



Iron budgets for three distinct biogeochemical sites around the Kerguelen Archipelago (Southern Ocean) during the natural fertilisation study, KEOPS-2

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Received: 19 November 2014 – Published in Biogeosciences Discuss.: 19 December 2014

Revised: 22 June 2015 – Accepted: 24 June 2015 – Published: 29 July 2015

Abstract. Iron availability in the Southern Ocean controls phytoplankton growth, community composition and the uptake of atmospheric CO₂ by the biological pump. The KEOPS-2 (Kerguelen Ocean and Plateau compared Study 2) “process study”, took place around the Kerguelen Plateau in the Indian sector of the Southern Ocean. This is a region naturally fertilised with iron on the scale of hundreds to thousands of square kilometres, producing a mosaic of spring blooms which show distinct biological and biogeochemical responses to fertilisation. This paper presents biogeochemical iron budgets (incorporating vertical and lateral supply, internal cycling, and sinks) for three contrasting sites: an upstream high-nutrient low-chlorophyll reference, over the plateau and in the offshore plume east of the Kerguelen Islands. These budgets show that distinct regional environments driven by complex circulation and transport path-

ways are responsible for differences in the mode and strength of iron supply, with vertical supply dominant on the plateau and lateral supply dominant in the plume. Iron supply from “new” sources (diffusion, upwelling, entrainment, lateral advection, atmospheric dust) to the surface waters of the plume was double that above the plateau and 20 times greater than at the reference site, whilst iron demand (measured by cellular uptake) in the plume was similar to that above the plateau but 40 times greater than at the reference site. “Recycled” iron supply by bacterial regeneration and zooplankton grazing was a relatively minor component at all sites (< 8 % of new supply), in contrast to earlier findings from other biogeochemical iron budgets in the Southern Ocean. Over the plateau, a particulate iron dissolution term of 2.5 % was invoked to balance the budget; this approximately doubled the standing stock of dissolved iron in the mixed layer. The ex-

change of iron between dissolved, biogenic particulate and lithogenic particulate pools was highly dynamic in time and space, resulting in a decoupling of the iron supply and carbon export and, importantly, controlling the efficiency of fertilisation.

1 Introduction

The concentration of carbon dioxide in earth's atmosphere, and therefore earth's climate, is highly sensitive to modification in the marine carbon (C) cycle due to the growth of phytoplankton in the Southern Ocean (Sarmiento and Gruber, 2006). These single-cell plants remove inorganic carbon from surface seawater during photosynthesis, and this inorganic carbon can be directly transferred into the deep sea when the plants die and sink, or indirectly through the food web. The Southern Ocean is responsible for 30 % of global ocean carbon export (Schlitzer, 2002). As first demonstrated over 20 years ago, phytoplankton growth in the Southern Ocean is limited by the availability of the micronutrient trace element iron (Fe; Martin, 1990). Low dissolved iron (dFe) availability limits the annual uptake of atmospheric carbon dioxide (CO₂) by the Southern Ocean (Boyd et al., 2000), shapes phytoplankton species composition and physiology (Assmy et al., 2013), the cycling of other nutrient elements (Moore and Doney, 2007), and thus the structure of the entire marine ecosystem (Boyd and Ellwood, 2010).

Artificial mesoscale ocean iron fertilisation experiments have unequivocally demonstrated the role of Fe in setting phytoplankton productivity, biomass and community structure in high-nutrient low-chlorophyll (HNLC) regions (de Baar et al., 2005; Boyd et al., 2007). However, the “carbon sequestration efficiency” of ocean fertilisation as a means to sequester atmospheric CO₂ (calculated as the additional (net) C that is exported from surface waters into the deep (> 1000 m) ocean for a given addition of Fe) varies widely between experiments and is considerably less than estimates from the early iron fertilisation experiments (see discussion in de Baar et al., 2008). This is due to a number of factors, including rapid grazing of phytoplankton in surface waters, the loss of added Fe by its precipitation and scavenging onto sinking particles, differences in estimated or assumed iron-to-carbon (Fe / C) ratios of the cells, and changes in wind mixed layer depth.

The natural resupply of iron to Fe-depleted waters is a more efficient process (Blain et al., 2007), although in part this depends on the mode of Fe delivery (e.g. from above, laterally or from below) and on the ability of organic ligands to keep the supplied Fe in solution (Gerringa et al., 2008), and for continued ocean fertilisation, it is in part reliant on the concurrent supply of other major nutrients. In the Indian sector of the subantarctic Southern Ocean, natural Fe supply from the Kerguelen Plateau (Blain et al., 2007) and Crozet Islands (Pollard et al., 2009) results in increased

phytoplankton biomass during summer, with chlorophyll levels increasing to more than 1 order of magnitude above the background, as revealed by NASA MODIS satellite chlorophyll climatology for January (2003–2010) (Westberry et al., 2013). Previous research on blooms in these localised “natural laboratories” has provided invaluable insights into mechanisms linking iron fertilisation and carbon cycling in the Southern Ocean, especially since studies of natural systems can address the effects of persistent, varying and multiple Fe sources that are not accessible through deliberate artificial mesoscale fertilisation experiments.

The KEOPS-1 (Kerguelen: Ocean and Plateau compared Study 1) project, which took place in the late austral summer of January–February 2005, demonstrated that this natural fertilisation of the Southern Ocean resulted in dramatic changes in the functioning of the ecosystem with large impacts on marine biogeochemical cycles (Blain et al., 2007, 2008a). These observations of the bloom were largely confined to the plateau region, where vertical upwelled supply from the plateau sediments (Blain et al., 2008b; Zhou et al., 2014) and lateral advection of water that had been in contact with the continental shelf of Heard Island to the south (Chever et al., 2010) were the dominant sources of dissolved and particulate Fe (as confirmed using rare earth element (REE) and radium (Ra) isotope tracers; van Beek et al., 2008; Zhang et al., 2008). The interaction of waters, islands and plateau of the Kerguelen Archipelago with several circumpolar fronts of the Southern Ocean allowed us to make a first attempt at placing our regional KEOPS-1 observations within a broader basin-scale context (Blain et al., 2007).

The KEOPS-2 project was designed to improve the spatial and temporal coverage of the Kerguelen region. During KEOPS-2, which was approved as a GEOTRACES process study¹, we studied the region above and downstream of the plateau and observed a massive natural iron fertilisation on the scale of hundreds of thousands of square kilometres. This produced a patchwork of blooms with diverse biological and biogeochemical responses, as detailed in the multiple studies in this special issue of Biogeosciences (volumes 11–12). KEOPS-2 was also carried out in the austral spring to document the early stages of the bloom and to complement the results of KEOPS-1 obtained in late summer during the start of the decline of the bloom, with a principal aim to better constrain the mechanism of Fe supply to surface waters earlier in the season.

Since Fe is actively taken up into phytoplankton and transferred throughout the food web, including removal by particle settling and remineralisation in deep waters, the assessment of its availability is quite complex and cannot be judged from dFe levels in surface waters alone (Breitbarth et al., 2010). Advances in chemical oceanographic techniques for trace elements through the GEOTRACES program (SCOR

¹<http://www.geotraces.org/cruises/cruise-summary/68-science/process-studies/206-geotraces-process-studies>

Working Group, 2007) now allow the measurement of Fe associated with different phases (dissolved and particulate), internal biological recycling and Fe export from surface waters. The results from earlier iron biogeochemical budgets for FeCycle-I (Boyd et al., 2005; Frew et al., 2006), KEOPS-1 (Blain et al., 2007; Chever et al., 2010), CROZEX (CROZet natural iron bloom and EXport experiment; Planquette et al., 2007, 2009) and SAZ-Sense (Sensitivity of the subantarctic zone to environmental change; Bowie et al., 2009) have highlighted that the dominant “new” Fe fluxes are associated with the particulate phase. Particles thus represent an important transport vector for trace metals in the marine ecosystem, although their bioavailability or transfer into a bioavailable fraction remains uncertain. Suspended particles have also been shown to be important aspects of sedimentary, boundary layer Fe sources and export processes (Tagliabue et al. 2009; Homoky et al., 2013; Marsay et al., 2014; Wadley et al., 2014), with particles being transported laterally over hundreds of kilometres in the ocean (Lam et al., 2006; Lam and Bishop, 2008). The biological cycling of particulate Fe may therefore be the most important aspect of the complete Fe biogeochemical cycle, especially since earlier budgets have demonstrated that biological Fe “demand” cannot be satisfied by the new Fe supply (Boyd et al., 2005; Blain et al., 2007; Sarthou et al., 2008; Bowie et al., 2009; de Jong et al., 2012). A simple one-dimensional vertical model that correctly represented the input of dFe to surface waters during KEOPS-1 did not accurately represent the supply of other geochemical tracers or particulate Fe (Blain et al., 2007; van Beek et al., 2008; Zhang et al., 2008), and the role of dissolved and particulate Fe earlier in the season (winter stock) in the Kerguelen region has yet to be quantified.

This paper presents a short-term (days to weeks) Fe budget for the period of KEOPS-2 for each of three process sites: (i) a “plateau” bloom site (A3) on the central Kerguelen Plateau studied during late summer on KEOPS-1 and reoccupied during spring on KEOPS-2; (ii) a “plume” bloom site (E) east of the Kerguelen Islands, which was located within a quasi-stationary, bathymetrically trapped recirculation feature near the polar front (PF); and (iii) a “reference” site (R-2) south of the PF and upstream (southwest) of the Kerguelen Islands in HNLC waters. We focus on mixed-layer integrated pools of dissolved Fe and particulate Fe (which we further separate into biogenic and lithogenic fractions using elemental normalisers), estimate the fluxes of Fe associated with new and recycled Fe sources, and compare Fe supply and demand with implications for bloom duration and magnitude. Our observations also include particulate measurements in both suspended-water-column (in situ pump; ISP) and sinking-export (free-floating sediment trap; “P-trap”) particles below the mixed layer, with linkage to food web processes via a discussion of Fe/C ratios. Finally, we present a seasonal comparison of our springtime budget for KEOPS-2 with late summer observations from KEOPS-1 and also make comparison with findings from other sectors of the Southern Ocean

subjected to natural Fe fertilisation (e.g. Frew et al., 2006 and Boyd et al., 2005 for FeCycle-I, and Ellwood et al., 2014 for FeCycle-II east of New Zealand; Bowie et al., 2009 for SAZ-Sense south of Tasmania; Planquette et al., 2011 for CROZEX near the Crozet Islands; and Zhou et al., 2010 for Blue Water Zone near the western Antarctic Peninsula). The observations of dFe (Qu  rou   et al., 2015) and particulate trace metals (van der Merwe et al., 2015) are detailed in companion papers in this special issue to allow the current paper to focus explicitly on the construction of iron budgets; however, the three papers should be seen as a collective whole.

2 Material and methods

2.1 Study area

The KEOPS-2 (Kerguelen Ocean and Plateau in compared Study 2) expedition was carried out in the Indian sector of the Southern Ocean in the vicinity of the Kerguelen Plateau between 7 October and 30 November 2011 on the RV *Marion Dufresne* (Fig. 1a). The plateau of the Kerguelen Archipelago is a northwest–southeast seafloor feature approximately 500 m deep and is constrained by the Kerguelen Islands to the north and the smaller volcanic Heard and McDonald islands to the south. Our study was conducted in early austral spring when phytoplankton biomass was developing rapidly and forming a mosaic of phytoplankton blooms in the region (Trull et al., 2015; Lasbleiz et al., 2014). Since sampling at the different stations took place at different times over the ~7-week study, our observations also provide a temporal sequence relative to the development of surface biomass.

The Kerguelen bloom has two main features: a northern branch that extends northeast of the island into waters both south and north of the PF and a larger bloom covering ~45 000 km² south of the PF and largely constrained to the shallow bathymetry of the Kerguelen Plateau (< 700 m) (Mongin et al., 2008; Supplement in Trull et al., 2015) (Figs. 1b and 2). Thirty-two stations were sampled during KEOPS-2, often with repeat visits. Here, we focus on three study sites, namely plateau A3, plume E and reference R-2 (Fig. 1). Two visits were made to A3 at the start (A3-1) and end (A3-2) of the voyage (28 days apart), and five visits were made to site E (over 21 days) to document the bloom development. Based on the trajectories of surface drifters, stations E-1, E-3 and E-5 were taken as tracking the middle of a recirculation region (d’Ovidio et al., 2015), so that they can be considered as pseudo-Lagrangian and their succession in time can be considered a first-order time series. Full details of other stations and sampling designed to document the meridional and zonal extensions of the blooms on the plateau and to the east of the Kerguelen Islands are contained in companion papers in this special issue of Biogeosciences.

