



Microbial Competition in the Subpolar Southern Ocean: An Fe–C Co-limitation Experiment

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Iron (Fe) is a paradox in the modern ocean – it is central to many life-critical enzymes but is scarce across most surface waters. The high cellular demand and low bioavailability of Fe likely puts selective pressure on marine microorganisms. Previous observations suggest that heterotrophic bacteria are outcompeted by small diatoms for Fe supply in the subantarctic zone of Southern Ocean, thereby challenging the idea of heterotrophic bacteria being more competitive than phytoplankton in the access to this trace metal. To test this hypothesis, incubation experiments were carried out at the Southern Ocean Time Series site (March-April 2016). We investigated (a) whether dissolved organic carbon (DOC), dissolved Fe, or both limit the growth of heterotrophic bacteria and, (b) if the presence of potential competitors has consequences on the bacterial Fe acquisition. We observed a pronounced increase in both bulk and cell-specific bacterial production in response to single (+C) and combined (+Fe+C) additions, but no changes in these rates when only Fe was added (+Fe). Moreover, we found that +Fe+C additions promoted increases in cell-specific bacterial Fe uptake rates, and these increases were particularly pronounced (by 13-fold) when phytoplankton were excluded from the incubations. These results suggest that auto- and heterotrophs could compete for Fe when DOC limitation of bacterial growth is alleviated. Such interactions between primary producers and nutrient-recyclers are unexpected drivers for the duration and magnitude of phytoplankton blooms in the Southern Ocean.

Keywords: iron, carbon, Southern Ocean, competition, heterotrophic bacteria, pico-nanoplankton, Fe uptake, bacterial production

INTRODUCTION

Geological timescales have enabled microbes to develop adaptive solutions to an evolving environment, and iron (Fe) had a fundamental role in the metabolic pathways that emerged (Falkowski and de Vargas, 2004; Hunter and Boyd, 2007). As life appeared in an oxygen-free environment, the primordial ocean provided sufficient concentrations of readily available

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Fe (Hunter and Boyd, 2007). However, the concentration of dissolved Fe (DFe) decreased drastically after two major irreversible oxygenation events (~2.4 billion years and ~542 million years ago, Ilbert and Bonnefoy, 2013) and Fe is found at trace levels in most of today's ocean surface (<0.5 nM, Johnson, 1997). Despite these changes, the Fe demand in marine microorganisms remained high. Nature retained this versatile metal, that can have a wide range of oxidation states, as an integral part for a wide range of proteins throughout evolution and many of these proteins are irreplaceable agents for vital cellular metabolic activities (oxygen transport, electron transport, DNA synthesis, etc., Morel and Price, 2003).

In phytoplankton, the photosynthetic transport chain is one of the most prominent Fe-dependent processes. One single copy of a photosystem requires 23–24 atoms of Fe, and overall 80% of Fe is allocated to the photosynthetic transport chain in a cell (Raven et al., 1999; Strzepek and Harrison, 2004; Behrenfeld et al., 2006). In heterotrophic bacteria the respiratory chain accumulates more than 90% of the intracellular Fe (Tortell et al., 1999; Andrews et al., 2003). Phytoplankton and bacteria play important roles in the ocean and have direct influence on global biogeochemical cycles. Considering that phytoplankton drive ocean $\rm CO_2$ sequestration via photosynthesis and downward export, while heterotrophic bacteria control much of the oceanic release of $\rm CO_2$ via respiration, the outcome of a competition for Fe could influence the direction and magnitude of carbon fluxes in the upper ocean.

Despite widespread interest in microbial Fe requirements (Sarthou et al., 2005; Twining and Baines, 2013; Blain and Tagliabue, 2016; Strzepek et al., 2019), there is no consensus regarding the minimum Fe requirements for phytoplankton or heterotrophic bacteria. There are two reasons to explain why this question has not been resolved. First, there have been few studies on the Fe requirements of heterotrophic bacteria compared with those for phytoplankton. Second, the wide range of Fe content relative to C biomass (Fe:C ratio) that exist for phytoplankton does not favor conclusive comparison (Blain and Tagliabue, 2016, and references herein). What is clear, however, is that DFe in the oceans is overwhelmingly complexed (99%, Rue and Bruland, 1997) by strong organic ligands with evidence of them containing Fe-binding functional groups consistent with biologically produced siderophores (Macrellis et al., 2001; Gledhill and Buck, 2012). Marine microbes have evolved different mechanisms to cope with the diversity of the Fe-binding ligand pool, and the capacity to acquire enough Fe for survival in a "highly diffusive" open ocean provides a competitive edge (Desai et al., 2012; Hopkinson and Barbeau, 2012; Toulza et al., 2012). The capacity to produce siderophores is generally confined to heterotrophic bacteria (Armstrong et al., 2004), but the ability to take up siderophores may be more widespread than previously thought, and extend to the phytoplanktonic realm (Hogle et al., 2016; Kazamia et al., 2018; McQuaid et al., 2018). Recent studies have revealed that distinct siderophores and strategies are being employed by heterotrophic bacteria (Boiteau et al., 2016, 2019; Bundy et al., 2018; Debeljak et al., 2019). But to date, no eukaryotic phytoplankton have been found to produce or release siderophores. For this reason heterotrophic bacteria are

commonly reported as highly efficient competitors, especially in severely Fe-limited environments (Braun and Killmann, 1999).

The Southern Ocean is the largest High Nutrient, Low Chlorophyll (HNLC) region in the world ocean, mainly because of Fe limitation. In the Southern Ocean, chlorophyll levels remain low year-round, but phytoplankton blooms occur in areas in the vicinity of land masses (Blain et al., 2007). During austral spring 2011, the KEOPS2 project aimed at exploring different phytoplankton blooms east of Kerguelen island. Over the course of the bloom, the release of dissolved organic carbon (DOC) derived from primary production increased the Fe demand of heterotrophic bacteria (Fourquez et al., 2015) which were Fe-limited (Obernosterer et al., 2015). The availability of a labile C source may have led to a higher bacterial Fe demand. These findings raised the hypothesis that labile organic carbon exacerbated the potential competition between smallsized phytoplankton cells (pico- and nanophytoplankton) and heterotrophic bacteria for Fe (Fourquez et al., 2015). The present work aimed to test the above hypothesis. For this, our experimental design was based on the joint assumption that (1) heterotrophic bacteria are outcompeted for Fe by pico-and nanophytoplankton and (2) that DOC availability to heterotrophic bacteria influences the strength of this relationship.

