA to FeD to be completed then FeA and FeD will both accumulate at the expense of silicate Fe. A comparison of the logarithmic mean FeA contents of the glacial (0.03 wt %) and iceberg (0.076 wt %) sediments and their errors suggests that melting/freezing effects, hereafter termed “ice processing”, could increase FeA contents by factor of 2.5, assuming similar initial FeA contents. These data provide the first, semi-quantitative, estimate of how deposition on to sea ice might enhance the FeA delivery from atmospheric dust. These changes may also be accompanied by other, poorly understood, chemical mechanisms that may further enhance Fe delivery from sea ice (Vancoppenolle et al., 2013).

### 3.3 Iceberg-hosted FeA fluxes

The iceberg-hosted FeA flux (Table 4) is based on sediment encased in icebergs and excludes sediments associated with seasonal ice (see later). The solid ice discharge from Antarctica has been determined as 1321 ± 144 km³ yr⁻¹ by Depoorter et al. (2013) for the period 1979–2010 and from Greenland as 524 ± 51 km³ yr⁻¹ for the period 1958–2010 by Bamber et al. (2012). Van Wychen et al. (2014) estimate that the contribution from other ice masses in Alaska, Svalbard, and the Russian and Canadian Arctic is 34.4 km³ yr⁻¹, for which we assume a 10% error (roughly the same as for the Greenland flux). Hence the total ice loss from the Arctic is 558 ± 55 km³ yr⁻¹ and from the Antarctic is 1321 ± 144 km³ yr⁻¹. Iceberg-hosted sediment FeA delivery can in theory be estimated from the product of ice mass loss, iceberg-sediment content, and FeA concentration but there are significant difficulties.

The ice mass loss does not represent the mass of icebergs delivered into coastal waters, as significant melting may oc-

Table 3. Composition of iceberg, glacial ice, and atmospheric dust samples (number of samples in brackets).

<table>
<thead>
<tr>
<th>Sample</th>
<th>wt %FeA Low</th>
<th>Mean</th>
<th>High</th>
<th>wt %FeD Low</th>
<th>Mean</th>
<th>High</th>
<th>(FeA + FeD) / FeT estimated range (see text)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Icebergs (51)</td>
<td>0.03</td>
<td>0.076</td>
<td>0.194</td>
<td>0.20</td>
<td>0.377</td>
<td>0.715</td>
<td>0.063–0.201</td>
</tr>
<tr>
<td>Glacial ice (16)</td>
<td>0.015</td>
<td>0.031</td>
<td>0.080</td>
<td>0.042</td>
<td>0.091</td>
<td>0.196</td>
<td>0.013–0.059</td>
</tr>
<tr>
<td>Atmospheric dust (15)</td>
<td>0.018</td>
<td>0.038</td>
<td>0.081</td>
<td>0.428</td>
<td>0.868</td>
<td>1.76</td>
<td>0.24–0.52</td>
</tr>
</tbody>
</table>

Low and high values each represent 1 logarithmic standard deviation from the logarithmic mean, except for (FeA + FeD) / FeT.
cur for glaciers that calve into long fjords (Hopwood et al., 2016). Such losses are relatively small in Antarctica where most icebergs are calved from massive, marine-terminating ice shelves and the remainder from outlet glaciers that calve directly into the sea (Silva et al., 2006; Diemand, 2008). However, the characteristics of Greenlandic glaciers vary. One endmember represents fast moving glaciers where the ice mass loss is mostly by calving into the ocean, and the other endmember represents glaciers entering long (up to 100 km) fjords where the ice mass loss is mainly by melting in the fjord (Straneo and Cedenese, 2015; Hopwood et al., 2016). For this latter endmember, fjord circulation patterns largely prevent iceberg-hosted sediments from being delivered directly to coastal waters (Hopwood et al., 2015, 2016). However, the five largest ice mass losses from Greenlandic glaciers occur from the Jakobshavn, Køge Bugt, Ikertivaq, Kangerdlugssuaq, and Helheim glaciers (together representing an ice mass loss of \( \sim 135 \text{ km}^3 \text{ yr}^{-1} \); Enderlin et al., 2014). The first three of these glaciers either calve directly into coastal waters or have relatively short fjord transit times or distances where melting losses should be low, while large icebergs have also been observed to drift > 150 km out of Sermeq Kujalleq (Helheim Glacier; Sutherland et al., 2014). The Jakobshavn, Køge Bugt, and Ikertivaq glaciers deliver approximately 68\% of the 135 km\(^3\) yr\(^{-1}\) directly to coastal waters. Data on fjord mass losses are urgently required but we will proceed by assuming that melting losses are negligible in Antarctica and are 50\% in the Arctic. Thus the ice discharge to the AO is estimated to be 279 ± 27 km\(^3\) yr\(^{-1}\) (Table 4).