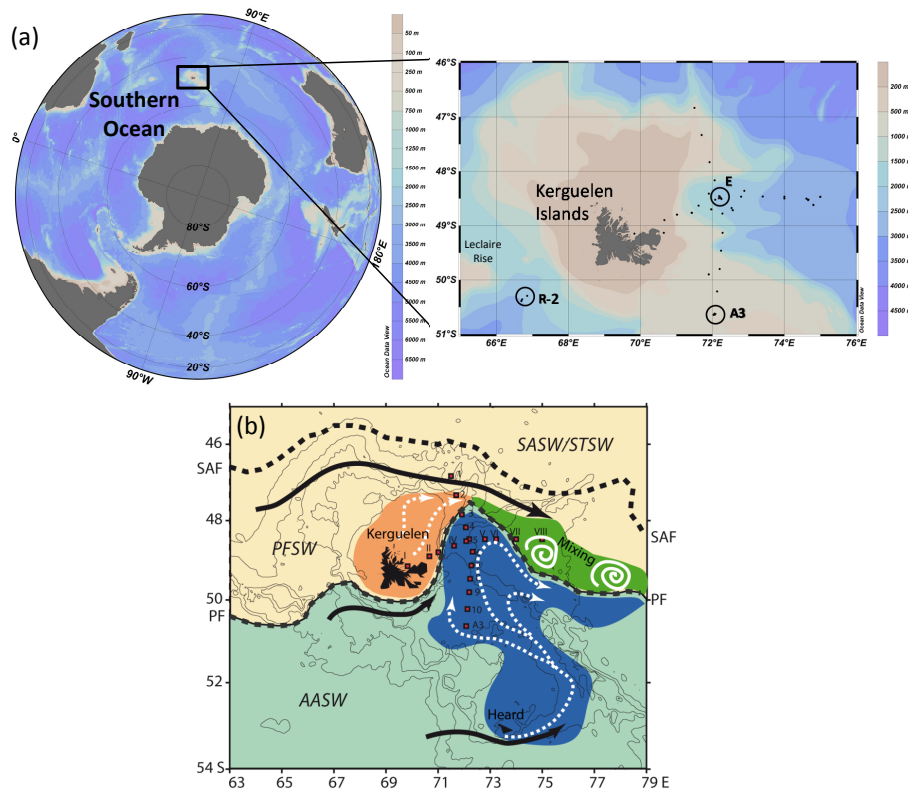


Figure 1. (a) The location of KEOPS-2 in the Indian sector of the Southern Ocean, showing bathymetry around the Kerguelen Archipelago. Our biogeochemical iron budgets focus on three process stations (open black circles): reference R-2 ($50^{\circ}2' S$, $66^{\circ}4' E$), plateau A3 ($50^{\circ}4' S$, $72^{\circ}0' E$) and plume E ($48^{\circ}3' S$, $72^{\circ}1' E$). Black dots mark the positions of the other stations visited, including N–S and E–W survey transects at the start of the KEOPS-2 expedition. (b) A schematic of the mean regional circulation of surface and subsurface waters around the Kerguelen Archipelago, indicating circumpolar Southern Ocean fronts, locations of stations along N–S and E–W transects, and pathways and origins of different water masses flowing on the plateau and offshore into the plume. The abbreviations are Antarctic Surface Water (AASW), Polar Front Surface Water (PFSW), Subantarctic Surface Water (SASW), Subtropical Surface Water (STSW), Subantarctic Front (SAF) and the polar front (PF) (reproduced with permission from Park et al. (2014a), courtesy of Isabelle Durand and Young-Hyang Park, LOCEAN/DMPA, MNHN, Paris).

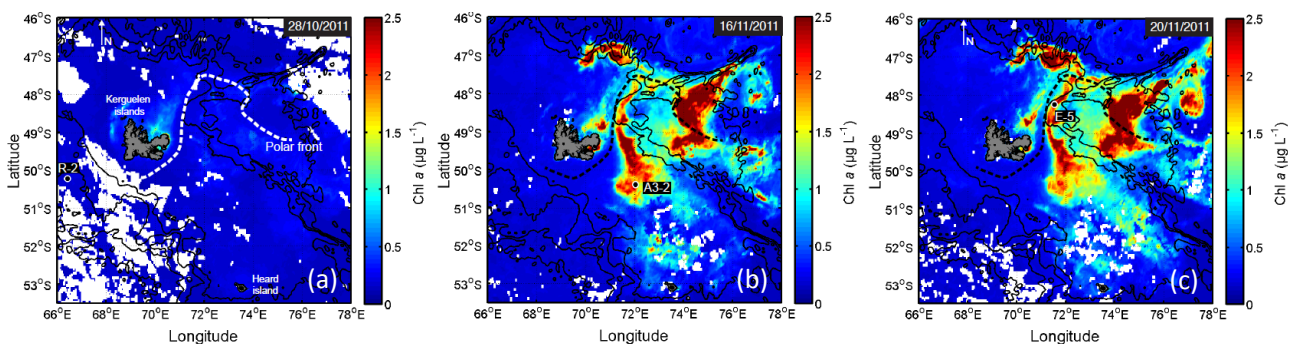


Figure 2. MODIS ocean-colour satellite images showing the development of the plateau and plume blooms during KEOPS-2. Surface chlorophyll ($\mu\text{g L}^{-1}$) biomass is shown for the nearest clear sky day to the final sampling day at stations R-2 (a), A3-2 (b) and E-5 (c). The PF is shown as a black dashed line in (b) and (c). Trull et al. (2015) discuss the timing of the stations relative to bloom development.

The hydrology and circulation around and above the Kerguelen Plateau have been described by Park et al. (2008a, b, 2014a), van Beek et al. (2008), Zhang et al. (2008) and

Zhou et al. (2014). The mean circulation is shown in Fig. 1b. Briefly, the Kerguelen Plateau constitutes a barrier to the eastward flowing Antarctic Circumpolar Current (ACC), the

main jets of which are the Subantarctic Front (SAF) and PF. Most of the ACC is deflected north of the Kerguelen Islands as Subantarctic Surface Water (SASW) but some filaments pass between the Kerguelen Islands and Heard Island (as the PF) and further south between Heard Island and Antarctica (Roquet et al., 2009). Above the plateau, the remainder of the ACC comes from the western part of the plateau. Currents of AASW travelling along the western flank of the plateau are deflected south and east of Heard Island as a branch of the Fawn Trough Current (FTC) (Sokolov and Rintoul, 2009) before travelling in a broadly northwest direction up along the eastern shelf break. The water flow is then deflected toward the east of the Kerguelen Islands, where there is an intense mixing zone consisting of mesoscale eddies which travel many thousands of kilometres in the ACC towards the Australian sector of the Southern Ocean.

2.2 Sampling

All trace metal sampling and analytical procedures followed recommended protocols in the cookbook² published by the international program GEOTRACES (Bishop et al., 2012; Cutter and Bruland, 2012; Planquette and Sherrell, 2012). All methods have been successfully used previously by this team during the KEOPS-1 (Blain et al., 2008b) and SAZ-Sense projects (Bowie et al., 2009). Subtle differences in methods employed during the earlier KEOPS-1 and SAZ-Sense projects are described in those papers and/or later in this manuscript.

2.2.1 Trace metal rosette (TMR)

Water column samples were collected using 10L externally closing, Teflon-lined Niskin-1010X bottles deployed on an autonomous 1018 intelligent rosette system (TMR – trace metal rosette, specially adapted for trace metal work; General Oceanics Inc.). The polyurethane-powder-coated aluminium rosette frame was suspended on Kevlar rope which passed through a clean block with a plastic sheave (General Oceanics) and was lowered to a maximum depth of 1300 m. Bottles were tripped at preprogrammed depths using a pressure sensor as the TMR was being raised through the water column at approximately 0.5 m s^{-1} .

All sample processing was carried out under an ISO class-5 trace metal clean laminar flow bench in a HEPA filtered-air clean container, with all materials used for sample handling thoroughly acid-washed. Samples were drawn through C-Flex tubing (Cole Parmer) and filtered in-line through $0.2 \mu\text{m}$ pore size acid-washed capsules (Pall Supor membrane Acropak 200 or Sartorius Sartobran 300 filters). The dissolved fraction is thus likely to contain colloids and small particles $< 0.2 \mu\text{m}$ in diameter (Bowie and Lohan, 2009). All transfer tubes, filtering devices and sample containers

were rinsed liberally with sample before final collection in 125 mL Nalgene LDPE bottles. Seawater samples were acidified within 24 h of collection using 2 mL of concentrated ultrapure hydrochloric acid (HCl, Seastar BASELINE grade) per litre of sample, resulting in an approximate final pH of 1.8, double bagged and stored for at least 24 h at ambient temperature until analysis.

2.2.2 In situ pumps (ISPs)

Suspended particles for trace elemental analysis were collected using 11 large-volume in situ pumps (McLane Research Laboratories WTS6-1-142LV and Challenger Oceanics pumps), suspended simultaneously at prechosen depths following methods reported in Bowie et al. (2009). Up to 2000 L of seawater was filtered across a 142 mm diameter stack (134 mm diameter active area) consisting of a $53 \mu\text{m}$ nylon pre-filter screen (NYTEX) followed by a QMA quartz fibre filter ($1 \mu\text{m}$ nominal pore size; Sartorius). The QMA filter was supported by a $350 \mu\text{m}$ polyester mesh, which was placed on top of the Teflon PFA grid of the pump housing. Prior to use, NYTEX screens were conditioned by soaking in 5 % H_2SO_4 , rinsed $3 \times$ with Milli-Q-grade water, dried at ambient temperature under a laminar flow hood and stored in clean plastic Ziploc[®] bags. QMA filters were conditioned for trace metal analysis (precombustion and acid cleaning) following Bowie et al. (2010). Upon recovery of the pumps, subsamples were taken from the QMA filters by using a circular plastic punch (14 mm diameter) and by cutting the nylon mesh using ceramic scissors. Filters were dried under a laminar flow bench and stored at -18°C in acid-washed PCR trays until further analysis in the home laboratory. The 1–53 and $> 53 \mu\text{m}$ size fractions were digested and analysed separately, and the particulate iron (pFe) reported here is the sum of both fractions. The ISPs were shown to be efficient in capturing large ($> 53 \mu\text{m}$) particles (Planchon et al., 2015).

2.2.3 Free-floating traps (P-trap)

Sinking particles for trace elemental analysis were collected using PPS3/3 free-floating sediment traps (Technicap, France), specially adapted for trace metals and deployed at 200 m. Traps were deployed for 5.3, 5.1, 1.9 and 1.5 days at stations E-1, E-3, A3-2 and E-5, respectively. The trap deployed at station R-2 was lost and not recovered. Traps drifted between 10 and 43 km over the course of the deployment. Full details of the trap deployments are given in Laurenceau-Cornec et al. (2015) and Planchon et al. (2015). Samples for trace elemental analysis were collected in three separate acid-washed cups (specifically for trace metals) containing a low trace metal brine solution (salinity ~ 60), each opened for either 1, 3, 8 or 12 h (depending on the station). Upon recovery, cups were taken to a clean room and particles filtered off-line onto a 47 mm diameter, $2 \mu\text{m}$ porosity polycarbonate filter under gentle vacuum using a Teflon PFA unit

²<http://www.geotraces.org/libraries/documents/Intercalibration/Cookbook.pdf>

(Savillex Corp., USA) equipped with a 350 μm pre-screen (to exclude zooplankton).

2.3 Analysis

2.3.1 Dissolved iron

Dissolved Fe (dFe) was determined shipboard by flow injection analysis with chemiluminescence detection (FI-CL) using in-line preconcentration on an 8-hydroxyquinoline chelating resin (adapted from Obata et al., 1993, de Jong et al., 1998 and Sarthou et al., 2003). Dissolved Fe data were quality controlled against the SAFe (Sampling and Analysis of Fe) standard reference materials (Johnson et al., 2007). Full data including certification results and analytical figures of merit are reported in Quéroué et al. (2015).

2.3.2 Particulate iron

Particulate Fe (pFe) was determined as follows. Sampled particles were acid extracted in 1 mL concentrated HNO_3 (Seastar Baseline) for 12 h on a DigiPREP HP Teflon hot-plate supplied with HEPA-filtered air (SCP Science) at 120 °C using 15 mL Teflon PFA Savillex vials. Digest solutions were diluted with 10 mL ultra high-purity water to 10 % HNO_3 and spiked with 10 ppb indium as internal standard prior to analysis by sector field inductively coupled plasma mass spectrometry (Finnigan ELEMENT 2, Thermo Scientific), following Bowie et al. (2010). Blanks from replicate analysis of filters treated identically to the sample filters but without large volumes of seawater passed through them, were typically 2–3 % and < 1 % of the pFe sample concentrations for the ISP deployments and P-trap deployments, respectively. Recoveries from the analysis of the Community Bureau of Reference plankton certified reference material BCR-414 were excellent, with a 101 % recovery ($n = 3$) for pFe. Full data are reported in van der Merwe et al. (2015).

2.3.3 Particulate organic carbon and nitrogen

For particulate organic carbon (POC) and particulate nitrogen (PN) analyses, QMA quartz filters from the ISPs were subsampled in a flow bench using a 14 mm diameter plastic punch and transferred to silver foil cups (Sercon brand *p/n* SC0037). Samples were also collected from the P-traps for POC and PN analyses (see Laurenceau-Cornec et al., 2015). Samples were treated with a 40 μL aliquot of 2 N HCl to remove carbonates (King et al., 1998), dried at 60 °C for 48 h and stored in a desiccator until analysis using a Thermo-Finnigan Flash EA1112 elemental analyzer (using sulfanilamide standards) at the Central Science Laboratory, University of Tasmania. The > 53 μm fraction was treated in the same way at the Vrije Universiteit Brussel after first transferring the material from one fourth of the screen, using pre-filtered seawater, onto 25 mm diameter, 1.0 μm pore size silver membrane filters (Sterlitech, Concord). Blank corrections

for the pump samples were estimated from filters prepared identically but not deployed on the ISPs; for the trap samples this was done by re-filtering the pre-filtered seawater. All blank corrections were less than 2 % for all samples. The subsampling introduces uncertainties of 5–10 % from inhomogeneous filter coverage that exceeds the analytical uncertainty of ~ 1 % in the POC analysis (Trull et al., 2015).

2.4 Biological iron cycling

2.4.1 Iron uptake

Trace metal clean seawater was collected from the mixed layer (20–40 m) using the TMR, was transferred into acid-washed polycarbonate bottles and 0.2 nmol L^{-1} (final concentration) of enriched ^{55}Fe as FeCl_3 was added (1.83×10^3 Ci mol^{-1} of specific activity, Perkin Elmer). Bottles were placed at in situ temperature in on-deck incubators continuously fed by surface seawater. Incubations were conducted for 24 h (sunrise to sunrise) at several light intensity levels (75, 45, 25, 16, 4 and 1 % of photosynthetically active radiation; PAR). For stations R-2, A3-1, E-1 and E-3, seawater was prefiltered on a 25 μm mesh size before ^{55}Fe was added. After incubation, 300 mL of seawater was passed through 0.2 μm pore size nitrocellulose filters (47 mm diameter, Nuclepore). To determine intracellular Fe uptake rates, ^{55}Fe not incorporated by cells was removed immediately after filtration using 6 mL of a Ti-citrate-EDTA washing solution for 2 min, then rinsed three times with 5 mL of 0.2 μm filtered seawater for 1 min (Hudson and Morel, 1989; Tang and Morel, 2006). The filters were placed into plastic vials and 10 mL of the scintillation cocktail “Filtercount” (Perkin Elmer) added. Vials were agitated for 24 h before the radioactivity on filters was counted with the Tricarb[®] scintillation counter (precision < 10 %). Controls were obtained with 300 mL of microwave-sterilised seawater (750 W for 5 min) incubated and treated the same way. Subsamples for enumeration by flow cytometry were collected from each bottle just before the filtration step. Cells were fixed in glutaraldehyde (1 %) and kept frozen (–80 °C) until processing and analysis. Data were corrected by blank subtraction and Fe uptake rates normalised to the concentration of Fe in each incubation (in situ dFe and ^{55}Fe added). Further details are given in Fourquez et al. (2015).