MATERIALS AND METHODS

Site Description

This study was carried out as part of the V02-IN2016 voyage of the R. V. Investigator (March 11 to April 17, 2016). During the expedition, we visited on two occasions (March 19 and 29) the Southern Ocean Time Series (SOTS, 47°S, 142°E) site that is located within a low current region in the subantarctic Zone (SAZ) north of the subantarctic Front (SAF) that marks the northern edge of the Antarctic Circumpolar Current (Figure 1). This area represents a large portion of the total area of the Southern Ocean that serves as a strong sink for atmospheric CO₂. Conditions at SOTS are typical and representative of the Indian sector SAZ, from ∼90 to 145°E. The absence of Fe is regarded as the primary cause that restricts primary production and constraints the biological pump in the area (Sedwick et al., 1997; Cassar et al., 2011; Trull et al., 2019). The relief of the Fe limitation can occur by aerosol Fe supply in summer in the region that differs in this way from mechanisms of deep mixing and/or sediment input/resuspension that enhances Fe concentrations in surface waters at the vicinity of subantarctic islands such as Kerguelen (Blain et al., 2008; Rembauville et al., 2015).

Experimental Strategy

The study involved two separate sets of incubation experiments: Experiment 1 and 2 (**Figure 2** and **Supplementary Figure 1**). Because there is a potentially confounding influence of Fe and C limitation on bacterial processes, the objective of Experiment 1 was to first determine whether Fe, C, or both are limiting or colimiting factors at SOTS while the objective of Experiment 2 was to determine whether the presence of larger cells (especially picoand nanoplankton) influences bacterial activities. In addition to

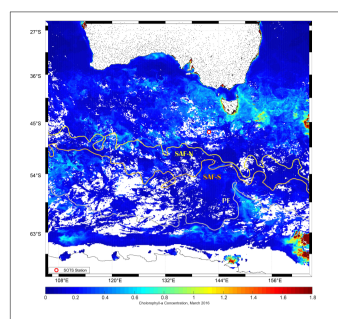


FIGURE 1 | Study site of R/V Investigator voyage 02. Dot represents location of the SOTS site where experiments were conducted (42°S, 47°S, 51°S, and 54°S at 141°E). SOTS was visited on two occasions on 19 (Experiment 1) and March 29, 2016 (Experiment 2). SAF-N, SAF-S stands for the northern and southern branches of the subantarctic front, and PF for polar front, respectively. Sea surface height was used to characterize fronts position (AVISO Satellite Data Altimetry). Sea surface Chl a composite image for March 2016 was derived from the Moderate Resolution Imaging Spectroradiometer aboard NASA's Aqua satellite and obtained from the ocean color data distribution site (https://oceandata.sci.gsfc.nasa.gov/MODIS-Aqua/Mapped/Monthly/4km/chlor_a/).

the one control that consisted in unamended nutrient seawater (no addition), the following three treatments were prepared as triplicates: +Fe, +C, and +Fe+C. Iron (+Fe) was added as FeCl₃ (final concentration of FeCl₃ 1 nM), and carbon (+C) was added as trace-metal clean glucose (final concentration of glucose 10 μ mol L⁻¹). Hence, the addition was 16.6 μ molFe molC⁻¹ to attain the bacterial Fe quota observed in Fe-replete bacterial cultures (16.1 \pm 2.3 μ molFe molC⁻¹, Fourquez et al., 2014). For Experiment 1, incubations were performed directly on unfiltered seawater (bacteria were incubated with micro and picoand nanoplankton communities). Additional incubations were performed on 20 µm-prefiltered seawater (<20 µm condition), and on 1 µm-prefiltered seawater (<1 µm condition) for Experiment 2. We use the term "condition" throughout the manuscript to refer to the different size fractionation treatments (summary in **Table 1**). The biological response of heterotrophic bacteria was monitored from sub-samples drawn from these incubation bottles, and analyzed for several parameters as described in sections below.

Sampling Procedures

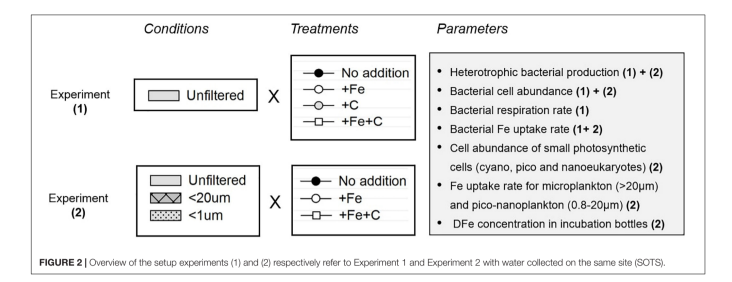
Seawater was pumped from the surface ocean (\sim 5 m depth) using a trace metal clean towed fish sampler. Samples were collected directly into a trace metal clean laboratory (clean room, ISO5) where 300 mL of seawater was dispensed into 500 mL

TABLE 1 | List of abbreviations used.

Abbreviation	Explanation
Unfiltered	Raw seawater. Microplankton, pico-nanoplankton and heterotrophic bacteria were incubated together
<20 μm	Seawater prefiltered on 20 µm mesh-size. Microplankton was excluded from incubation. Pico-nanoplankton and heterotrophic bacteria were incubated together
<1 μm	Seawater prefiltered on 1 μ m mesh-size. Heterotrophic bacteria were incubated alone

acid-washed polycarbonate (PC) bottles under a laminar flow hood (ISO class 5). Overall, 12 (for Experiment 1) and 54 (for Experiment 2) independent replicates were prepared in bottles capped and sealed with Parafilm. All plastic materials used were acid-washed following GEOTRACES procedures in our home laboratory (GEOTRACES cookbook Cutter et al., 2017). Briefly, PC bottles were soaked for 1 week in the alkaline detergent Decon 90, then rinsed four times with deionized water and three times with ultrapure water. They were subsequently filled with 10% hydrochloric acid (Suprapur, Merck) for 1 week. After that time, bottles were rinsed five times with highly purified water. The PC bottles were dried, and UV sterilized for 15–30 min under a laminar flow hood and then stored in triple plastic bag before being used.