Raiswell et al. (2006) and Death et al. (2014) point out that the sediment content of icebergs is poorly constrained but use a value of 0.5 g litre\(^{-1}\), similar to the mean sediment content of river water. Death et al. (2014) cite a range of 0.4–0.8 g L\(^{-1}\) for Antarctic icebergs and a range 0.6–1.2 g L\(^{-1}\) has been inferred by Shaw et al. (2011) based on the sediment load needed to produce the excess \(^{234}\text{Ra}\) activity in the vicinity of icebergs in the Weddell Sea. Substantially larger concentrations (0.2–200 g L\(^{-1}\)) have been found by Dowdeswell and Dowdeswell (1989). Here we use the conservative estimate of 0.5 g litre\(^{-1}\) of sediment but this value may be a significant source of error. The mean wt % FeA content of icebers is 0.076 % with a variability of 0.030 to 0.194 % (Table 3). Deriving the product of the ice mass loss, sediment load, and FeA content (Table 4) shows that the flux of iceberg-hosted FeA to the AO ranges from 0.7 to 5.5 Gmol yr\(^{-1}\), with a mean of 1.9 Gmol yr\(^{-1}\), and to the SO is 3.2 to 25 Gmol yr\(^{-1}\), with a mean of 9.0 Gmol yr\(^{-1}\). The estimated ranges span an order of magnitude and hence all flux values hereon are only quoted to two significant figures.

### 3.4 Atmospheric dust composition

Mineralogy is a key factor in comparing particulate sources, and use of the ascorbic acid extraction technique for the iceberg sediments and atmospheric dust enables their ferricydrite contents (as the most readily soluble and potentially bioavailable Fe mineral) to be compared. The atmospheric dust sample set is relatively small and mainly includes samples that are unlikely to be delivered to the polar regions, although Patagonian dust is a possible source to the SO (e.g. Schulz et al., 2012). Our Patagonian dust sample set is small but a Student’s \( t \) test indicates that there are no significant differences in the concentrations of FeA and FeD between the Patagonian dust and the other dust analysed here. Consistent with this we note that the range of total Fe values (2.9 to 4.3 wt %) for the Patagonian aeolian dust analysed by Gaiero et al. (2007) overlaps the range in our dust (2.8–4.5 wt %; Table S4) and the mean value of 3.5 wt % commonly assumed for atmospheric dust (e.g. Gao et al., 2003; Shi et al., 2012).

Our dust wt % FeA contents are low (mean 0.038 %; range 0.018 to 0.081 %) and are comparable to the wt % FeA contents of the sediments present in glacial ice, but significantly lower (\( p<1 %\)) than the iceberg-hosted sediments (Table 3). Assuming a dust total Fe (FeT) of 3.5 wt %, the range in wt % FeA corresponds to a fractional solubility of \( \sim 1\%\). These data provide a justification for the commonly used fractional

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**Table 4.** Fluxes of FeA derived from iceberg-hosted sediment by melting.