2.4.2 Iron remineralisation

Since iron regeneration was not measured directly by experiment during KEOPS-2, we used the following approach to calculate iron regeneration fluxes. Bacterial Fe regeneration was estimated from bacterial turnover times determined from bacterial production and biomass (Christaki et al., 2014), assuming all loss of bacterial biomass through viral lysis and flagellate grazing resulted in the regeneration of Fe (Strzeppek et al., 2005) and using a bacterial iron quota of 7.5 μmol

Fe (mol C)⁻¹ (Tortell et al. 1996). The mesozooplankton grazing contribution to Fe regeneration was assumed to be equal to the experimentally determined Fe regeneration during KEOPS-1 (Sarhou et al., 2008). The regeneration rates per mesozooplankton individual determined in Sarhou et al. (2008), were then multiplied by mesozooplankton abundance, calculated from the number of cells captured in a daily haul over 200 m during KEOPS-2 (Carlotti et al., 2015; values reported in Table 6 in Laurenceau-Cornec et al., 2015).

3 Results and discussion

3.1 Biogeochemical settings at our three study sites

Full descriptions of the dFe and pFe distributions can be found in Qu erou e et al. (2015) and van der Merwe et al. (2015), respectively, with further presentation of the distributions of other micronutrient trace elements (Mn, Co, Ni, Cu, Cd, Pb) from KEOPS-2 to be presented elsewhere. However, briefly our subset of stations used for the iron budgets can be described as follows.

3.1.1 Reference station R-2

In the upper 100 m, we observed a salinity minimum (33.8) and temperature maximum (2.2  C) characteristic of Antarctic Surface Water (AASW) overlying a layer of winter water (WW) at 180–200 m (T_{\min} of 1.6  C) (Fig. 3a). Deeper in the water column, a T_{\max} of 2.5  C at 500 m (associated with an oxygen minimum; not shown) was indicative of upper Circumpolar Deep Water (UCDW) overlying a salinity maximum of 34.8 at 1830 m in lower Circumpolar Deep Water (LCDW). Phytoplankton abundance was low (0.2  g Chl *a* L⁻¹; Lasbleiz et al., 2014) and dominated by diatoms, in waters with relatively high surface nitrate concentrations (> 25  mol L⁻¹; Blain et al., 2015), typical of Southern Ocean HNLC conditions (Lasbleiz et al., 2014).

Dissolved Fe concentrations were very low at the surface (< 0.1 nmol L⁻¹) and increased with depth, averaging 0.3 nmol L⁻¹ in LCDW and broadly tracking the nitrate profile. The pFe profile showed a similar structure to the dFe profile but with surface and deep water concentrations between 0.3 and 1.1 nmol L⁻¹ (the deepest sample was 148 m above the seafloor). The exception was at 500 m, where, interestingly, we observed a dFe and pFe peak of 0.4 and 1.6 nmol L⁻¹, respectively. Whilst this maximum may have arisen due to the enrichment of Fe in UCDW delivered from further south, we hypothesise that the Fe supply may have originated from subsurface sediments of the nearby Leclaire Rise (also known as Skiff Bank; Kieffer et al., 2002), a large seamount which rises to 250 m at 49 50'S, 65 00'E (approximately 140 km northwest of station R-2). Similar lithogenic inputs were also observed for other dissolved (Mn; F. Qu erou e, personal communication, 2014, data not shown)

and particulate (Mn, Al; van der Merwe et al., 2015) trace elements.

The dFe profile at the KEOPS-2 reference station R-2 is similar to the KEOPS-1 reference station C11 (with the exception of the R-2 enrichment in the 200–700 m depth strata; Fig. 4a), but it should be noted that the location of C11 was quite different – in HNLC waters to the southeast of the Kerguelen Plateau (51 39'S, 78 00'E) – and we had only 1 dFe data point in UCDW at C11. In contrast to the similarity of the dFe profiles, the pFe profile at C11 was generally lower than at R-2, with mean values through the water column of 0.2 ± 0.14 nmol L⁻¹ (Andrew Bowie, unpublished data) compared to 0.53 ± 0.35 nmol L⁻¹ for station R-2.

3.1.2 Plateau station A3

Stations A3-1 (Fig. 3b) and A3-2 (Fig. 3c) were in relatively shallow waters on the central plateau, and were impacted by plateau sediments and possibly fluvial and glacial runoff from the basaltic rocks of Heard Island ~ 300 km upstream (van der Merwe et al., 2015; M. Grenier, personal communication, 2014). A pycnocline was observed at ~ 190 m, above which the salinity (33.9) and nitrate (~ 29  mol L⁻¹) were relatively constant. The mixed layer shoaled (from 165 to 123 m) and increased in temperature (from 1.7 to 2.2  C) between the two visits to A3, consistent with springtime warming of surface waters. We believe that the water masses at A3-1 and A3-2 are comparable since surface waters move slowly in this region (Park et al., 2008, 2014a; Zhou et al., 2014); this was confirmed by REE data which indicated similar waters at both stations marked with fresh continental supplies and only modified by biological processes (M. Grenier, personal communication, 2014).

Surface chlorophyll images revealed that during the 28 days between the first and second visits to A3, a large diatom spring bloom developed mostly dominated by lightly silicified *Chaetoceros* spp. (surface Chl *a* increasing from 0.2  g L⁻¹ at A3-1 to 1.3  g L⁻¹ at A3-2; Lasbleiz et al., 2014), which likely resulted in the drawdown of dFe (mean mixed layer values decreasing from 0.3–0.4 nmol L⁻¹ at A3-1 to 0.1–0.2 nmol L⁻¹ at A3-2). The peak of biomass had passed by the time we sampled at A3-2, with the bloom starting to fade (Trull et al., 2015). Below the mixed layer, similar dFe profiles were observed during both visits to A3, with expected significant increases at depth towards the plateau floor (e.g. to 1.30 nmol L⁻¹ at 480 m at A3-2; note that, due to operational constraints, there was no dFe data deeper than 340 m at A3-1). Such enrichments at depth were also observed in dissolved Mn and Co profiles (F. Qu erou e, personal communication, 2014; data not shown) and dFe profiles from the occupations of station A3 during KEOPS-1 (Fig. 4b), indicative of plateau sedimentary supply.

The pFe profiles at A3 showed a similar structure to the dFe profile, with lower values at the surface (< 10 nmol L⁻¹ at A3-1 and < 4 nmol L⁻¹ at A3-2) and increasing with depth

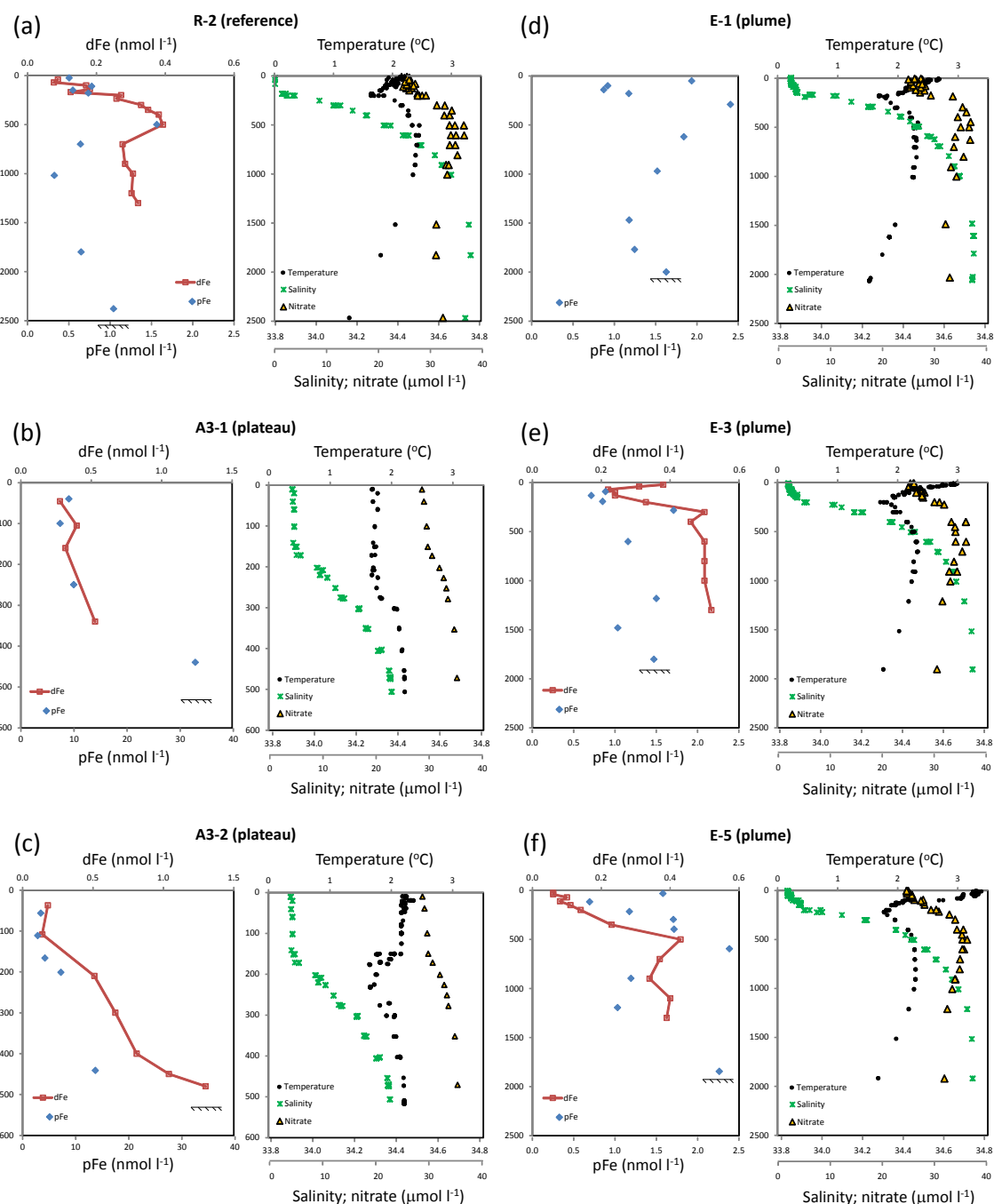


Figure 3. (a) Vertical profiles of dissolved iron (dFe) and particulate iron (pFe), potential temperature, salinity, and nitrate at reference station R-2. The seafloor depth at 2528 m is shown. (b, c) Vertical profiles of dFe and pFe, potential temperature, salinity, and nitrate at plateau stations A3-1 (b) and A3-2 (c). The seafloor depth at ~ 530 m is shown. Note different scales for dFe and pFe compared to R-2 and E stations. (d, e, f) Vertical profiles of dFe and pFe, potential temperature, salinity, and nitrate at plume stations E1 (d), E3 (e) and E5 (f). The seafloor depth ranging from 1905 m (E3) to 2057 m (E1) is shown.

due to enrichment from bottom sediments (up to 33 and 14 nmol L^{-1} at 440 m at A3-1 and A3-2, respectively), and were on average 10 times greater than dissolved concentrations through the water column. The mixed layer pFe con-

centrations changed remarkably between the two visits, and the full water column integrated pool was $\sim 70\%$ lower at A3-2 than at A3-1. Interestingly, this change was also associated with a shift of particles from the 1–53 μm size range

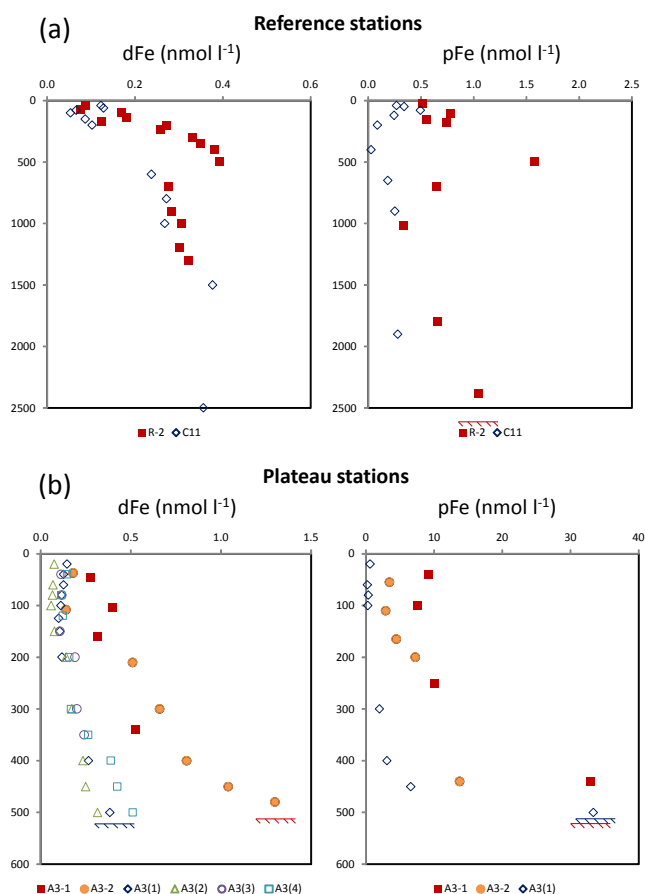


Figure 4. (a) Comparison of dFe and pFe at reference stations for KEOPS-1 (station C11, open blue diamonds) and KEOPS-2 (station R-2, closed red squares). The water depths were 3110 m at C11 and 2530 m at R-2. (b) Comparison of dFe and pFe at A3 plateau stations for KEOPS-1 (open symbols) and KEOPS-2 (closed symbols). Data are shown for all visits to A3 on both KEOPS cruises. Note difference in scale for dFe and pFe between (a) and (b).

to the $> 53 \mu\text{m}$ size range, with the larger class tripling in size (van der Merwe et al., 2015). The development of the large bloom between our two visits to A3, which consisted of a diatom community 50–210 μm in size (Trull et al., 2015), was likely responsible for converting the pFe within the surface mixed layer from the smaller size class to the larger size class. This may have been due to either (i) physical aggregation of the particles onto diatom aggregates and/or (ii) microbially driven conversion of small lithogenic Fe (1–53 μm) to bioavailable forms and incorporation into the large ($> 53 \mu\text{m}$) diatoms as biogenic Fe, with potentially some fraction of these larger particles exported to depths below the mixed layer, as previously discussed by Lam et al. (2006), Frew et al. (2006) and Planquette et al. (2011).