To minimize risks of potential contamination of samples with metals or dissolved organic matter as an artifact of filtration in preparation for Experiment 2, seawater was filtered at very low pressure (<5 Hg). All incubations were performed at in situ temperature (13.5°C). Bottles were placed in a water bath within a controlled temperature room (13.5°C) to avoid temperature fluctuations. For Experiment 1, incubations were performed in total darkness. For Experiment 2, incubations were conducted under 12:12 light-dark condition, and we employed neutral density screens to attenuate the light intensity. Low levels of light can reduce rates of photosynthesis and the release of DOC associated and may alter Fe uptake rates in photosynthetic cells. Therefore, we opted to use low light intensity (average 4.5 μ mol photon m⁻² s⁻¹, ~1W) in Experiment 2 to (a) increase potential competition for Fe stocks between autotrophic picoand nanoplankton cells and heterotrophic bacteria, and to (b) avoid stimulation of large phytoplankton growth and subsequent organic enrichment artifacts. For subsampling, incubation bottles were transferred to the clean container and opened under a laminar flow hood (ISO class 5). Subsamples from each triplicate for bacterial abundance and production were taken at T0, T1 (+24 h), T2 (+48 h), T3 (+72 h) and at T4 (+110 h) for Experiment 1 and at T0, T1 (+36 h), and T2 (+120 h) for Experiment 2. The time points for Experiment 2 were chosen following results collected during Experiment 1. For both experiments bacterial heterotrophic production, cell abundance, and Fe uptake were measured at several time points. We also measured heterotrophic bacterial respiration at the end of the incubation in Experiment 1 to estimate the bacterial growth efficiency in the different treatments (Figure 2). Methods for each parameter measured are detailed in sections below.



Cell Abundance

For each biological replicate, 4.5 mL subsamples were fixed with glutaraldehyde (0.5% final concentration), kept in the dark at $4^{\circ}\mathrm{C}$ for 20 min and then shock-frozen in liquid nitrogen. The samples were stored at $-80^{\circ}\mathrm{C}$ until analyses by flow cytometry. Flow cytometry was performed following the protocols in Marie et al. (1997, 2005). Frozen samples were rapidly thawed in a water bath at 70°C for 3 min and aliquots taken for autotrophic and/or prokaryote cell counts. Sample aliquots were kept on ice in the dark and promptly analyzed on a Becton Dickinson FACScan flow cytometer fitted with a 488 nm laser. Ultrapure water was used as sheath fluid for all analyses. Before and after each run, samples were weighed to \pm 0.0001 g to determine the volume of sample analyzed.

Samples for autotrophic cell abundance were prepared by aliquoting 998 μL of sample into a clean 5 mL polycarbonate tube, with 2 μL of PeakFlow Green 2.5 μL beads (Invitrogen) added as an internal fluorescence and size standard. Each sample was run for 5 min at a high flow rate of $\sim\!40~\mu L$ min $^{-1}$. Autotrophic cell populations were separated into regions based on their chlorophyll autofluorescence in red (FL3) versus orange (FL2) bivariate scatter plots. Synechococcus cells were determined from their high FL2 and low FL3 fluorescence. Pico- and nano-phytoplankton communities were determined from their relative cell size in side scatter (SSC) versus FL3 fluorescence bivariate scatter plots. Final cell counts in cells L^{-1} were calculated from event counts in the identified regions and analyzed volume.

Samples for prokaryote cell abundance were prepared by aliquoting 995 μL of sample to a clean 5 mL polycarbonate tube. Samples with high prokaryote cell counts were diluted to 1:10 with 0.2 μm filtered seawater (FSW) to remove underestimation of cell concentration from coincidence (100 μL sample in 900 μL FSW). Cells were stained for 20 min with 5 μL of SYBR Green I (Invitrogen) at a final dilution of 1:10,000. An additional 2 μL of PeakFlow Green 2.5 μL beads (Invitrogen) was added to the sample as an internal fluorescence and size standard. Each sample was run at a low flow rate of

 ${\sim}12~\mu L~min^{-1}$ for 3 min and prokaryote cell abundance was determined from bivariate scatter plots of SSC versus green (FL1) fluorescence. Final cell counts in cells L^{-1} were calculated from event counts in the identified regions and analyzed volume.

Heterotrophic Bacterial Production

Bacterial production was estimated by [³H] leucine incorporation applying the centrifugation method (Martinez et al., 1996) as described in Obernosterer et al. (2008). Briefly, 1.5-mL samples were incubated with a mixture of radioactive leucine, L-[3,4,5-3H(N)] (PerkinElmer, specific activity 123.8 mCi.mol⁻¹) and non-radioactive leucine at final concentrations of 20 nM. Duplicates plus one "killed sample" were incubated in the dark at the respective in situ temperatures for 2-3 h. Linearity of leucine incorporation over this time period was tested in parallel and at two occasions (at the beginning of Experiment 1 and 2). Incubations were terminated by the addition of trichloroacetic acid (TCA; Sigma) to a final concentration of 5%. To facilitate the precipitation of proteins, bovine serum albumin (BSA; Sigma, 100 mg L⁻¹, final concentration) was added prior to centrifugation at 16,000 g for 10 min (Van Wambeke et al., 2002). After discarding the supernatant, 1.5 mL of 5% TCA solution was added and the samples were subsequently vigorously shaken on a vortex and centrifuged again. The supernatant was discarded again and 1.5 mL of UltimaGoldTM uLLt (PerkinElmer) was finally added. The radioactivity incorporated into bacterial cells was counted in Hidex 300SL Liquid Scintillation Counter. A factor of 1.55 kg C mol leucine⁻¹ was used to convert the incorporation of leucine to carbon equivalents, assuming no isotope dilution (Kirchman et al., 1993). Isotopic dilution ranged from 1.0 to 1.3 as determined on three occasions using a kinetic approach.