<table>
<thead>
<tr>
<th>Source/notes</th>
<th>Arctic</th>
<th>Antarctic</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ice discharge km(^3) yr(^{-1})</td>
<td>279 ± 27 (a)</td>
<td>1321 ± 144 (b)</td>
</tr>
<tr>
<td>Sediment content g L(^{-1})</td>
<td>0.5 (c)</td>
<td>0.5 (c)</td>
</tr>
<tr>
<td>FeA wt %</td>
<td>0.03–0.076–0.194</td>
<td>0.03–0.076–0.194</td>
</tr>
<tr>
<td>FeA flux Gmol yr(^{-1})</td>
<td>0.7–1.9–5.5</td>
<td>3.2–9.0–25</td>
</tr>
</tbody>
</table>
solvibility range of 1–2 % (see earlier), which is known to be an arbitrary choice (Boyd et al., 2010). However, our ascorbic acid fractional solubility data are difficult to compare with literature values because a wide range of extractions have been used, few of which have been calibrated against ferrihydrite (see earlier). Conway et al. (2015) measure fractional solubility based on the ratio between Fe extracted at pH 5.3 by meltwater and total Fe. A median fractional solubility value of 6 % was found for dust (deposited during the Last Glacial Maximum on ice at Dome C, East Antarctica) that was high in total Fe (8 wt %), possibly due to enrichment in smaller particles as a consequence of long-range transport. Rather lower fractional solubility values (∼3 %) were found at Berkner Island (closer to the South American dust sources) and these data are comparable to the FeA range of our dust data, assuming similar extraction behaviour.

Dust wt % FeD values (mean 0.87 %; range 0.43 to 1.76 %) are significantly higher (p<0.1 %) than in both iceberg and glacial ice sediments. These data suggest that the net effect of weathering and atmospheric/cloud processing (Shi et al., 2015) on our atmospheric dust has more than doubled Fe (oxyhydr)oxides present as the less reactive FeD. The influence of weathering effects alone on soils (potential dust precursors) has been studied by Shi et al. (2011), who showed that the ratio (FeA + FeD) / FeT increased from 0.1–0.2 to 0.5–0.6 in highly weathered samples from areas with relatively high rainfall and temperatures. The (FeA + FeD) / FeT values for the atmospheric dust in Table 3 range from 0.24 to 0.52, which is clearly achievable by weathering alone in the source area. Values of (FeA + FeD) / FeT for the glacial (range 0.013 to 0.059) and iceberg (range 0.063 to 0.201) sediments can also be estimated assuming FeT = 4.2 % (mean value for glacial sediments from Poulton and Raiswell, 2002). These values also suggest a trend of increasing weathering intensity from the glacial to the iceberg sediments (resulting from ice processing effects, see earlier) and to the atmospheric dust. Further data from atmospheric dust delivered to the polar regions are clearly needed to substantiate this conclusion.

3.5 Atmospheric dust FeA fluxes

This FeA flux is based on dust transported through the atmosphere where there is potential for processing (see above) and excludes soils. Localised areas of the Ross Sea are subject to large dust inputs from local terrestrial sands and silts but these appear to be only minor contributors to productivity (Chewings et al., 2014; Winton et al., 2014). Here we proceed cautiously on the basis that the FeA content of our atmospheric dust represents mineral dust (with small to negligible contributions from combustion sources) delivered to the polar regions. Dust deposition fluxes to the SO have been variably estimated as 0.1 to 27 Tg yr⁻¹ (Gao et al., 2003; Mahowald et al., 2005; Jickells et al., 2005; Li et al., 2008). The new flux estimates derived here are based on the Community Earth System Model (Albani et al., 2014), which produces a value of 0.84 Tg yr⁻¹ for dust deposition to the SO. The model version we use has been extensively compared to observations, with the sources modified to best match dust fluxes at high altitude (Albani et al., 2014). In the absence of ice processing, atmospheric dust delivered to the SO with an FeA wt % ranging 0.018 to 0.081 % produces a flux of <0.01 to 0.02 (mean 0.01) Gmol yr⁻¹ (Table 5). This corresponds to a flux of 0.14 to 0.64 µmol m⁻² yr⁻¹ (assuming an area of 19 × 10⁶ km² for the SO).