The spring (Oct–Nov) KEOPS-2 Fe profiles at station A3 showed a similar structure to those from the late summer (Jan–Feb) during KEOPS-1, with surface depletion, concen-

trations increasing with depth and enrichment just above the plateau seafloor (Fig. 4b). Through the water column, dFe was between 2 and 5 times greater during KEOPS-2 than KEOPS-1 and pFe was ~ 10 times greater during KEOPS-2 (with the exception of the deepest samples). The lower values during KEOPS-1 were likely the result of biological uptake in surface waters and the export of Fe during the spring bloom prior to our arrival at the study site, combined with seasonal changes in the strength of the supply mechanisms to deeper waters at A3 (discussed in van der Merwe et al., 2015).

3.1.3 Plume E stations

The E stations within the bathymetrically trapped complex recirculation system showed similar hydrographic and nutrient distributions below the mixed layer (Fig. 3d, e and f), which shoaled from 64 m at E-1 to 32 m at E-3 to 39 m at E-5, with some internal variability in water column structure at mid-depths. Surface waters warmed from 2.7 to 3.4 $^{\circ}\text{C}$ between the occupations of E-1 and E-5, although no significant nitrate drawdown was observed (Blain et al., 2015). Below AASW, a subsurface temperature minimum (T_{min} , $\sim 1.7^{\circ}\text{C}$) was observed between 180 m (E1) and 220 m (E5), characteristic of WW. The T_{min} feature is associated with waters south of the PF, although the recirculation feature probably also received SAZ waters mixed in from the north (d’Ovidio et al., 2015). T , S and O_2 characteristics indicated the presence of UCDW (~ 600 – 700 m) and LCDW (deeper than ~ 1300 m) deeper in the water column above the seafloor (Qu erou e et al., 2015). Water parcel trajectories calculated from altimetry based geostrophic currents indicated that it took generally > 2 months for Fe-rich waters from the plateau to travel to the downstream plume site associated with the recirculation feature (E stations) (d’Ovidio et al., 2015). However shorter transport times are also possible due to episodic transport across the PF (Sanial et al., 2015).

Waters at the plume stations showed the largest spatial heterogeneity in surface biomass as revealed by the evolution of a mosaic of complex blooms seen in satellite images (see Supplement in Trull et al., 2015). We observed moderate surface Chl a levels ranging from 0.3–0.4 $\mu\text{g L}^{-1}$ at E-1 and E-3 to 0.5–0.9 $\mu\text{g L}^{-1}$ at E-5 (Lasbleiz et al., 2014), noting that as much as 50% of the chlorophyll was below the mixed layer at the plume stations due to stratification of the upper water column in the warm, spring conditions. Unlike the plateau bloom dominated by large cells $> 53 \mu\text{m}$, the community in the plume E stations was more mixed (Laurenceau-Cornec et al., 2015), with cells present in both the 5–20 and 50–200 μm size classes (Trull et al., 2015). The E stations showed the highest C export fluxes of all regions as estimated from Th deficits, nitrate depletions and free-drifting sediment trap observations (Planchon et al., 2015; Trull et al., 2015; Laurenceau-Cornec et al., 2015).

Due to operational constraints, no dFe data were available at station E-1. The dFe vertical profiles at E-3 and E-5 were quite different, with a distinct surface enrichment to 0.4 nmol L^{-1} at E-3 above a minimum of 0.2 nmol L^{-1} at 100 m. This feature was absent at station E-5, where dFe was depleted to $< 0.1 \text{ nmol L}^{-1}$ at the surface, likely due to biological Fe uptake, which was highest at E-5 ($1745 \text{ nmol m}^{-2} \text{ d}^{-1}$) compared to A3-2 ($1120 \text{ nmol m}^{-2} \text{ d}^{-1}$) (Table 1) and E-4E ($880 \text{ nmol m}^{-2} \text{ d}^{-1}$; data not shown), despite lower POC and primary production (see discussion below and Fourquez et al., 2015). Deeper in the water column ($> 500 \text{ m}$) at E stations, dFe was broadly uniform ($0.3\text{--}0.5 \text{ nmol L}^{-1}$).

The pFe distributions at the three E stations were similar with a surface (35–40 m) enrichment ($1.6\text{--}1.9 \text{ nmol L}^{-1}$), a minimum at $\sim 100\text{--}200 \text{ m}$ below the mixed layer ($0.7\text{--}0.9 \text{ nmol L}^{-1}$; broadly consistent with the T_{min} layer), a maximum at 280–600 m ($1.7\text{--}2.4 \text{ nmol L}^{-1}$), and with evidence of enrichment near the seafloor at depths $> 1800 \text{ m}$ (up to $1.5\text{--}2.3 \text{ nmol L}^{-1}$). By applying biogenic (using P) and lithogenic (using Al) normalisers to the data (see Sect. 3.2 below), surface pFe enrichment was roughly equally composed of biogenic and lithogenic Fe, whilst the 300–600 m maximum was predominantly composed of lithogenic Fe (> 100 -fold greater than biogenic Fe at these depths). This lithogenic Fe was most likely from waters enriched by sediments and transported laterally eastward off the Kerguelen Plateau which sits at $\sim 530 \text{ m}$ below the sea surface. There was no obvious change in pFe in surface or deep waters during the bloom evolution at the pseudo-Lagrangian E stations.

KEOPS-1 only occupied one station in the plume east of the Kerguelen Islands (A11 at $49^{\circ}09' \text{ S } 74^{\circ}00' \text{ E}$). Dissolved Fe at A11 ranged from 0.09 nmol L^{-1} at the surface to 0.17 nmol L^{-1} at 1500 m (Blain et al., 2008b), and pFe ranged from 0.07 nmol L^{-1} at the surface to 0.81 nmol L^{-1} at 1500 m (Andrew Bowie, unpublished data); thus, it was much lower than our KEOPS-2 observations at the E site (noting that different sampling and digestion methods for pFe were used for the two cruises).

3.2 Construction of iron budgets

The primary aim of this work was to use our observations of Fe pools and fluxes to understand the sources, sinks and biological Fe cycling and to evaluate whether Fe supply could meet demand in both the high-Fe and low-Fe environments in the vicinity of the Kerguelen Archipelago during KEOPS-2. Iron budgets have been constructed for previous studies in waters fertilised with Fe both naturally (Sarhou et al., 2008; Bowie et al., 2009; Chever et al., 2010; Ellwood et al., 2014) and artificially (Bowie et al., 2001) as well as low-Fe conditions (Price and Morel, 1998; Boyd et al., 2005). These budgets have combined geochemical and chemical components to demonstrate that the dominant long-term fluxes of Fe are associated with the particulate pool (dust supply and

particle export), whilst studies on Fe uptake and microbial cycling have shown that short-term fluxes within the “ferrous wheel” are dominated by biological uptake and remineralisation (Strzepek et al., 2005). Here, we follow a similar approach to that used by Bowie et al. (2009) for the SAZ-Sense study south of Tasmania (Australia) at our three study sites. Since all parameters in our iron budget calculations were only measured at stations R-2, A3-2 and E-5, discussion will focus on these stations. Data for stations A3-1, E-1 and E-3 are given to provide a context for spatial and temporal changes in the Fe pools and fluxes during KEOPS-2, and they are collated in Table 1.

3.2.1 Iron pools

Iron and carbon pools were calculated by integrating the dissolved and particulate profiles down to the base of the surface mixed layer, defined as the depth where the potential density equalled the potential density at $10 \text{ m} + 0.02 \text{ kg m}^{-3}$ (Park et al., 2014a). The mixed layer varied from 165 m at station A3-1 to 32 m at station E-3, consistent with the seasonal shoaling as surface waters warmed, but it remained deep ($> 120 \text{ m}$) on the plateau throughout the study due to deep mixing as a result of several passing storms.

Integrated pools of both dissolved ($\sim 5\times$) and particulate ($\sim 10\times$) Fe were significantly greater on the plateau (station A3) compared to in the plume (station E), with stocks at the reference station R-2 lower still. Horizontal dFe supply from the plateau to the plume was either or both via (i) a geostrophic path looping along the northern side of the PF and then back into the recirculation feature (d’Ovidio et al., 2015) and (ii) direct Ekman flux transport of Fe-rich coastal water across the PF driven by westerly winds, as indicated by Ra tracers (Sanial et al., 2015). The latter process is supported by Lagrangian trajectories of water parcels derived from altimetry, which showed the PF was not a strong barrier to water mass movement, with transport of waters across the front taking place on timescales of days to weeks but being highly variable in space and time (d’Ovidio et al., 2015). The pFe pool showed the same variability as the dissolved pool at our three study sites and exceeded the dFe stocks at all sites by factors of approximately 19–26 (A3), 31 (E) and 6 (R-2), although it is estimated that only $\sim 2\text{--}3\%$ of the particulate pool can be converted into bioavailable forms by physically or biologically mediated dissolution (Schroth et al., 2009). If we assume that station A3-1 represented pre-bloom conditions and the integrated mixed layer pool of $54 \mu\text{mol dFe m}^{-2}$ was a good estimate of the winter stock, observations show that only 4 weeks later at station A3-2, almost 60% of the winter stock had been drawn down to $21 \mu\text{mol m}^{-2}$. If annual variability is low, which may not always be the case (Grenier et al., 2015), by late summer $> 90\%$ of the winter stock had been used with only $4.7 \mu\text{mol dFe m}^{-2}$ remaining in the surface mixed layer at A3 (KEOPS-1 data; Blain et al., 2007). We note that this drawdown is probably a con-

Table 1. Summary of iron standing stocks and fluxes for the upper mixed layer at KEOPS-2 process station sites R-2 (reference), A3 (plateau) and E (plume). For full details of the calculations, see text. Error bounds are provided where available. Due to logistical constraints resulting in missing data at some stations, we will focus on R-2, A3-2 and E-5 in the discussion. Data for stations A3-1, E-1 and E-3 are given to provide a context for spatial and temporal changes in the pools and fluxes during KEOPS-2.

Region Station	Reference R-2	Plateau A3-1	A3-2	Plume E-1	E-3	E-5
Location	50°21.53' S, 66°42.44' E	50°37.88' S, 72°4.99' E	50°37.47' S, 72°03.35' E	48°27.44' S, 72°11.26' E	48°42.13' S, 71°58.1' E	48°24.69' S, 71°53.99' E
Mixed layer depth (m) ¹	76	165	123	64	32	39
Bottom depth (m)	2528	533	530	2057	1905	1920
Iron pools, integrated over the mixed layer ($\mu\text{mol m}^{-2}$, unless otherwise stated)						
dFe	7 ± 1	54 ± 10	21 ± 4	n.d. ²	12 ± 0	2 ± 0
pFe	43 ± 0	1392 ± 195	401 ± 52	117 ± 1	n.d. ³	61 ± 1
Biogenic pFe	9	13	14	11	n.d.	9
Lithogenic pFe	12	892	265	33	n.d.	9
POC (mmol m^{-2})	124 ± 11	239 ± 33	274 ± 24	198 ± 10	n.d.	150 ± 12
Iron fluxes ($\text{nmol m}^{-2} \text{d}^{-1}$, unless otherwise stated)						
(a) Diffusion	2	42	93	n.d.	1	0.5
(b) Upwelling	35	200	250	n.d.	330	140
(c) Entrainment	57	769	769	n.d.	330	330
(d) Total vertical dFe supply (a + b + c)	94	1011	1112	n.d.	661	471
(e) Lateral advective dFe supply	0		180		2400 ± 600	
Ratio of lateral-to-vertical supply (e/d)	0		0.2		4–5	
Atmospheric total Fe deposition				500 ± 390		
(f) Atmospheric soluble Fe deposition				50 ± 39		
Downward total pFe export flux	1302 ± 586 ⁴	n.d.	5746 ± 1198	4579 ± 1376	1890 ± 286	895 ± 358
(g) Downward non-lithogenic pFe export flux	–	–	2797 ± 583	–	–	541 ± 216
Downward POC export ($\text{mmol m}^{-2} \text{d}^{-1}$)	1.8 ± 0.9 ⁵	n.d.	2.2 ± 0.7	7.0 ± 2.3	4.9 ± 1.5	2.0 ± 1.0
(h) Iron uptake ⁶	40 ± 6	2528 ± 704	1120 ± 389	n.d.	743 ± 194	1745 ± 350
(i) Iron remineralisation ⁷	10 ± 2	19 ± 6	71 ± 12	27 ± 2	23 ± 2	31 ± 2
Fe/C ratios (mmol mol^{-1})						
(j) Mixed layer Fe/C cellular uptake ratio ⁸	n.d.	n.d.	0.007 ± 0.004	n.d.	n.d.	0.021 ± 0.002
Suspended mixed layer particulate $\text{Fe}_{\text{tot}}/\text{C}$ ratio ⁸	0.2 ± 0.1	3.3 ± 0.4	1.5 ± 0.2	0.5 ± 0.1	n.d.	0.4 ± 0.1
Sinking $\text{Fe}_{\text{tot}}/\text{C}$ export ratio	n.d.	n.d.	2.6 ± 1.0	0.7 ± 0.5	0.4 ± 0.3	0.5 ± 0.1
Iron supply vs. demand (for reference R-2, plateau A3-2 and plume E-5 stations only) ($\text{nmol m}^{-2} \text{d}^{-1}$)						
Total iron supply from new sources (d + e + f) ⁹	144	–	1342	–	–	2921
(k) Additional iron requirement to balance the dissolved budget (d + e + f – h + i) ¹⁰	114	–	293	–	–	1207
(l) Biological uptake of new iron (d + e + f – g) ¹¹	–1158	–	–1455	–	–	2380
<i>fe</i> ratio (l/h) ¹²	–	–	–	–	–	1.4
Fe ratio (g/h) ¹²	–	–	–	–	–	0.3
Estimated vs. observed production ($\text{mmol C m}^{-2} \text{d}^{-1}$)						
Potential new primary production (l/j) ¹³	–	–	–	–	–	132
Observed net primary production ¹⁴	11 ± 0	n.d.	158 ± 15	44 ± 4	57 ± 8	79 ± 9

n.d.: no data

¹ The mixed layer depths were calculated on the density plane to allow for heave (internal tides driven by topography) and other localised events.