Fe Uptake Rates

Following the subsampling for bacterial production, Fe uptake experiments were initiated by adding $0.2 \text{ nmol } L^{-1}$ at final

concentration of 55Fe (as 55FeCl₃, PerkinElmer specific activity 2.46×10^3 Ci mol⁻¹) after 36 and 120 h of incubations (independent replicates). After a 24-h incubation with ⁵⁵Fe, microorganisms were filtered through a stack of nitrocellulose filters (Whatman) of 0.2 and 0.8 µm porosity, separated with 20 µm mesh filters. These filter porosities were chosen to separate phytoplankton (including Synechococcus, >0.8 µm) from heterotrophic bacteria (0.2-0.8 µm). Before running dry, the filters were rinsed with 0.2-µm filtered Ti(III) citrate EDTA solution (Tovar-Sanchez et al., 2003) for 2 min to dissolve any extracellular Fe, followed by three consecutives rinses with 5 mL of 0.2 μm filtered seawater for 1 min (Fourquez et al., 2012, 2015). The filters were placed into plastic vials and 10 mL of the scintillation cocktail Filtercount (Perkin Elmer) was finally added. Vials were agitated for 24 h before the radioactivity was counted with the Hidex 300SL scintillation counter. Radioactivity on filters was corrected for background (55Fe adsorbed into the filter and/or onto particles and not being efficiently washed out by the washing solution) using ⁵⁵Fe-radiotracer medium with dead cells, also called "killed control". Killed controls were treated the same as above but microorganisms were fixed with 1% of glutaraldehyde (left for 1 h at 4°C) prior addition of ⁵⁵Fe. DFe concentration in each incubation bottle was also assessed prior incubation with ⁵⁵Fe. Subsamples (~40 mL) were measured by flow injection with online preconcentration and chemiluminescence detection (adapted from Obata et al., 1993). An internal acidified seawater standard was measured every day in order to control the stability of the analysis. The detection limit was 40 pmol kg⁻¹ and the accuracy of the method was controlled by analyzing the SAFe S (0.110 \pm 0.036 nmol kg^{-1} (n = 3); consensus value 0.093 ± 0.008 nmol kg^{-1}), and SAFe D1 (0.66 \pm 0.06 nmol kg⁻¹ (n = 4); consensus value 0.67 ± 0.04 nmol kg⁻¹) seawater standards. DFe concentration were employed to correct 55Fe uptake estimates from cold DFe present in incubation bottle at the time of the measurement. In incubation bottles, the DFe concentration changed over time also suggesting that there was remineralization taking place. To correct for this, we have multiplied the results from these experiments by the calculated proportion of 55Fe from the total DFe. Calculation details can be found in Fourquez et al. (2015).

Bacterial Respiration

Rates of respiration were determined from dissolved oxygen consumption in 24 h dark incubations at the end (T4, + 110 h) of Experiment 1 using Winkler titration method. In order to keep the bacteria only for the measurement, all samples were carefully pre-filtered onto 0.8 μ m acid-washed PC filter. Two out of the three biological replicates belonging to each treatment (control, +Fe, +C, and +Fe+C) were used for measurement. Last replicate was employed as a T0 by adding manganese chloride followed by alkaline iodide prior incubation. To estimate the consumption in dioxygen (O₂), the amount of O₂ measured in bottles after 24 h of incubation was subtracted from T0 measurement.

All incubation bottles were opened and gently shaken under flow laminar hood to optimize O₂ level inside and homogeneity between treatments. After these steps, the samples were subsequently transferred into cleaned and acid-washed glass biological oxygen demand stoppered bottle. Incubations lasted for 24 h in the dark in a temperature-controlled incubator set at 13.5°C (in situ temperature). At the end of the incubation time, subsampling (1.8 mL) for flow cytometry analysis was taken quickly just prior to add the reactive. Dissolved oxygen concentration was measured based on the whole-bottle modified Winkler titration of Carpenter (1965) plus modifications by Culberson (1991). Bacterial respiration rates were normalized to C biomass by considering the bacterial cell abundance in each incubation bottle.

Carbon Biomass and Conversion Factor

Direct carbon contents for pico-nanoplankton and microplankton were estimated from particulate organic carbon (POC) measurements. In total, 3 L of seawater sample were first filtered through 20 μm and subsequently passed through 1.2 μm (diameter 25 mm) Sterlitech silver membrane filters, and dried at 60°C. The samples were acidified, dried and analyzed by high-temperature combustion (1000°C) to determine POC. The analysis for total nitrogen, carbon, and hydrogen was determined using a Thermo Finnigan EA 1112 Series Flash Elemental Analyzer.

In parallel, the carbon content was also indirectly estimated using conversion factors. Photosynthetic pico-nanoplankton cell abundance was converted to carbon biomass using constant cell-to-carbon conversion factors based on the literature. Conversion factors used were, respectively 255 and 2590 fgC cell⁻¹ for the cyanobacterium *Synechococcus* and for picoeukaryotes (Buitenhuis et al., 2012), and 183 fgC cell⁻¹ for nanoeukaryotes (Caron et al., 1994, 2017). We assume the cyanobacterium *Synechococcus* represented the majority of the resident cyanobacteria. For heterotrophic bacteria, the carbon content was estimated using 12.4 fgC cell⁻¹ as reported by Fukuda et al. (1998).

Statistical Analyses

All statistical comparisons were performed using one-way analysis of variance (ANOVA) and a post hoc Tukey test. Differences were considered statistically significant at p < 0.05. To evaluate the differences between treatments, statistics were individually performed between nutrient unamended (control) and amended treatments (+Fe, +C, and +Fe+C). We also evaluated statistical differences between conditions by comparing unfiltered (control) to other conditions (<20 μ m or <1 μ m).

RESULTS

Environmental Settings of the Study Site

As is typical for HNLC regions, at SOTS site the C biomass was dominated by small cells ($<20~\mu m$) which represented

 $1.51~\mu mol~L^{-1}$ of the $1.90~\mu mol~L^{-1}$ total POC in surface waters. The concentration of DFe was $0.081~\pm~0.02~nM$ at 5 m depth. Concentration of dissolved organic carbon (DOC) were not measured during the cruise. Nevertheless, the Southern Ocean surface waters exhibit a DOC concentration range of $\sim\!40{\text -}50~\mu molC~L^{-1}$ (Hansell et al., 2009). If we consider the upper range of 50 $\mu molC~L^{-1}$ to be representative of what can be found at SOTS, the resulting DFe:DOC molar ratio was $\sim\!1.62~\mu mol~mol^{-1}$.

Experiment 1: Responses of Bacteria to Fe and C Additions

Cell-Specific Bacterial Production

In Experiment 1, we investigated the responses of bacteria to Fe, C and concomitant Fe and C additions. Sole additions of Fe did not result in any significant enhancement of bulk nor cellspecific BP (Figure 3). In accordance with these results, bacterial abundance in the Fe-amended treatment did not differ from the control (Figure 3A). However, single (+C) and combined (+Fe+C) additions of C significantly stimulated bulk and cellspecific BP. We note a pronounced response in cell-specific BP to single (+C) and combined (+Fe+C) additions of carbon over the time of the experiment (1.3-56-fold and 1.6-26-fold higher than control in +C and +Fe+C, respectively, Figure 3B). The results from treatments +Fe+C and +C were not statistically different from each other apart from T3 (+ 72 h, p = 0.012). The bacterial cell abundance also increased, however, the magnitude of the stimulation was less than that measured for the BP. Cell abundance is a complex function between growth and mortality rates; this decoupling is therefore not surprising. At the end of the experiment, the enhancement of these parameters was still detectable in the +C and +Fe+C treatments but a decrease in cell-specific BP was also observed after 72 h of incubation.