Comparisons with other Fe flux estimates are difficult due to the different methodologies used. Edwards and Sedwick (2001) measured Fe soluble at pH 2 from snow samples from East Antarctica, deriving a deposition flux of 0.3 to 2.0 µmol m⁻² yr⁻¹. Winton et al. (2015) used an acetic acid plus hydroxylamine hydrochloride extraction (at pH 2) to estimate a flux of 0.64 to 2.5 µmol m⁻² yr⁻¹ for dust being delivered to a sector of the SO >45° S. Both sites are believed to sample clean air with little addition from combustion sources. Our FeA data are at the low end of these estimates (consistent with the higher pH of our ascorbic acid extraction) and suggest that our FeA data provide a reasonable benchmark to compare mineral dust (in the absence of combustion addition) and iceberg fluxes delivered to the SO.

However, the SO is more than 80 % covered by sea ice during winter (declining to a minimum of ∼16 %), which has residence time of 1–2 years (Vancoppenolle et al., 2013). Studies of sea ice show that it can be enriched in Fe by up to 2–3 orders of magnitude relative to the underlying seawater, and the melting edge is commonly associated with plankton blooms (Lannuzel et al., 2007, 2008, 2014). This Fe is derived from more sources than that in icebergs and includes atmospheric dust deposited on the ice surface (augmented by lithogenic dust in near-shore regions) and Fe scavenged from seawater during sea ice formation (Vancoppenolle et al., 2013; Wang et al., 2014). Studies of sea ice in Antarctica have shown high concentrations of Fe that are accompanied by EPS able to solubilise and complex Fe (Lannuzel et al., 2014). We suggest that atmospheric dust deposited on sea ice be processed by melting/refreezing cycle(s) in a similar fashion as dust deposited on icebergs where dust is dissolved (at low pH and aided by EPS) and photoreduced. Our comparison between glacier and iceberg wt % FeA contents (Table 3) indicates that this ice processing has the potential to increase mean wt % FeA contents by a factor of 2.5 from 0.038 to 0.095 wt %. Simulations with the Community Earth System Model (Albani et al., 2014) representing the annual cycle of sea ice show that 0.6 Tg yr⁻¹ of atmospheric dust are deposited on sea ice that melts (enabling ice processing to occur), which produces a mean rate of FeA delivery of 0.01 Gmol yr⁻¹ with a range from <0.01 to 0.02 Gmol yr⁻¹. A further 0.24 Tg yr⁻¹ are deposited on open water (no ice processing), which supplies only small amounts of FeA (<0.01 Gmoles yr⁻¹). Together the delivery
Table 5. Atmospheric dust FeA fluxes.

<table>
<thead>
<tr>
<th>Source</th>
<th>Arctic</th>
<th>Antarctic</th>
<th>Community Earth Systems Model (Albani et al., 2014).</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mass flux Tg yr(^{-1})</td>
<td>5.1</td>
<td>0.84</td>
<td></td>
</tr>
<tr>
<td>FeA wt % (no ice processing)</td>
<td>0.018–0.038–0.081</td>
<td>0.018–0.038–0.081</td>
<td>Based on 15 dust samples from the Atlantic, Mediterranean, and Patagonia with little combustion inputs.</td>
</tr>
<tr>
<td>FeA flux Gmol yr(^{-1})</td>
<td>&lt;0.01–0.01–0.02</td>
<td>&lt;0.01–0.01–0.02</td>
<td>Combustion inputs may range up to similar levels.</td>
</tr>
<tr>
<td>FeA wt % (with ice processing)</td>
<td>0.045–0.095–0.203</td>
<td>0.045–0.095–0.203</td>
<td>Assuming ice processing increases concentrations 2.5 times.</td>
</tr>
</tbody>
</table>