² Due to logistical reasons there was no TMR cast for dFe at station E-1.

³ Due to ISP failure, there were no mixed layer samples for pFe at station E-3.

⁴ The P-trap was lost at R-2. We therefore estimated the pFe export flux using the ²³⁴Th flux in suspended particles at 200 m ($449 \pm 203 \text{ dpm m}^{-2} \text{d}^{-1}$; from Table 1 in Planchon et al., 2014) and a mean Fe/Th ratio collected in the upper 200 m above the trap ($2.9 \pm 1.3 \text{ nmol dpm}^{-1}$).

⁵ Estimated using the ²³⁴Th flux and Fe/C ratio in suspended particles at 200 m.

⁶ For stations R-2, A3-1, E-1 and E-3, seawater for iron uptake experiments were conducted for small cells filtered through a 25 μm mesh. This size fraction represented between 77 and 91 % of the total POC pool. At stations A3-2 and E-5, we also used unfiltered seawater for our uptake experiments. Similar results were obtained for both the 0.2–25 μm and unfiltered fractions at station A3-2.

⁷ Includes bacterial and mesozooplankton contributions.

⁸ Mean of all samples collected in the mixed layer.

⁹ Assumes that only the soluble iron atmospheric supply is available (see text).

¹⁰ A negative value indicates an additional iron requirement.

¹¹ At stations R-2 and A3-2, the negative values most likely occurred due to differences in the timescales of observations and calculations of fluxes (parameters were decoupled in time). The iron budget was based on an instantaneous picture of different fluxes that were not strictly measured at the same time (i.e. export fluxes operated on a different time frame to the iron supply (vertical, lateral and atmospheric) and were very large at R-2 and A3-2).

¹² *fe*: uptake of new / (uptake of new + regenerated iron F_e): biogenic iron export / (uptake of new + regenerated iron) (Boyd et al., 2005). Note the *fe* and F_e ratios have considerable plasticity due to uncertainties in the lithogenic vs. biogenic fraction of exported particulate iron and the missing iron source at A3-2.

¹³ Calculated using the biological uptake of new iron (k) and molar Fe/C cellular uptake ratio (j).

¹⁴ Net primary production (NPP) integrated within the euphotic zone down to 1 % PAR, based on ¹³C incorporation (Cavagna et al., 2014).

servative estimate since the winter dFe stock was probably an underestimation (as evidenced by lower dFe in the mixed layer compared to deep waters at A3-1; Qu erou e et al., 2015). Biogenic iron (defined as the Fe associated with living phytoplankton and phytoplankton biodetritus) was calculated by assuming that all particulate phosphorus (P) was of biogenic origin and multiplying the mean particulate P concentration in the mixed layer at each station by a maximum intracellular Fe / P ratio of $1.9 \text{ mmol mol}^{-1}$ for natural phytoplankton assemblages measured by Twining et al. (2004) for Fe-replete conditions. These calculations follow methods reported in Planquette et al. (2013) and assume that particulate P is not (in part) derived from local rock weathering. Van der Merwe et al. (2015) have tested this assumption using Fe / P ratios in the Kerguelen Islands basalts and the upper continental crust and note that the ~ 1000 -fold increase in pP within suspended particles can only be explained by pP produced in situ within the mixed layer from dissolved PO_4^- . Lithogenic Fe was calculated by assuming that all particulate aluminium (pAl) was of lithogenic origin and by multiplying the mean pAl concentration in the mixed layer by a lithogenic Fe / Al ratio of $0.36 \text{ mol mol}^{-1}$, which is the mean value based on basaltic rocks from the Crozet region (0.51; Gunn et al., 1970) and crustal materials (0.2; Wedepohl, 1995). The selected Fe / Al ratio of $0.36 \text{ mol mol}^{-1}$ is also very similar to that of 0.33 used extensively in earlier calculations (Taylor and McLennan, 1989) and reported for the deep Atlantic Ocean (Sherrell and Boyle, 1992). This approach is dependent not only on the chosen Fe / Al ratio but also assumes that processes other than biological assimilation, such as adsorption and scavenging onto organic particles, photoreduction, surface precipitation, and chemically and biologically driven dissolution, are not significant (Measures et al., 2008; Planquette et al., 2011, 2013; Ellwood et al., 2014). Since biogenic and lithogenic Fe were calculated independently, their sum may be less than the observed total particulate Fe concentration. This is likely due to plasticity in the chosen Fe / Al and Fe / P ratios and differential remineralisation rates for Fe, Al and P. Nevertheless, our estimates of biogenic and lithogenic Fe provide a perspective on the relative contributions to the total pFe pool.

Reference and plume waters contained roughly an equal fraction of biogenic and lithogenic Fe. The origin of this biogenic Fe pool will be a combination of biological uptake of dFe, physical adsorption onto suspended biological particles, and conversion from the lithogenic fraction (likely driven by microbes), with these processes operating on different timescales (Boyd et al., 2005; Frew et al., 2006; Planquette et al., 2011). By contrast, the plateau stations contain 19–69 times more lithogenic Fe than biogenic Fe, consistent with the supply from the nearby sediments of the plateau and Heard Island, as suspected by Zhang et al. (2008), van Beek et al. (2008) and discussed in Chever et al. (2010). The measurement of other geochemical “fingerprint” particulate tracers (such as Al, Mn) on the plateau confirmed the provenance

of Fe supplied from the Kerguelen shelf sediments in the particulate phase (van der Merwe et al., 2015).

3.2.2 Internal iron supply

Vertical fluxes were calculated as follows. A vertical diffusivity (K_z) at the base of mixed layer of $10^{-5} \text{ m}^2 \text{ s}^{-1}$ was used for the plume and reference site, and a K_z of $3 \times 10^{-4} \text{ m}^2 \text{ s}^{-1}$ was used for the plateau site, estimated from the Shih parameterisations (Shih et al., 2005) using the Thorpe scale method (Park et al., 2014b). These values are comparable to K_z values estimated for KEOPS 1 using the Osborn model ($4 \times 10^{-4} \text{ m}^2 \text{ s}^{-1}$; Osborn, 1980) (Park et al., 2008a). Vertical diffusivity was multiplied by the vertical dFe gradient for each profile, which was determined using the linear part of the vertical profiles corresponding to the 150–200 m depth strata in Fig. 3a–f, consistent with calculations for KEOPS-1 (Blain et al., 2007). Vertical diffusivity of Fe was negligible at reference and plume sites but significant on the plateau due to both the higher vertical diffusivity and the steeper Fe gradient between 150 and 200 m.

Upwelling was defined as the vertical velocity (w_{ek}) multiplied by the dFe concentration at 200 m, which corresponds to the depth of the remnant winter water. The magnitude of vertical velocity in this region has recently been studied by Rosso et al. (2014), who used the MIT general circulation model to examine the sensitivity of the vertical velocity to the horizontal resolution. They found clear differences in w_{ek} due to the development of near-surface sub-mesoscale frontal structures that only their highest-resolution model was able to resolve. Rosso et al. (2014) reported vertical velocities for individual water parcels in excess of 100 m day^{-1} in the Kerguelen region, with w_{ek} stronger in the downstream plume. Both the horizontal and vertical circulations were much weaker over the plateau since it acts as a natural barrier to the strong ACC fronts coming from the west. Unfortunately, no seasonal cycle was included in the model forcing. Therefore the temporal root mean square of the vertical velocity reported in Fig. 12b of Rosso et al. (2014) was used for the plateau and plume sites ($w_{\text{ek}} = 0.5$ and 1 m d^{-1} , respectively), and a conservative value of $w_{\text{ek}} = 0.13 \text{ m d}^{-1}$ for the open Southern Ocean (used by de Baar et al., 1995; originally reported in Gordon et al., 1977) was chosen for our reference station. Although w_{ek} was lower on the plateau compared to the plume, the higher dFe concentration at 200 m resulted in comparable estimates of upwelled Fe (Table 1).

Entrainment of Fe by episodic (intraseasonal) deepening of the mixed layer has rarely been taken into account in field studies (Frants et al., 2013) due to the absence of data characterising the short-term variability of the mixed layer depth, yet a recent compilation of observations (Nishioka et al., 2011; Tagliabue et al., 2014) and modelling studies (Mongin et al., 2008) suggests that entrainment could be a major vertical supply mechanism fuelling surface biomass (Carranza et al., 2015). We used more than 6000 vertical profiles of

salinity and temperature collected in the KEOPS-2 regions of interest to estimate the seasonality of the mixed layer depth and its variability (Supplement Fig. 3). We derived the vertical supply of Fe by entrainment via hypotheses regarding the relation between the size of mixed layer depth excursions and their frequency (see the Supplement). Entrainment data based on transient deepening of the mixed layer was not available for station R-2; therefore we calculated this by multiplying the dFe concentration in winter water (which reflects the dFe concentration of the winter mixed layer) by the winter mixed layer depth (MLD) and assume that this entrainment event happens once per year. “Detrainment” at R-2 was accounted for by multiplying this new entrainment flux by the summer-to-winter MLD ratio.

For the vertical fluxes, in spring on KEOPS-2, entrainment was the dominant vertical Fe flux term on the plateau, delivering $\sim 70\%$ of the total vertical supply and tripling the total vertical flux in comparison to budgets that neglect this process. At the plume and reference sites, entrainment was comparable to the upwelling flux. Vertical diffusion accounted for 4–8% of the total vertical supply on the plateau. In contrast, the contribution from dFe entrainment was much reduced in late summer on KEOPS-1 (42 and $8.8 \text{ nmol m}^{-2} \text{ d}^{-1}$ for plateau and plume, respectively) due to the deepening and weakening of the ferricline (Fig. 4b). The relative magnitude of the total vertical Fe supply terms at the three study sites was plateau > plume > reference (Table 1, row d).

For the lateral fluxes, the horizontal supply at reference station R-2 was assumed to be zero since HNLC waters upstream and downstream of this station contained similar dFe and pFe concentrations, and as phytoplankton growth and biomass was low at this site, there would be little biogenic Fe exported below the mixed layer. On the plateau, Fe supply at station A3 was taken from the steady-state box model of Chever et al. (2010), which used the horizontal dFe gradient and current velocities from Park et al. (2008a) to calculate the lateral flux of $180 \text{ nmol m}^{-2} \text{ d}^{-1}$ in the 0–150 m depth band above the plateau; note that this model used KEOPS-1 data. Lateral transport into the plume E stations was assumed to originate from Fe-fertilised plateau waters that were advected offshore (d’Ovidio et al., 2015). This value was estimated by assuming that horizontal stirring occurs in a Lagrangian framework and using altimetry-derived geostrophic velocities to determine transport across the plateau boundary. We also used a depth band of 0–150 m, considered as the winter mixed layer in the plume over the season. These estimates were combined with direct measurements of the dFe content of three different types of on-plateau stations to calculate the lateral flux over a 3-month supply period prior to the spring bloom, namely (i) two coastal stations near the Kerguelen Islands occupied on KEOPS-2 (stations TEW-1 and TEW-2), (ii) one coastal station close to Heard Island occupied during KEOPS-1 (station C1) and (iii) the central plateau station A3 considered here. This resulted in $5.4 \times 10^7 \text{ mol Fe}$ per day being injected into a plume size

(defined at a threshold of $> 0.3 \mu\text{g Chl } a \text{ L}^{-1}$ and identified from satellite images) of $2.5 \times 10^{11} \text{ m}^2$ over 90 days in spring (full details of the calculations are contained in d’Ovidio et al., 2015). This equated to a lateral flux into the plume of $2400 \text{ nmol m}^{-2} \text{ d}^{-1}$ in the October–November period.

By combining our in situ Fe measurements with estimated ages of the water bodies in the plume, we calculate a first-order exponential scavenging removal constant between 0.041 and 0.058 d^{-1} , which equated to a residence time of 17 to 24 days, consistent with estimates based on the Fe inventory and Fe export in free-floating traps (15–79 days; Laurenceau-Cornec et al., 2015). Since the total of the vertical and lateral fluxes in the plume were more than double those on the plateau, this may imply that the source waters supplying the plume from the northern Kerguelen Islands shelves (which had a uniquely narrow *T-S* class in surface waters; Grenier et al., 2015) were richer in Fe than the plateau further south at A3. This is supported by observations of dFe in the surface ocean at stations TEW-1 and TEW-2 ($1.2\text{--}1.8 \text{ nmol L}^{-1}$), which were close to Hillsborough Bay in waters only 86 m deep (Qu  rou   et al., 2015).

Considering only internal processes (diffusion, upwelling, entrainment, lateral transport) in supplying Fe to the surface mixed layer, the vertical terms dominated at the reference station; vertical terms were 6-fold greater than lateral terms on the plateau, whereas lateral advection was the dominant term in the plume (4–5-fold greater than the vertical terms). Since the particulate Fe stocks were abundant in surface waters (above the winter temperature minimum layer) and significantly higher than the dissolved pools (most notably on the plateau), it is likely that a fraction of the suspended lithogenic pFe from Heard Island or the Kerguelen Plateau sediments also contributed to the internal dFe supply and fuelled biological responses. This is discussed in more detail later.

3.2.3 External iron supply

Data on atmospheric Fe fluxes through dust deposition and the solubility of Fe in the dust for all three study sites were taken from the nearby land-based sampling site “Jacky” ($49^\circ 18' 42.3'' \text{ S}$, $70^\circ 07' 47.6'' \text{ E}$; altitude 250 m) on the Kerguelen Islands, as reported in Heimburger et al. (2012, 2013a). Mean total Fe fluxes taken over the period 24 November 2008 to 07 September 2010 were $500 \pm 390 \text{ nmol m}^{-2} \text{ d}^{-1}$ (Heimburger et al., 2013a), which was comparable to the Crozet region upstream ($895 \text{ nmol m}^{-2} \text{ d}^{-1}$; Planquette et al., 2007) and the Southern Ocean sector south of Australia ($288\text{--}488 \text{ nmol m}^{-2} \text{ d}^{-1}$; Bowie et al., 2009) but greater than the flux value estimated by Wagener et al. (2008) during KEOPS-1 ($14\text{--}46 \text{ nmol m}^{-2} \text{ d}^{-1}$). The remoteness of the Kerguelen region means it receives low quantities of atmospheric material (Heimburger et al., 2012; Wagener et al., 2008), the majority of which is crustal in origin, such as desert dust from South America, South Africa or Australia (Prospero et al., 2002;

Mahowald, 2007; Bhattachan et al., 2012), although local anthropogenic activities, rock outcrops and exposed soil may also impact dust fluxes.