Bacterial Respiration and Growth Efficiency

Bacterial respiration (BR) rates were measured at the end of the Experiment 1. BR varied from 0.39 ± 0.15 and 1.63 ± 0.34 fmolO₂ cell⁻¹ d⁻¹ (standard error, SE; n = 2). It was intriguing to note that the highest BR rate was measured for +Fe addition alone, which shows also the lowest (3.2%) bacterial growth efficiency (BGE) due to low BP rate (**Table 2**). The highest BGE estimation was measured for the +Fe+C treatment with 57% (**Table 2**).

TABLE 2 | Specific bacterial production, cell specific bacterial respiration, and bacterial growth efficiency at the end of the incubation (Experiment 1).

Treatment	Specific BP (fmol C cell ⁻¹ d ⁻¹)	Specific BR (fmol C cell ⁻¹ d ⁻¹)*	BGE (%)
Control	0.04 ± 0.02	0.39 ± 0.15	10 ± 0.1
+Fe	0.05 ± 0.01	1.63 ± 0.34	3.2 ± 1.5
+C	0.77 ± 0.21	0.66 ± 0.21	52 ± 15
+Fe+C	0.54 ± 0.10	0.40 ± 0.09	57 ± 10

*Bacterial respiration rates have been converted from O_2 into C units using a RQ value of 1. Samples were prefiltered on 1 μ m and kept in the dark to measure specifically the consumption of O_2 by heterotrophic bacteria. Means values \pm SE of two biological replicates are given for specific BR and BGE. Means values \pm SD of three biological replicates are given for specific BP.

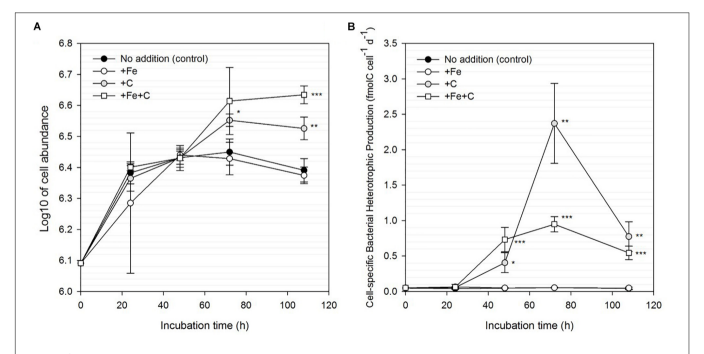


FIGURE 3 | Changes in bacterial cell abundance **(A)** and cell-specific bacterial heterotrophic production **(B)** for unamended (control) and amended treatments (+Fe, +C, and both +Fe +C) over time for Experiment 1. Treatments with an asterisk are significantly different from the control (one-way analysis of variance (ANOVA) and post hoc Tukey test; *p < 0.05, **p < 0.01, ***p < 0.001).

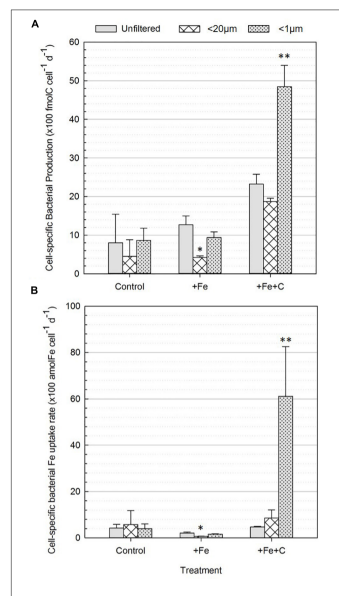


FIGURE 4 | Cell-specific heterotrophic production **(A)** and Fe uptake **(B)** by bacteria (0.2–0.8 μm fraction-size) for the three different treatments (control, +Fe, +Fe+C) and conditions (unfiltered, <20 μm , <1 μm). Values represent average number of three biological replicates after 36 h incubation (Experiment 2). Error bars represent the standard deviation of the three biological replicates. Treatments with an asterisk are significantly different from the unfiltered condition (one-way analysis of variance (ANOVA) and *post hoc* Tukey test; *p < 0.05, **p < 0.01).

Experiment 2: Responses of Bacteria When Phytoplankton Is Absent or Present

To make the section "Results" concise and easy to follow, all results presented in figures and text description correspond to the first time point for Experiment 2 (+ 36 h of incubation). Similar conclusions can be formulated from the second time point (+ 120 h of incubation) and data are accessible in the **Supplementary Materials**.

Bacterial Production in Presence and Absence of Phytoplankton

Cell-specific BP rates across all treatments and conditions are shown in Figure 4A. In the control treatment (no nutrient addition), the cell-specific BP ranged from 4 to 9 (×100 fmolC cell⁻¹ d⁻¹) and no significant difference was found between unfiltered and $<20 \mu m$ nor $<1 \mu m$ conditions, which is evidence that the presence of phytoplankton did not affect cell-specific BP. However, in the nutrient amended treatments two differences were significant: a decrease in BP in the +Fe treatment and an increase in the +Fe+C treatment, respectively. In the +Fe treatment, the cell-specific BP was highest when the whole community was present (unfiltered) and the lowest when bacteria were incubated solely with pico- and nanoplankton (<20 μm). The difference between these two conditions was significant (p = 0.04, Figure 4A). Cell-specific BP in +Fe+C treatment were higher than all other treatments, ranged from 19 to 48 $(\times 100 \text{ fmolC cell}^{-1} \text{ d}^{-1})$ and was the highest when bacteria were incubated solo. Cell-specific BP was about two times higher in <1 µm compared to unfiltered condition, and this difference was highly significant (p = 0.005, **Figure 4A**).

Significant differences were also found when comparing $<\!20~\mu m$ and $<\!1~\mu m$ conditions in +Fe and +Fe+C treatments. Indeed, the $<\!1~\mu m$ condition showed significantly higher rates compared to the $<\!20~\mu m$ condition with, respectively 9 \pm 1 versus 4 \pm 0.4 and 48 \pm 6 versus 19 \pm 1 (×100 fmolC cell $^{-1}$ d $^{-1}$) for +Fe and +Fe+C treatment. Overall, cell-specific BP was negatively affected by the presence of pico- and nanoplankton cells in the +Fe and +Fe+C treatments while no effect was found in the control (no addition) treatment.