New dust Fe flux estimates to the AO (5.1 Tg yr\(^{-1}\)) are also derived from the Community Earth System Model (Albani et al., 2014) as before. In the absence of ice processing, a mass flux of 5.1 Tg yr\(^{-1}\) dust delivers a range of 0.02 to 0.07 (mean 0.03) Gmol yr\(^{-1}\) of FeA (Table 5). Sea ice in the Arctic has a maximum extent of < 60 % with a residence time of 1–7 years (Vancoppenolle et al., 2013). That part of the dust flux that falls on sea ice (2.1 Tg yr\(^{-1}\)) may be altered by ice processing which increases the wt % FeA by a factor of 2.5 (see above) before being released by melting, as with the SO. Ice-processed dust delivery to the AO provides a mean FeA flux of 0.03 Gmol yr\(^{-1}\) with a range of 0.02 to 0.08 Gmol yr\(^{-1}\). The 3.0 Tg yr\(^{-1}\) of dust delivered to open water supply a mean FeA flux of 0.02 Gmoles yr\(^{-1}\) (range 0.01 to 0.04 Gmol yr\(^{-1}\)) and the total delivery (Table 5) to the AO is the sum of both fluxes (mean 0.05 Gmol yr\(^{-1}\); range 0.03 to 0.12 Gmol yr\(^{-1}\)).

Table 6. Summary data for the main sources of iron to the Arctic and Southern oceans.

<table>
<thead>
<tr>
<th>Source</th>
<th>FeA flux range Gmol yr(^{-1})</th>
</tr>
</thead>
<tbody>
<tr>
<td>Iceberg sediments</td>
<td>Arctic Ocean 0.7–5.5 Southern Ocean 3.2–25</td>
</tr>
<tr>
<td>Atmospheric dust</td>
<td></td>
</tr>
<tr>
<td>Ice processed</td>
<td>0.03–0.12 &lt; 0.01–0.03</td>
</tr>
<tr>
<td>No ice processing</td>
<td>0.02–0.07 &lt; 0.01–0.02</td>
</tr>
</tbody>
</table>

Sources of variation in Tables 4 and 5 relate both to the estimates of mass fluxes as well as the Fe analytical data but improved mass flux estimates may be difficult to achieve given their temporal and spatial variability. Table 6 and Fig. 2 summarise the flux ranges. At first sight there appear to be broad similarities in the magnitude of these Fe sources to the polar oceans but we list below three limitations to the current data set.

The new iceberg and atmospheric dust data presented here provide a valuable insight into the iceberg and dust Fe sources to the polar oceans. They substantiate the view that iceberg sediments have the potential to be a significant source of bioavailable Fe as ferrihydrite (Table 6). We provide a context for the iceberg sediment flux data by using the global shelf flux value of Dale et al. (2015) to derive an order of magnitude estimate of shelf sources (thought to be a dominant source in the SO, see earlier). The Arctic and Antarctic shelf areas represent 11.5 and 7.3 % of the global shelf area (< 200 m depth; Jahnke, 2010). Combining these area percentages with the global shelf flux dFe value of 72 Gmol yr\(^{-1}\) (Dale et al., 2015) suggests shelf sources are approximately 8.3 Gmol yr\(^{-1}\) to the AO and 5.3 Gmol yr\(^{-1}\) to the SO. The shelf areas of the AO and SO that are able to source shelf fluxes of iron are unknown and the values suggested here may be an overestimation. Furthermore, shelf dFe (largely colloidal or nanoparticulate Fe of unknown composition) and FeA as ferrihydrite may not be of similar bioavailability. Nevertheless, the ranges of the shelf and iceberg suggest that both are comparably important sources.

To sea ice and open water supplies a mean of 0.01 Gmol yr\(^{-1}\) with a range from < 0.01 to 0.03 Gmol yr\(^{-1}\) (Table 5).
FeA is present as ferrihydrite, which is potentially bioavailable to phytoplankton although acquisition rates are unknown and may vary substantially between organisms and with local environmental factors (Shaked and Lis, 2012).