Atmospheric fluxes were dominated by wet deposition (Heimburger et al., 2012). Heimburger et al. (2013b) calculated the mean “soluble” Fe deposition flux (defined as $< 0.2 \mu\text{m}$) using a median solubility of $82 \pm 18\%$ in rainwater on the Kerguelen Islands. These high solubilities were attributed to the remoteness of the sampling location from dust sources resulting in strong cloud chemical processing during transport. However, the solubility of Fe dissolved in seawater at higher pH will be much lower (Schroth et al., 2009; Sedwick et al., 2007). Hence a conservative value of 10% of Fe that is released into seawater was chosen (Baker et al., 2006; Mackie et al., 2006) for our budgets here, resulting in a soluble Fe atmospheric deposition flux to the Kerguelen region of $50 \text{ nmol m}^{-2} \text{ d}^{-1}$ (Table 1, row f). This value was lower than the internal vertical supply on the plateau (~ 20 -fold) and plume (~ 10 -fold), insignificant compared to the lateral supply to the plume but comparable to the lateral supply on the plateau. Although volcanic ash has not been considered here for atmospheric Fe supply, this term may have played an important role for primary productivity on the Kerguelen Plateau during the middle Miocene climate transition (Abrajevitch et al., 2014).

3.2.4 Iron export

Downward Fe and C fluxes were measured directly in free-floating sediment P-traps at the plateau (A3-2) and plume (E-1, E-3, E-5) stations and estimated using the ^{234}Th fluxes and Fe/Th ratios at the reference site (R-2) (Planchon et al., 2015). The sinking of pFe was by far the greatest loss term in our budgets, with $5746 \text{ nmol m}^{-2} \text{ d}^{-1}$ of total Fe exported from the mixed layer on the plateau, between 895 and $4579 \text{ nmol m}^{-2} \text{ d}^{-1}$ exported at the plume stations and $1302 \text{ nmol m}^{-2} \text{ d}^{-1}$ exported at the reference station. The flux of sinking pFe decreased from station E-1 to E-3 to E-5, concurrent with the seasonal progression of the bloom and indicating the mixed layer assemblages were efficiently recycling Fe under strong grazing pressure (Laurenceau-Cornec et al., 2015). The downward total pFe fluxes were greater than the sum of the vertical, lateral and atmospheric dFe supply on the plateau but were generally less in the plume.

Aluminium was used as a normaliser to estimate the fraction of lithogenic Fe in the exported material. The percentage lithogenic fraction of total pFe exported at the E stations remained much the same at each deployment (34–39%), whereas the lithogenic fraction was a much larger component at A3-2 (51%), reflecting the close proximity to sources of particulate material rich in Fe. The Fe/Al ratio of exported material was higher at E stations (1.0–1.1) and on the plateau A3-2 (0.87) compared to the Fe/Al ratio of lithogenically dominated particles (0.2; Wedepohl, 1995), confirming that a significant amount of exported Fe was biogenic in origin.

Interestingly, the Fe/Al export ratios were similar to those associated with suspended particles at E stations (0.9–1.2) but lower than the Fe/Al of suspended particles at A3-2 (1.2). This suggests that the biota associated with the plateau bloom at A3 were more capable of efficiently recycling and retaining biogenic particulate Fe in the mixed layer (through rapid turnover to prevent aggregation and sinking) compared to lithogenic particulate Fe, which had a shorter residence time and was preferentially exported to depth. This may be due to greater ballasting of the lithogenic particles (Ellwood et al., 2014) and is consistent with other export studies which have shown that biologically processed particles have longer residence times than lithogenic particles in the mixed layer (Lamborg et al., 2008a). Since P may be lost from exported particles much faster than Fe due to bacterial remineralisation and zooplankton consumption (Schneider et al., 2003; Lamborg et al., 2008b), it was not appropriate to apply a biogenic normaliser to the P-trap data as this may underestimate the biogenic Fe component of particles captured in the traps.

Iron export fluxes were greater during the spring study of KEOPS-2 compared to the late summer study of KEOPS-1 (Table 2). This difference between the KEOPS studies was also observed in Fe uptake rates (Fourquez et al., 2015). Such observations may be simply related to the seasonal supply; in other words, greater Fe supply in spring resulted in greater Fe uptake and export. Determined pFe sinking fluxes were also greater than those observed during CROZEX study (Planquette et al., 2011), the SAZ-Sense expedition south of Tasmania (Bowie et al., 2009) and the FeCycle-I expedition east of New Zealand (Frew et al., 2006) and of similar magnitude to those reported by Bowie et al. (2001) during the Southern Ocean Iron Release Experiment (SOIREE) ($5.2 \mu\text{mol m}^{-2} \text{ d}^{-1}$), but they were much lower than Ellwood et al. (2014) reported for FeCycle-II.

The export of particulate organic carbon (POC) into our P-traps followed the same trend as that of pFe at the E stations, decreasing from 7.0 ± 2.3 at E-1 to $2.0 \pm 1.0 \text{ mmol m}^{-2} \text{ d}^{-1}$ at E-5 (Table 1). Despite the higher pFe vertical fluxes at A3-2, POC export was lower than the E stations. The C export fluxes at 200 m at A3-2 using our P-traps (Laurenceau-Cornec et al., 2015) were similar to results estimated from ^{234}Th deficits by Planchon et al. (2015; 2.3 ± 0.7 and $3.8 \pm 0.8 \text{ mmol C m}^{-2} \text{ d}^{-1}$, respectively). The comparison of these POC fluxes to results (for the A3 plateau site only) obtained during KEOPS-1 illustrates highly dynamic variations, reflecting the rapid decline in biomass during autumn (Blain et al., 2007). Specifically, P-trap measurements of POC fluxes at 200 m during KEOPS-1 decreased from 3.7 ± 0.3 to $1.3 \pm 0.3 \text{ mmol C m}^{-2} \text{ d}^{-1}$ over two visits to A3 in February 2005 (Trull et al., 2008), whereas estimates based on ^{234}Th at this time, reflecting the previous ~ 30 days of export, suggested much higher values ($25 \pm 7 \text{ mmol C m}^{-2} \text{ d}^{-1}$; Savoye et al., 2008). These variations illustrate the difficulty of constraining budgets in temporally evolving systems, providing a cautionary note to our efforts. Addi-

tional discussion of temporal and spatial export flux variations during KEOPS-2 is provided in Laurenceau-Cornec et al. (2015) and Planchon et al. (2015).

3.2.5 Biological iron recycling

Intracellular Fe uptake by phytoplankton and bacteria $> 0.2 \mu\text{m}$ (Fourquez et al., 2015) was measured at stations A3-2 and E-5 when the bloom was rapidly growing (Cavagna et al., 2014). Iron uptake fluxes were similar on both the plateau (A3-2) and in the plume (E-5), ranging between 1120 and 1745 $\text{nmol m}^{-2} \text{d}^{-1}$. If we assume that the Fe uptake rate of 28.1 $\text{pmol L}^{-1} \text{d}^{-1}$ measured at E-5 (Fourquez et al., 2015) was conservative at E stations, 0.17 nmol L^{-1} of Fe could have been consumed in surface waters between the occupations of stations E-4E and E-5. This is consistent with the observed decrease in surface dFe concentrations from 0.19 to 0.06 nmol L^{-1} at E-4E and E-5, respectively (F. Quéroué, personal communication, 2013). The net and gross demand calculated at A3 during KEOPS-1 (204 and 408 $\text{nmol m}^{-2} \text{d}^{-1}$, respectively; Sarthou et al., 2008) is approximately 3–5 times smaller than the intracellular Fe uptake at A3-2 during KEOPS-2 for a similar C biomass (mean value of 12.7 and 10.3 $\mu\text{mol L}^{-1}$ POC in surface at KEOPS-1 and KEOPS-2, respectively; Cavagna et al., 2014), perhaps indicating luxury uptake as well as important differences in community composition and activity (primary production). These studies provide the opportunity to compare KEOPS-2 to KEOPS-1 data and generate a general picture of the seasonal progress from early spring to late summer, assuming that interannual and spatial variability is low, which may not be the case (Grenier et al., 2015).

The bacterial and mesozooplankton contributions to Fe regeneration were calculated separately (Table 3). Volumetric values varied between 0.06 and 0.59 $\text{pmol Fe L}^{-1} \text{d}^{-1}$ and between 0.02 and 0.08 $\text{pmol Fe L}^{-1} \text{d}^{-1}$ for bacterial and mesozooplankton Fe regeneration, respectively. The mesozooplankton rates were much lower than for KEOPS-1 because there were much fewer individuals (0.26–0.56 per L, compared to about 1–6 individuals per L for KEOPS-1; see Fig. 2 in Carlotti et al., 2008). Total Fe regeneration fluxes ranged from 10 (R-2) to 71 (A3-2) $\text{nmol m}^{-2} \text{d}^{-1}$.

A similar Fe regeneration calculation was also performed based on the C budget by using the percentage of gross community production (GCP) that is remineralised for KEOPS-2 and results from Fe uptake experiments described above. This yielded higher Fe regeneration estimates in the range of 1–11 $\text{pmol L}^{-1} \text{d}^{-1}$. Specifically, for station A3-2, 23 % of GCP was remineralised, and therefore the Fe regeneration flux in the mixed layer was 1119 $\text{nmol m}^{-2} \text{d}^{-1}$. Similarly, for station E-5, 34 % of GCP was remineralised, resulting in an Fe regeneration flux of 504 $\text{nmol m}^{-2} \text{d}^{-1}$. Since the Fe regeneration fluxes based on the C budget are much greater (~ 16 times) than those calculated using the first approach,

this suggests that the remineralisation efficiency for Fe regeneration appears to be less than that of C.

Iron regeneration fluxes can be compared with those from KEOPS-1 using the same first approach above. For station A3 on KEOPS-1, this resulted in a Fe regeneration flux of 1 $\text{pmol L}^{-1} \text{d}^{-1}$ in surface waters. Malits et al. (2014) also calculated the release of bacterial bound Fe by viral lysis (0.42 $\text{pmol L}^{-1} \text{d}^{-1}$), which was the dominant loss term during KEOPS-1 (Brussaard et al., 2008). This value compared to 1.5 $\text{pmol L}^{-1} \text{d}^{-1}$ determined in zooplankton grazing experiments (Sarthou et al., 2008), suggesting that grazing and microbial Fe cycling were in a similar range, and the total Fe regeneration was between 2–3 $\text{pmol L}^{-1} \text{d}^{-1}$ for KEOPS-1.

Importantly, Fe regeneration was much lower during the early compared to late bloom stage and was dominated by bacterial regeneration in spring (60–90 % of total Fe regeneration). Strzepek et al. (2005) estimated Fe regeneration rates (during FeCycle-II) for herbivores (16.5–18.4 $\text{pmol L}^{-1} \text{d}^{-1}$), bacterivores (15–25.5 $\text{pmol L}^{-1} \text{d}^{-1}$) and viruses (0.4–28 $\text{pmol L}^{-1} \text{d}^{-1}$), which is equivalent to a total Fe regeneration rate of 1435–3236 $\text{nmol m}^{-2} \text{d}^{-1}$ for a 45 m mixed layer. Bowie et al. (2009) estimated Fe regeneration to be 261–1206 $\text{nmol m}^{-2} \text{d}^{-1}$ for the SAZ-Sense study. So our determined KEOPS-2 mixed-layer Fe regeneration rates (71 and 31 $\text{nmol m}^{-2} \text{d}^{-1}$ at A3-2 and E-5, respectively) were on the lower end of the range reported in other sectors of the Southern Ocean, and clearly insufficient to meet demand (measured as Fe uptake) at all stations, indicating a reliance on new Fe supply. This is discussed in more detail below.

3.3 Sequestration efficiencies: iron-to-carbon ratios

The mixed-layer phytoplankton intracellular Fe / C uptake ratios were calculated directly from deck board incubations for stations A3-2 (0.007 mmol mol^{-1}) and E-5 (0.021 mmol mol^{-1}) (Table 1). These values are similar to those reported for other natural and artificial iron fertilisation studies in the Southern Ocean, including for Fe-limited conditions during the Southern Ocean Iron Experiment (SOFEX; 0.01 mmol mol^{-1} ; Twining et al., 2004), and those inside the KEOPS-1 plateau bloom (0.005 $\text{mmol Fe mol C}^{-1}$; Sarthou et al., 2008) but lower than those reported for SAZ-Sense (0.06–0.07 mmol mol^{-1} ; Bowie et al., 2009).

Suspended mixed layer Fe / C ratios (Table 1) were significantly higher than phytoplankton intracellular uptake ratios. This finding is likely the result of the contribution of lithogenic and detrital Fe to suspended material Fe / C ratios and is consistent with the removal of C at a faster rate than that of Fe and with Fe being added through new sources after phytoplankton uptake. Differences may also arise because of luxury uptake, the timescale of integration in deck board experiments compared to Fe / C ratios in ocean suspended and sinking particles (which are broadly similar – see below), and/or the fact that our system was not in steady state.

Table 2. Fluxes of iron and carbon (mean and standard deviation, SD) exported in sinking particles (trap deployed at 200 m) and ratio of Fe / C in sinking (traps) and suspended mixed layer (ISP) particles at stations A3-2 and E-stations. There was no successful trap deployment at station R-2. A comparison to previous studies is provided.