Microbial Fe Uptake

To investigate whether heterotrophic bacteria compete for Fe with other members of the microbial community, the bacterial Fe uptake rates were determined for incubations where microplankton (<20 μm condition) or both micro- and pico- and nanoplankton (<1 μm condition) were excluded from incubation. Results were compared to the treatment where all the members of the microbial community were present (unfiltered condition). During Experiment 2, we also compared the contribution of two size-fractions to Fe uptake. **Figure 4B** shows results for bacteria on a cell-specific basis, and **Figure 5** combines data for phytoplankton (<0.8 μm) and bacteria (0.2–0.8 μm) on the volumetric basis for better comparison. Data used to create **Figure 5** can also be found in detail in **Table 3**.

Cell-Specific Fe Uptake by Bacteria

The response of bacteria to size-fractionation (condition) and nutrients amendments (treatment) were overall similar for cell-specific Fe uptake to those presented for bacterial production (**Figure 4**). While no significant differences were found for the control treatment, the uptake of Fe by bacteria was, respectively lowered and enhanced in +Fe and +Fe+C treatments. In +Fe treatment, Fe uptake by bacteria is three times lower in presence of pico-nanoplankton (<20 μ m condition) and decreased from 2.1 \pm 0.5 to 0.6 \pm 0.1 (×100 amolFe cell⁻¹ d⁻¹) compared to the unfiltered condition. This difference was significant (p = 0.006).

Across all datasets, the Fe uptake by bacteria was the highest for the +Fe+C treatment and the $<1~\mu m$ condition. This result makes precise the sense in which the availability of C together with the removal of potential competitors had the greatest effect on the uptake of Fe by bacteria. Interestingly, the concomitant addition of Fe and C did not enhance the bacterial Fe uptake for the other conditions (unfiltered and $<20~\mu m$). Values were, respectively 4.7 ± 0.3 and $8.5\pm3.5~(\times100~amolFe~cell^{-1}~d^{-1})$ for unfiltered and $<20~\mu m$ conditions while it reached up to $61\pm21~(\times100~amolFe~cell^{-1}~d^{-1})$ for the $<1\mu m$ condition. Considering all data together, the size fractionation used in the incubation (i.e., presence or absence of phytoplankton) had a greater effect than addition of growth-limiting nutrients.

Fe Uptake by Phytoplankton

Iron uptake by cells larger than 0.8 μ m is presented as the Fe uptake by phytoplankton in **Figure 5** and specific Fe uptake by pico-nanoplankton (0.8–20 μ m) is given in **Table 3**. If we consider the unfiltered condition with no nutrient addition to be the closest representation of the natural system, phytoplankton contributed to 66 \pm 6% of the total Fe uptake at the SOTS site. In terms of percentage contribution, phytoplankton Fe uptake increased to more than 80% when Fe or Fe plus C was added (82 \pm 1% and 83 \pm 1% for +Fe and +Fe+C treatment, respectively).

In the +Fe treatment, the Fe uptake by phytoplankton was slightly lower in $<\!20~\mu m$ condition compared to unfiltered condition which is explained by the removal of about 20% of

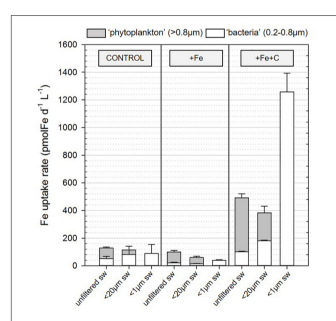


FIGURE 5 | Contribution of the two size-fraction (>0.8 μ m and 0.2–0.8 μ m) to Fe uptake in incubation for the three conditions (unfiltered, and prefiltered on <20 μ m or <1 μ m seawater) and for the three treatments [control (no addition), +Fe, +Fe+C]. Bars and errors bars represent, respectively the average and the standard deviation of biological triplicates for the Fe uptake measured following 36 h (T1, Experiment 2) of incubation in conditions described in section "Materials and Methods." sw, seawater.

the phytoplankton initial biomass. However, the concomitant addition of Fe and C clearly led to the increase of Fe uptake by phytoplankton. As the Fe uptake is higher by 22% in unfiltered conditions compared to $<20~\mu m$, which is again explained by the removal of larger cells, we consider that the microphytoplankton also benefited from the +Fe+C treatment in some ways (e.g., Fe regenerated by bacteria).

Carbon Biomass and Fe:C Ratios

Contribution to Carbon Biomass at Initial Conditions

We examined the carbon (C) biomass partitioning of the pico-and nanoplankton communities across treatments and size fractions, and in relation to the heterotrophic bacteria. First, we compare estimates of C biomass of small photosynthetic cells based on conversion factors and flow cytometry numbers with direct measurements of POC as described before. We found 1.49 \pm 0.05 μ molC L^{-1} (estimate) versus 1.51 μ molC L^{-1} (measure). Given the comparable results we are confident in using conversion factors to investigate variations in carbon biomass in our incubation bottles.

Among the pico-and nanoplankton community, picoeukaryotic cells were the most abundant in surface waters at SOTS and represented $71 \pm 0.2\%$ (n = 9) of C biomass while cyanobacteria made up $26 \pm 0.3\%$ (n = 9) and photosynthetic nanoeukaryotes $2 \pm 0.1\%$ (n = 9). After 36 h of incubation, no notable differences in these contributions were found when comparing size fractions and treatments. In the case of total carbon biomass, heterotrophic bacteria were dominant (averaging 58%) at the start of the experiment, with values ranging from 1.85 to $2.33~\mu$ mol L^{-1} .

Contribution to Fe Uptake for Bacteria and Pico-Nanoplankton

In this section, we only consider values measured in $<\!20~\mu m$ condition as we did not directly measure the carbon content of larger cells (microplankton) so that we cannot evaluate accurately their contribution. Heterotrophic bacteria represented about 52% (control), 56% (+Fe), and 59% (+Fe+C) of the total C biomass (**Table 3**). However, their contribution to the total uptake of Fe did not reflect this dominance (**Figure 5**). For the unfiltered condition, heterotrophic bacteria were only responsible for 25, 20, and 32% of the total Fe uptake in control, +Fe and +Fe+C treatments, respectively.