Iceberg-derived FeA is a major source of Fe to both the AO and the SO that will likely increase as iceberg delivery increases with climate warming in the polar regions (Table 6 and Fig. 2). Our measurements of iceberg FeA contents are based on a substantial data set, although Antarctic data are still poorly represented. It is clear that iceberg FeA is a major source of potentially bioavailable Fe as ferrihydrite, unless the errors associated with the estimates of iceberg sediment contents exceed an order of magnitude (Raiswell et al., 2008; Death et al., 2014; Hawkings et al., 2014). Modelling the impact of iceberg FeA delivery on surface water dFe concentrations will be complex and will require kinetic models that incorporate scavenging, complexation, dissolution, and sinking (e.g. Tagliabue and Volker, 2011; Raiswell and Canfield, 2012). FeA attached to coarse material will settle out of surface waters quickly, but FeA present mainly as fine-grained material (or nanoparticles) may be held in suspension for long periods in the wake of icebergs. The basal and sidewall melt from icebergs creates complex patterns of upwelling and turbulence producing a persistent water column structure that may last for several weeks and whose influence extends for tens of kilometres and from the surface to 200–1500 m depth (Smith et al., 2013). Furthermore, giant icebergs (>18 km in length) have a disproportionately large areal influence (compared to smaller bergs) which may last for longer than a month (Duprat et al., 2016). The proportion of the FeA found within this area of influence will clearly have a prolonged residence time that may be a key factor in its dispersion and utilisation away from iceberg trajectories into areas where other Fe supplies are limited.

Atmospheric dust fluxes are estimated to be a minor source of FeA to both the AO and the SO, compared to iceberg-hosted sediment, although substantially larger to the AO (Table 6). The dust database used here is small but appears to be globally representative of mineral dust in that the range of wt % FeD contents (2–5 %) overlaps that found in other studies (e.g. Lafon et al., 2004, 2006). There are no comparable data for potential dust sources to the polar regions although Patagonia atmospheric dust (Gaiero et al., 2007) has wt % total Fe values ranging from 2.9 to 4.3 wt % (which overlaps the 3.5 wt % total Fe value commonly used as a global average). Our mineral dust flux estimates could be significantly increased by combustion sources, estimates of which are very dependent on the flux model assumptions, especially those for Fe solubility. Luo et al. (2008) show global maps of the ratio (soluble Fe from combustion)/(total soluble Fe) which ranges from 10 to 40 % in the SO (>60° S) and 20 to 60 % in the AO (>60° N). Ito (2015) also shows that soluble Fe from dust makes up ~50 % of the total soluble Fe. Table 5 acknowledges that combustion sources could be as large as that from dust in some areas of the AO and the SO.

The important features of the new FeA and FeD dust data presented here is that they are closely tied to mineralogy, with FeA measuring the content of fresh ferrihydrite, which is the most reactive and potentially bioavailable Fe mineral. Thus these data enable direct comparison with iceberg sediment FeA delivery. Furthermore we have estimated a potential role for ice processing which appears to enhance FeA contents of dust delivered to sea ice. Mean dust FeA concentrations of 0.095 wt % (if ice processed) approximate to the mean concentration in icebergs (0.076 wt %), which indicates that the former will dominate in areas where dust mass fluxes exceed iceberg sediment delivery, assuming both types of particulates have similar residence times in the ocean. Additional atmospheric dust samples from the polar regions are needed to support these cautious conclusions and to clarify the role of combustion sources. Wet deposition is thought to be the main mechanism of deposition to the SO but fluxes are poorly known (Mahowald et al., 2011). Very high soluble Fe contents (Heimburger et al., 2013) have been found in wet deposition samples from the Kerguelen Islands (at 48° S, which lies outside our SO area) and a similar flux to the area >60° S would represent a major contribution.

Information about the Supplement

All the data used in this manuscript are included in the Supplement.
The Supplement related to this article is available online at doi:10.5194/bg-13-3887-2016-supplement.

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