Site	PFe flux ($\mu\text{mol m}^{-2} \text{d}^{-1}$)		POC flux ($\text{mmol m}^{-2} \text{d}^{-1}$)		Fe/C (sinking) (mmol mol^{-1})		Fe/C (suspended) (mmol mol^{-1})
	mean	SD	mean	SD	mean	SD	mean
KEOPS-2							
A3-2	5.75	1.20	2.23	0.68	2.57	0.97	1.51
E-1	4.58	1.38	7.02	2.28	0.65	0.52	0.49
E-3	1.89	0.29	4.87	1.54	0.39	0.29	0.39
E-5	0.90	0.36	2.00	1.00	0.45	0.13	0.33
KEOPS-1 ¹							
A3-initial	0.33	0.05	3.60	0.43	0.09	–	–
A3-final	0.20	0.02	1.36	0.39	0.15	–	–
C5	1.51	0.32	1.57	0.08	0.96	–	–
CROZEX ²							
North	0.84	–	15.9	–	2.55	–	0.69
South	0.23	–	12.9	–	0.57	–	0.31
SAZ-Sense ³							
P1	0.17	0.09	3.34	1.81	0.05	0.04	0.04
P2	0.07	0.01	2.11	0.88	0.04	0.02	0.06
P3	0.21	0.05	0.86	0.38	0.25	0.13	0.03
FeCycle-1 ⁴							
F1-80 m	0.22	0.03	n.d.	–	n.d.	–	0.04
F1-120 m	0.36	0.05	2.09	0.03	0.17	–	–
F2-80 m	0.55	0.06	2.51	0.17	0.22	–	–
F2-120 m	0.35	0.03	2.10	0.01	0.17	–	–
FeCycle-2 ⁵							
A1-100 m	5.0	0.7	11	–	0.45	–	0.78
A1-200 m	7.3	1.6	5.8	–	1.26	–	–
A2-100 m	10	1.0	42	–	0.24	–	1.12
A2-200 m	10	9	6.8	1.8	1.47	–	–
A3-100 m	17	2	12	2	1.42	–	0.63
A3-200 m	10	1	14	–	0.71	–	–
A4-100 m	20	8	9.3	0.9	2.15	–	0.86
A4-200 m	15	6	6.1	1.8	2.46	–	–
Other literature data							
Mixed plankton assemblages ⁶	–	–	–	–	–	–	0.01–0.05
Iron-limited algae ⁷	–	–	–	–	–	–	0.01
Iron-replete algae ⁷	–	–	–	–	–	–	0.02–0.05
Southern Ocean synthesis ⁸	–	–	–	–	0.01–0.06	–	–

n.d.: no data

¹ Data for particles > 0.2 μm (Blain et al., 2007; Bowie et al., unpublished data).

² Data for > 53 μm particles only (Planquette et al., 2011). Downward Fe fluxes were estimated from samples collected from in situ pumps using ²³⁴Th depletions and Fe / Th ratios in sinking particles. Waters to the north of Crozet Island were downstream of the islands and iron fertilised, whilst those to the south were upstream HNLC conditions. The Fe / C from bioassay culturing experiments conducted during CROZEX was 0.25 mmol mol^{-1} (Moore et al., 2008).

³ Data for particles > 1 μm (Bowie et al., 2009).

⁴ Data for particles > 0.4 μm (Frew et al., 2006). Only one mixed layer Fe / C ratio was reported. The biogenic Fe / C mixed layer ratio was estimated to be 0.004–0.012 mmol mol^{-1} .

⁵ Data for particles > 0.4 μm , except deployment A1 (> 2 μm) (Ellwood et al., 2014). The mixed layer Fe / C ratios were calculated from Table 4 using the sediment trap deployment periods reported in Table 3 in the original publication.

⁶ Estimates of Fe / C for diatoms and whole plankton assemblages compiled by de Baar et al. (2008), with optimal ratios for growth tending towards the upper end of the range.

⁷ Intracellular ratio reported for HNLC polar water south of New Zealand during SOFeX (Twining et al., 2004).

⁸ Ratio of dFe supply to POC export, synthesis by Morris and Charette (2013).

Table 3. Iron regeneration rates based on bacterivore and herbivore contributions.

Site	Bacterial ($\text{pmol L}^{-1} \text{d}^{-1}$)	Mesozooplankton ($\text{pmol L}^{-1} \text{d}^{-1}$)	Total Fe regeneration ($\text{pmol L}^{-1} \text{d}^{-1}$)	Percentage bacterial contribution	Total integrated mixed layer Fe regeneration ($\text{nmol m}^{-2} \text{d}^{-1}$)
R-2	0.06 ± 0.01	0.04	0.10	61	10
A3-1	0.10 ± 0.03	0.02	0.12	87	19
A3-2	0.43 ± 0.07	0.03	0.46	93	71
E-1	0.33 ± 0.02	0.04	0.37	88	27
E-3	0.54 ± 0.04	0.06	0.60	90	23
E-5	0.59 ± 0.03	0.08	0.67	88	31

Also, since a Ti-citrate-EDTA wash was used to remove extracellular surface Fe during the incubation experiments but not from particles collected in the ISPs and P-traps, our suspended and sinking pFe concentrations include Fe present within cells, adsorbed to cell walls, detrital Fe and lithogenic Fe. This would tend to increase Fe / C in suspended particles. Differences between intracellular and suspended mixed-layer Fe / C ratios may also derive from the C term, since the ISP sampling includes detrital material as well as living cells. We also note that suspended pFe data are the sum of 1–53 and > 53 μm size fractions collected by ISPs and thus may also include some sinking particles. This may affect the suspended Fe / C ratios.

In addition to the ratio of “total” particulate (biogenic + lithogenic) Fe over POC (Fe / C) in suspended particles discussed above, we also calculated the ratio of biogenic Fe over POC (i.e. $\text{Fe}_{\text{bio}} / \text{C}$) following methods discussed in Sect. 3.2.1. Profiles are shown in Fig. 5. Suspended Fe / C ratios were remarkably similar at all E stations and station R-2 but higher on the plateau at A3 stations (Table 1). We also observed generally surface-to-deep increases in Fe / C ratios in suspended particles at all stations (Fig. 5), consistent with earlier findings (Frew et al. 2006). The vertical profiles of $\text{Fe}_{\text{bio}} / \text{C}$ showed a similar structure at the three study sites, with a general decreasing trend from the surface to sea floor (opposite to that of Fe / C); note that a constant Fe / P was used to estimate the Fe_{bio} component. These findings indicate that Fe is preferentially retained within, and adsorbed to, sinking particles (i.e. scavenging drives the total Fe / C ratio), but biogenic Fe is recycled at a faster rate compared to C, similar to macronutrients N and P. A preferential loss of C relative to Fe from sinking material implies that an external input of Fe is required to sustain a downward flux of carbon.

At station R-2, the Fe / C ratio peaked at 500 m, most likely due to lithogenic particulate Fe input (and not C) from the Leclaire Rise (see above) (note this peak was not seen in the $\text{Fe}_{\text{bio}} / \text{C}$ ratios). At E stations, the Fe / C ratio showed maximum values in mesopelagic intermediate waters in the 600–1000 m depth range. We also believe this was due to the lateral transport of lithogenic particulate Fe (and not C)

from the plateau (seafloor at ~ 600 m) into the plume. This is supported by the absence of this feature in the $\text{Fe}_{\text{bio}} / \text{C}$ ratios for E stations. Fe / C ratios in deep waters were much higher at A3 stations ($26\text{--}38 \text{ mmol mol}^{-1}$) compared to R-2 (4 mmol mol^{-1}) and E stations ($5\text{--}7 \text{ mmol mol}^{-1}$), indicating enrichment of lithogenic particulate Fe above the plateau. Some fraction of this lithogenic Fe will be accessible to the biota and then be incorporated into the biogenic Fe pool. This is confirmed by modification in the Fe / Al ratio (van der Merwe et al., 2015). Inclusion of the biologically available fraction of the lithogenic Fe flux is therefore required to calculate fully the yield of carbon exported per unit Fe injected, consistent with Planquette et al. (2011) and Pollard et al. (2009).

Interestingly, although Fe / C ratios varied greatly between stations ($0.2\text{--}37 \text{ mmol mol}^{-1}$), the $\text{Fe}_{\text{bio}} / \text{C}$ ratio fell within a narrow band ($0.01\text{--}0.08 \text{ mmol mol}^{-1}$ for all stations and depths), which encompasses the elemental ratios of Fe-replete ($0.04 \text{ mmol mol}^{-1}$) and Fe-limited ($0.01 \text{ mmol mol}^{-1}$) large diatoms (Sunda and Huntsman, 1995; de Baar et al., 2008). This highlights the tight coupling between Fe_{bio} and POC in the absence of new sources of Fe and allow us to estimate the relative remineralisation efficiencies for Fe versus C. The $\text{Fe}_{\text{bio}} / \text{C}$ data contrast with the findings of Planquette et al. (2011) for the CROZEX study who observed variable $\text{Fe}_{\text{bio}} / \text{C}$ ratios to the north of Crozet (Fe-fertilised region), which were on average much higher than those found to the south (Fe-limited region). The fraction of Fe_{bio} relative to lithogenic Fe in particles collected below the mixed layer also depends on the stage of the bloom, the nature and magnitude of the supply of new lithogenic particles, and the rate of conversion from lithogenic-to-biogenic Fe (Lam et al., 2006; Frew et al., 2006; Lam and Bishop, 2008). These factors are highly variable in the Kerguelen region and this explains the wide range of Fe_{bio} -to-total Fe values in particles observed during KEOPS-2.

The Fe / C export ratio of sinking particles in our traps were similar to suspended mixed-layer ratios for the E stations but slightly higher at A3-2 (Fig. 5), possibly due to the sinking of recently supplied lithogenics over the plateau.

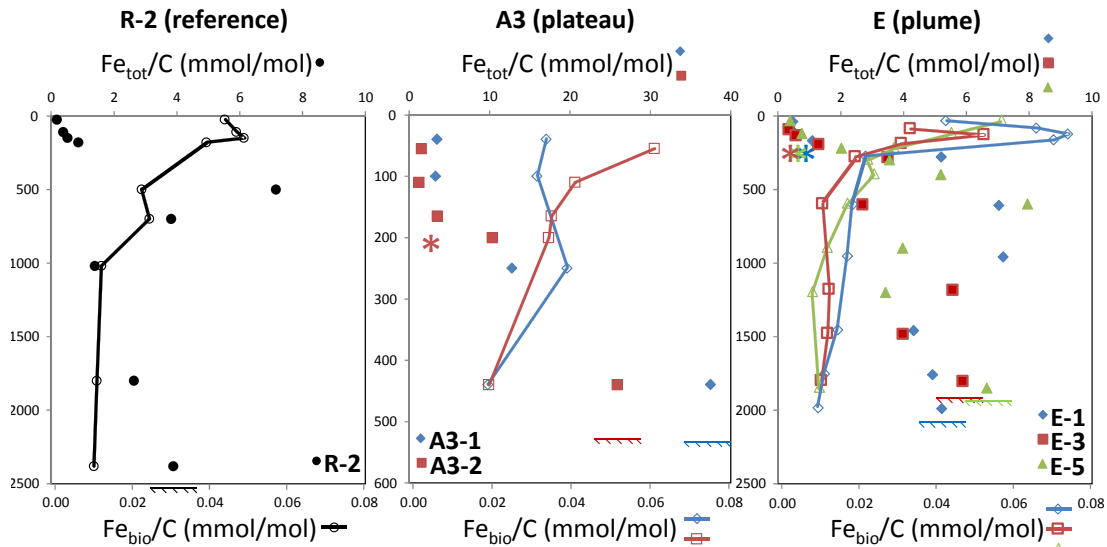


Figure 5. Vertical profiles of Fe / C ratios in suspended (ISP) and sinking (P-trap) particles. Solid symbols indicate total Fe / C (i.e. ratio of biogenic + lithogenic Fe over POC) and joined open symbols indicate Fe_{bio} / C (i.e. ratio of biogenic Fe only over POC; calculated using P as a normaliser). The asterisk markers (*) show the export total Fe / C ratio (P-traps). Note the different scale on the x axis for Fe / C at A3 stations.

Both pFe and POC export fluxes decreased during bloom development at E stations, indicating the mixed layer became more retentive for both Fe and C. This is consistent with the picture that emerges from the E time series from primary and export production estimates, which show that production was moderate and matched by the moderate export during our visits (Planchon et al., 2015; Trull et al., 2015; Cavagna et al., 2014).

Since POC export fluxes during spring (KEOPS-2) were similar to late summer (KEOPS-1), but pFe export fluxes were higher in spring compared to summer (Table 2), this resulted in a generally higher carbon sequestration efficiency (lower Fe / C) during late summer, consistent with a rapidly exporting ecosystem during bloom decline. The exported particles may have been dominated by more lithogenics and may have been much more processed in KEOPS-2 compared to KEOPS-1, where the system had already run out of Fe. It was also expected that growing communities during KEOPS-2 would retain dFe through luxury uptake, which may also result in observed generally higher Fe / C ratios in sinking particles during the spring bloom (KEOPS-2, FeCycle-II) compared to austral summer conditions (KEOPS-1, CROZEX, FeCycle-I; Blue Water Zone; Morris and Charette, 2013) (Table 2 and Fig. 6).

Morris and Charette (2013) presented a detailed synthesis of ^{234}Th -derived POC export and dFe budgets in studies where natural iron fertilisation fuels the substantial phytoplankton blooms observed in the Southern Ocean. Where data are available to calculate the seasonal Fe / C ratios, 1 order of magnitude variation (0.006–0.06) is observed between different Southern Ocean regions. It is likely that Fe / C ra-

tio variations (Table 2) reflect both experimental methodologies, different calculation approaches, observational limitations and system complexities. Le Moigne et al. (2014) have also recently shown that variability in the carbon sequestration efficiency is related to the mode of Fe delivery.

3.4 Iron supply vs. demand

Using calculated flux estimates, a comparison of Fe supply and demand at the three sites around the Kerguelen Archipelago in spring was possible (Fig. 7). In our short-term iron biogeochemical budgets, the total dFe supply from new sources (calculated as the sum of diffusion, upwelling, vertical and lateral advection, and atmospheric dust) to surface waters of the plume was more than twice that above the plateau and > 20 times greater than at the reference station (Table 1). The Fe demand (measured by cellular Fe uptake) in the plume was similar (1.5 times greater) to the plateau but > 40 times greater than at the reference station. New Fe supply was 14–94 times greater than recycled Fe supply (“iron remineralisation”; row i in Table 1) from bacterial regeneration and zooplankton grazing. This contrasts with the findings of Bowie et al. (2009) for SAZ-Sense who reported recycled fluxes that were broadly comparable with new Fe supply in the SAZ in summer at study sites further from natural iron fertilisation.