Fe:C Ratio

We normalized Fe uptake rate to carbon biomass and the resulting Fe:C ratio is presented in **Table 3**. Comparison of these Fe:C ratios for bacteria among the different treatment and condition indicate that heterotrophic bacteria had a higher Fe content in the +Fe+C treatment. However, this Fe:C ratio also varied from 46 to 591 μ molFe molC $^{-1}$ (+Fe+C treatment, **Table 3**) which shows that heterotrophic bacteria can assimilate a substantial amount of Fe when phytoplankton is removed from the experiment. For instance, the Fe:C ratio of heterotrophic bacteria in the <20 μ m condition was nearly twofold higher than in the unfiltered condition, and it was close to 13-fold higher in the <1 μ m condition. These high Fe:C ratios also indicate

TABLE 3 | Fe uptake rates, C biomass, and C-normalized Fe uptake rates (Fe:C ratio) of photosynthetic cells (cyanobacteria plus pico- and nanoeukaryotes) and heterotrophic bacteria.

Condition	Treatment	Fe uptake rate (pmolFe L^{-1} D^{-1})		C biomass (μmolC ⁻¹ L ⁻¹)		Fe:C ratio (μmolFe molC ⁻¹)	
		Phot. cells	H. bacteria	Phot. cells	H. bacteria	Photo. cells	H. bacteria
Unfiltered	Control	103 ± 5.4¥	52 ± 16	1.84 ± 0.34	1.29 ± 0.19	51 ± 0.2¥	41 ± 15
	+Fe	79 ± 9.3¥	21 ± 3.6	1.65 ± 0.23	1.06 ± 0.12	49 ± 10 [¥]	20 ± 4
	+Fe+C	393 ± 23 [¥]	101 ± 5.1	1.79 ± 0.16	2.20 ± 0.19	231 ± 24 [¥]	46 ± 3
$<$ 20 μ M	Control	114*	38*	1.44 ± 0.90	1.54 ± 0.90	59*	15*
	+Fe	60 ± 10	15 ± 2.1	1.86 ± 0.04	2.34 ± 0.09	32 ± 6	6 ± 1
	+Fe+C	383 ± 47	181 ± 21	1.56 ± 0.28	2.37 ± 0.67	250 ± 50	83 ± 34
$<1~\mu M$	Control	n/a	89 ± 65	n/a	2.10 ± 0.86	n/a	39 ± 20
	+Fe		40 ± 3		2.62 ± 0.16		15 ± 2
	+Fe + C		1258 ± 136		2.25 ± 0.55		591 ± 208

Values represent the average \pm standard deviation of the three independent biological replicates for each treatment and condition after 36 h of incubation. *Only one replicate available. *\frac{\pi}{2}Calculation} were done on the basis that photosynthetic cells for size fraction comprised between 0.8 and 20 \(\mu\)m represented 80% of Fe uptake (as it was measured they represented 80% of POC in unfiltered condition).

that heterotrophic bacteria have rapidly upregulated their Fe acquisition machinery relative to C to acquire more dissolved Fe or that their uptake systems were already activated.

The Fe:C ratios were also estimated for photosynthetic pico- and nanoplankton cells (including cyanobacteria). As for heterotrophic bacteria, we observed a pronounced increase of Fe:C ratio in the +Fe+C treatment, but no notable difference between unfiltered and <20 μ m condition (**Table 3**). These estimates ranged from 32 to 59 μ molFe molC⁻¹ for both control and +Fe treatments considered versus a range of 231–250 μ molFe molC⁻¹ in +Fe+C treatment.

Overall, our calculation of Fe:C ratios show that picoand nanoplankton constitute a larger fraction of biogenic Fe compared to heterotrophic bacteria in all incubations. However, the highest Fe:C ratio measured in this study was when bacteria were incubated alone (591 \pm 208, n = 3; +Fe+C treatment) and it was more than twofold higher than the maximum we calculated for pico-and nanoplankton (250 \pm 50, n = 3; +Fe+C treatment for <20 μ m condition).

DISCUSSION

In environmental science, the concept of bioavailability for one or several resources is generally associated with chemical features, in particular in the case of Fe. This micronutrient is present in multiple chemical forms and redox states (Morel and Price, 2003). However, we show here that biological interactions matter as well. To discuss the results of this study, we first comment on the concept of co-limitation. Next, we discuss the nature of the interspecific interactions that most likely explain our results. Finally, we close this section on implications for future perspectives of research.

Does Carbon Availability Offset Fe Limitation in Heterotrophic Bacteria?

One particular feature of the Southern Ocean is that both bioavailable Fe and organic carbon can be at growth-limiting

concentrations for heterotrophic bacteria in surface waters (Church et al., 2000; Obernosterer et al., 2015). At first sight, our results suggest that heterotrophic bacteria were primarily limited by organic carbon at SOTS. Iron could have a role, however, in affecting BP and bacterial metabolism when the supply of organic carbon is adequate and Fe concentrations are low. Indeed, while these results lead us to the conclusion that heterotrophic bacteria were firstly C-limited, the argument for Fe limitation of heterotrophic bacteria is not so clear. A simple comparison between in situ molar DFe:DOC ratio (2.61 μ mol mol⁻¹) and Fe:C ratios of Fe-limited cultures (e.g., 0.43 ± 0.1 and 7.52 ± 1.65 µmolFe molC⁻¹ for oceanic strains in Fourquez et al., 2014 and Tortell et al., 1996, respectively) would suggest that both nutrients may become limiting. In the present study, Fe alone had no effect on rates of BP or cell abundance. But Fe did affect these variables when added together with glucose.

Consistent with the hypothesis that bacteria may have been co-limited by Fe and C, we observed that cell-specific BP was positively correlated ($r=0.98,\ n=6,\ p=0.000373,$ Pearson correlation) with bacterial Fe uptake in the +Fe+C treatment. In contrast, there was no significant correlation in the control ($r=-0.33,\ n=6,\ p=0.58$), and in the +Fe treatment ($r=0.29,\ n=6,\ p=0.56$). Such a high correlation in studies using natural communities is uncommon. An explanation for our results is that bacterial growth became Fe-limited, but only after C-limitation was alleviated by the addition of glucose. This can be explained by the increase in the C demand induced by cellular remodeling to support growth and maintenance under Fe stress conditions (Kirchman et al., 2000; Fourquez et al., 2014).