Since Fe supply from new sources was greater than the Fe demand (uptake minus remineralisation as a recycled Fe source) at all stations (R-2, A3-2 and E-5), this resulted in a positive value for row k in Table 1 (i.e. there was no additional Fe required to balance the dissolved budget). This find-

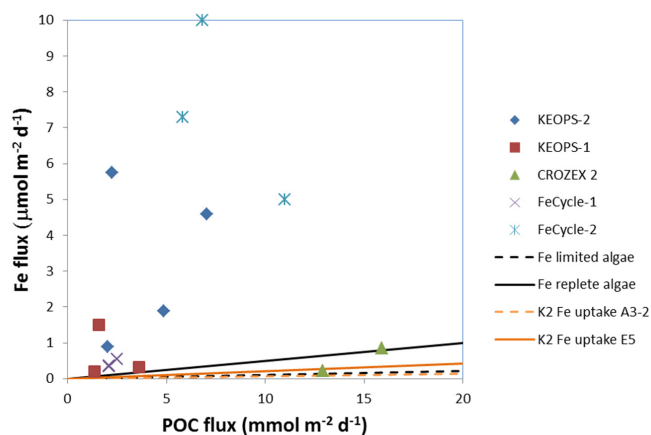


Figure 6. A comparison of export fluxes of pFe versus POC in sinking particles for natural iron fertilisation studies in the Southern Ocean. For details of the sampling methods, refer to Table 2 and the original articles. The lines indicate Fe / C ratios for Fe-limited (black dashed) and Fe-replete (black solid) phytoplankton (Twining et al., 2004) and the mean mixed-layer intracellular Fe / C ratios at stations A3-2 (orange dashed) and E5 (orange solid) on KEOPS-2 (taken from Table 1). FeCycle-II had complex biogeochemical dynamics due to a storm event and subsequent deep-water mixing (during sediment trap deployment at their station A3), splitting the study into two phases (“eddy centre” and “eddy periphery”). To aid interpretation of Fe / C export data in the context of iron fertilisation, only data from the pseudo-Lagrangian phase 1 (i.e. deployments A1 and A2 during bloom development and export) from that study are included in this plot (Ellwood et al., 2014).

ing is consistent with other observations at both the plateau and plume sites which were Fe replete in early spring but somewhat surprising for the HNLC reference site R-2. This may partly be a result of an overestimate of the atmospheric supply used in calculations presented here from literature data. Another explanation is that the parameters used in our “short-term” iron budget calculations are decoupled in time (e.g. there will be an offset between the mechanisms for organism acquisition of Fe and the processes resulting in Fe-laden particles leaving the upper ocean), and the short-term Fe budget is based on an “instantaneous picture” of different fluxes that were not in steady state.

Interestingly, at station A3-2, the sink processes (Fe export and uptake) are so large and the regenerated Fe flux so small that the total (dissolved + particulate) Fe losses are far greater than the net dFe supply (Fig. 7a). In other words, to a first order the budget is not balanced with known sources of Fe insufficient to account for the downward flux, even if we only accounted for the non-lithogenic particulate Fe export flux (row 1 in Table 1). Assuming all flux calculations to be correct within the estimated error bounds in Table 1, this implies that there is a missing flux term in the budget at A3 and this is likely lithogenic pFe from the Kerguelen Plateau and/or Heard Island (and this may be converted to biogenic Fe). Currently, we do not invoke a lithogenic pFe

to dFe transfer in the budget, which could increase the Fe supply on the plateau significantly, although at present we do not know what fraction of particulate material is converted into the dissolved form. This will vary largely with the mineralogy (Schroth et al., 2009), provenance of the particles and seawater characteristics (e.g. organic complexation; Shaked and Lis, 2012). Indeed, Thuróczy et al. (2012) previously measured organic complexation in Antarctic waters and discussed the role of ligands in transporting and dissolving pFe into dFe, using theoretical data provided by Borer et al. (2005).

Applying a solubility of 2.5 % used for KEOPS-1 at A3 to enable Fe supply to meet demand (Blain et al., 2007) would provide an extra 10–34 $\mu\text{mol m}^{-2}$ of dFe to the mixed layer over the plateau in KEOPS-2, approximately doubling the dFe standing stock. These values are comparable to our observations and suggest that particulate material plays a major role in the supply of dFe (van der Merwe et al., 2015). Further, if we assume pFe from glacial melt is delivered over a 3-month period, this would provide an additional 111–387 $\text{nmol m}^{-2} \text{d}^{-1}$ to the mixed layer at A3, values of similar magnitude to the individual vertical and lateral supply terms. Whilst a dissolution estimate of 2.5 % may be considered to be at the upper extent of the range, Schroth et al. (2009) have reported that 2–3 % of Fe is soluble in glacial flour, which can remain suspended in surface water for several months after delivery from the Kerguelen or Heard islands.

The release of Fe to biota via the conversion of lithogenic to biogenic Fe has been previously suggested (Lam et al., 2006; Frew et al., 2006; Borer et al., 2009; Planquette et al., 2011) and the present work strongly supports this hypothesis, with our data (Fig. 5) indicating that biogenic Fe has a longer residence time in the upper ocean than lithogenic Fe which is not accessed by biota. The role of pFe in supplying bioavailable Fe is also supported by the similarity of the pFe and dFe profile shapes in Fig. 3, which infer that pFe may be contributing to the control of dFe, either by supplying it or because biogenic particles are controlling both.

Finally, our estimation of Fe supply and regeneration allowed us to estimate an *fe* ratio, defined by Boyd et al. (2005) as $fe = \text{uptake of new Fe} / (\text{uptake of new} + \text{regenerated Fe})$. For the plume region, *fe* was 1.4 (Table 1). This was higher than the *fe* ratio calculated for KEOPS-1 (0.49; Sarthou et al., 2008), which at that time was comparable to the average *f*-ratio for nitrogen of 0.41 (corresponding to NO_3^- uptake / (NO_3^- uptake + NH_4^+ uptake); Mosseri et al., 2008), indicating that both NH_4^+ and regenerated Fe could support export production. Conversely, the KEOPS-2 *f*-ratio was higher (up to 0.9; Cavagna et al., 2014), indicating that primary production was mainly sustained by nitrate uptake. The *fe* ratios for both KEOPS studies were much higher than the *fe* ratio estimated during FeCycle-I (0.17, Boyd et al., 2005) and SAZ-Sense (0.06–0.16; Bowie et al., 2009). This confirms that, in the Kerguelen region, there are sufficient new

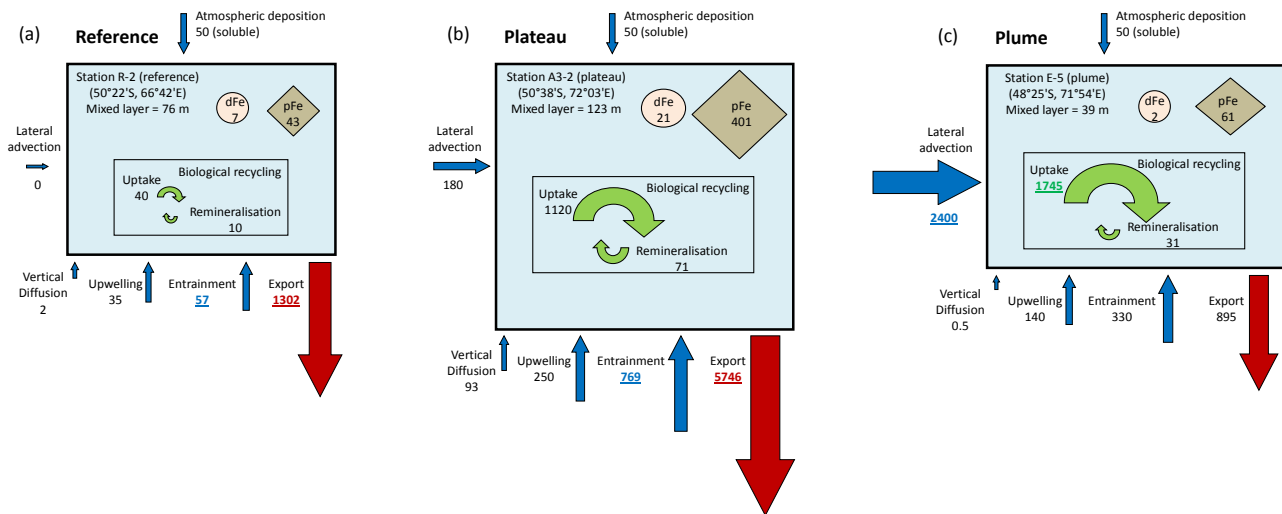


Figure 7. Biogeochemical iron budgets for the reference (R-2, **a**), plateau (A3-2, **b**) and plume (E-5, **c**) stations. Iron pools are given in $\mu\text{mol m}^{-2}$ and iron fluxes in $\text{nmol m}^{-2} \text{d}^{-1}$. Iron sources are shown as blue arrows, sinks as red arrows and the green arrows indicate biological Fe cycling. The size of the arrows is roughly proportional to the magnitude of the Fe fluxes, with major fluxes shown as bold underlined text.

sources of Fe delivered on a seasonal timescale (predominantly via intraseasonal entrainment, winter mixing, lateral transport and particulate Fe dissolution) available to sustain the massive bloom observed in spring.

4 Conclusions

The complex regional circulation, multiple iron sources, and transport pathways above and downstream of the naturally fertilised Kerguelen Plateau region results in a mosaic of phytoplankton blooms. The budgets presented here result from direct measurements of the Fe inventories and fluxes between different pools. The system was not in steady state during the period of the KEOPS-2 observations, and the exchange of Fe between the dissolved, biogenic and lithogenic pools was highly dynamic in time and space. Our analysis highlights the important role of pFe, the inherent heterogeneity and biogeochemical differences associated with particulates within and exported below the mixed layer and the lithogenic to biogenic conversion pathways.

This study also highlights the significance not only of the mode of Fe fertilisation on the plateau (predominantly vertical) versus the plume (predominantly lateral) but also of the relative magnitude. Importantly, since the Fe supply from new sources to the plume was more than double that above the plateau, this implies that the waters that supply the plume are not the same as those at station A3 on the southern plateau, and the plume must be supplied with water from the northern part of the plateau or Kerguelen coastal waters, which are richer in dFe (Qu  rou   et al., 2015; Trull et al., 2015). This source of Fe, which will contain a large frac-

tion of particulate material (van der Merwe et al., 2015) that is transported off the Kerguelen Plateau, is therefore an important but previously unquantified contribution to the downward flux of Fe exiting the upper ocean in the plume. Moreover, the KEOPS-2 results are tightly linked to the mode of Fe supply that is different from dust deposition or purposeful additions and to the concomitant supply of major nutrients, and this has consequences for the carbon sequestration efficiency of the system. When Fe supply is predominantly vertical (as it is at station A3), then the C sequestration efficiency is lower (i.e. higher Fe / C) as C would be resupplied to the mixed layer as well as Fe. This coupling has important implications for geoengineering schemes that propose to increase the supply of Fe to surface waters by pumping waters from below.

Future efforts should focus on the quantification of the full seasonal cycle of Fe delivery, which will be fundamental to closing the iron budget around the Kerguelen Archipelago on annual timescales. This will allow the assessment of the important longer-term climatic and ecosystem implications with changes in the nature and strength of Fe supply with physical (weakening overturning circulation, warming, increased stratification) and chemical (ocean acidification, deoxygenation) environmental forcings, together with increases in glacial melt, rainfall and dust deposition on a warming planet.

The Supplement related to this article is available online at [doi:10.5194/bg-12-4421-2015-supplement](https://doi.org/10.5194/bg-12-4421-2015-supplement).

Author contributions. A. R. Bowie designed the iron budgets, performed the calculations and prepared the manuscript with contributions from all co-authors. P. van der Merwe, F. Quéroué, G. Sarthou, F. Chever and A. T. Townsend collaborated on trace metal sampling, analyses and interpretation; M. Fourquez and I. Obernosterer were responsible for biological cycling, T. Trull for carbon dynamics and the P-trap deployments, F. Planchon for Th-based export and J.-B. Sallée for vertical flux estimates. S. Blain designed the overall KEOPS-2 project and helped with budget calculations.

Acknowledgements. We thank the captain B. Lassiette, officers and crew of RV *Marion Dufresne*, Pierre Sangiardi (Institut Paul-Émile Victor) and the Institut National des Sciences de l'Univers for voyage logistics and their support of the science, and voyage leader Bernard Quéguiner (Institut Méditerranéen d'Océanologie) and chief scientist Stéphane Blain (LOMIC, Université Pierre et Marie Curie) for leading the KEOPS-2 expedition. We acknowledge Thomas Rodemann (UTAS) for CHN analysis in the Central Science Laboratory, together with members of the ISP and P-trap teams for support at sea. Michael Ellwood (Australian National University) kindly loaned the TMR for the project. Access to ICP-MS was provided through Australian Research Council LIEF funds (LE0989539). The altimeter and colour and temperature products for the Kerguelen area were produced by Ssalto/Duacs and CLS with support from CNES and kindly prepared by Emmanuel Laurenceau-Cornec (UTAS) and Francesco d'Ovidio (LOCEAN – IPSL, Université Pierre et Marie Curie). We thank Isabella Rosso (Australian National University) for useful discussion.

This KEOPS-2 project was supported by the French Research program of INSU-CNRS LEFE-CYBER (“Les enveloppes fluides et l’environnement” – “Cycles biogéochimiques, environnement et ressources”), the French ANR (“Agence Nationale de la Recherche”), SIMI-6 program, ANR-2010-BLAN-614 KEOPS2 and, ANR-10-JCJC-606 ICOP), the French CNES program (“Centre National d’Etudes Spatiales”) and the French Polar Institute IPEV (Institut Polaire Paul-Émile Victor). The Australian participation in the project was supported by the Antarctic Climate and Ecosystems Cooperative Research Centre and a University of Tasmania “Rising Stars” award to the lead author.

We thank two anonymous reviewers for their very constructive comments, which improved our manuscript.

Edited by: B. Quéguiner

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