There is growing evidence that the expression of alternative pathways is a widespread strategy for heterotrophic bacteria in low Fe environments (Fourquez et al., 2014; Beier et al., 2015; Koedooder et al., 2018; Debeljak et al., 2019). A comparative proteomics approach revealed that Fe limitation leads cells to utilize C through the glyoxylate cycle (Fourquez et al., 2014). This alternative pathway not only bypasses two important

Fe-containing enzymes in the Krebs's cycle, but also the two steps where carbon is lost as CO₂. Redirection of glucose into the Entner-Doudoroff pathway also allows Fe-limited cells to supply the Krebs's cycle with substrates while bypassing the first step of glycolysis which is ATP-consuming (Fourquez et al., 2014). Other biomass recycling processes such as amino and organic acid catabolism contribute as well to the regulation of energy production (Fourquez et al., 2014). These underlying mechanisms may explain why BP was stimulated by C addition, strongly stimulated by concomitant Fe and C additions, but not stimulated by the addition of Fe alone in the present study. It also indicates that Fe–C co-limitation for heterotrophic bacterial growth is a predominant feature in the Southern Ocean, but that it can be masked by conventional experimental approaches.

A Minimum Fe:C Quota to Support Bacterial Growth?

In our study we explicitly examined the bacterial Fe uptake together with the BP. As we observed there is a good agreement in the trends for each response variable, bacterial growth and Fe uptake rates are likely to be related. However, the link between growth and nutrient uptake is not straightforward in natural communities, and variation in maximum growth rate and minimum cell quota can greatly complicate this relationship. Here we propose a threshold value to reconcile these two variables. Based on the correlation between cellspecific Fe uptake and BP, we derived a minimum Fe:C quota for heterotrophic bacteria of 37 μmolFe molC⁻¹ that we propose as a threshold limit value to define Fe or C limitation (Supplementary Figure 2). Based on this assumption, Fe is the primary limiting element for a cellular quota that is below 37 μ molFe molC⁻¹, and C is the primary limiting element for a quota above to this value. To understand the boundaries of implication for this threshold limit, it is important to consider how heterotrophic bacteria utilize Fe and organic substrates to gain energy. However, the vastness of biogeochemical gradients both spatially and temporally - that govern the composition of microbial communities, and the plethora of metabolic strategies among taxa (Hopkinson and Barbeau, 2012; Hogle et al., 2016; Debeljak et al., 2019) require similar studies in other ocean regions in order to investigate the spectrum of heterotrophic bacterial Fe:C quotas.

Carbon Availability Increases Fe Demand: The Starting Point of Competition?

When DOC is no longer a limiting resource, the competition for Fe between autotrophic pico- and nanoplankton and heterotrophic bacteria negatively affect the latter (Fourquez et al., 2015, this study). There were also intriguing results in the outcomes of the experiment regarding phytoplankton. In the present study, the uptake of Fe by phytoplankton was similar for the control and the +Fe treatment but increased by nearly five times in the +Fe+C treatment. We have two explanations for this intriguing finding.

Phytoplankton Had a Higher Fe Uptake Rate in +Fe +C Treatment Because Fe Regenerated by Bacteria Became Available

Iron availability influences the growth and abundance of autoand heterotrophic microorganisms, and heterotrophic bacteria can modify its speciation by the synthesis of organic ligands (Rue and Bruland, 1997; Gerringa et al., 2008). In this context, heterotrophic bacteria could act either as competitors with phytoplankton (Kirchman, 1994; Thingstad, 2000), or on the contrary, facilitate their assimilation of Fe in maintaining Fe solubility within the ecological niche they share (Amin et al., 2009; Hopkinson and Morel, 2009).

The high rates of BP in +Fe+C treatment could in part be due to remineralization of Fe during the incubation. The regenerated Fe may become available for phytoplankton. Since the DFe concentration was measured in the incubation bottles at the beginning, after + 36h and at the end of the incubation (120 h, **Supplementary Figure 3**), we were able to directly compare these values with total Fe uptake by microorganisms. The DFe concentrations decreased over time during the incubations (**Supplementary Figure 3**). We used a simple approach to (1) verify that remineralization occurred during the incubation and (2) to provide an estimate of the Fe regeneration rate. The regeneration rate of Fe was estimated by subtracting the amount of Fe consumed by the entire microbial community (phytoplankton and bacteria) from the initial Fe concentration as follows:

$$DFe_{regenerated} = DFe_{mesured} - DFe_{expected}$$

With DFe_{expected} = DFe_{initial} - [Total Fe_{uptake}
$$\times t$$
])

Where DFe initial is the concentration of DFe at the start of the experiment, Total Fe uptake is the amount of Fe consumed by phytoplankton and bacteria during the incubation, and DFe measured is the DFe concentration at the end of the incubation.

If we consider the amount of "missing" DFe as the regenerated Fe, we obtain rates of Fe regeneration of 0.48 to 0.92 nmolFe L^{-1} d⁻¹ (respectively unfiltered and <1 μ m condition, **Supplementary Table 1**).

Iron regeneration within the microbial loop (also termed the "ferrous wheel"; Kirchman, 1996) represents a key term in the Fe budget (Strzepek et al., 2005; Boyd and Ellwood, 2010). For instance, in the SAZ (FeCycle voyage, see Strzepek et al., 2005), it was found that between 30 and 100% of the microbial Fe demand could be met by Fe regeneration mediated by grazers (Boyd et al., 2005). In the naturally Fe fertilized waters off Kerguelen Island, Fe regeneration accounted for roughly 50% of the Fe demand (Sarthou et al., 2008). Heterotrophic bacteria and viruses contribute as much as grazers to Fe recycling (Poorvin et al., 2004; Obernosterer et al., 2008). Many of the metabolites originating from microorganisms can possess Fe-binding properties that can exert strong control on Fe speciation (Poorvin et al., 2004; Dalbec and Twining, 2009; Boyd et al., 2010). Unlike larger cells of phytoplankton (e.g., diatoms), pico- and nanoplankton are equally adept to heterotrophic bacteria at accessing either new or regenerated Fe (Boyd et al., 2012). Ultimately, niche differentiation of bacteria

TABLE 4 | Total intracellular Fe uptake rates in this study compared to published studies of natural and Fe-fertilized surface waters in the Southern Ocean.

Experiment location (acronym)	Fe uptake rate (pmolFe L^{-1} d^{-1})	References
SAZ waters – SOTS	96–156	This study – unfiltered control
	847–1258*	This study – $<1 \mu m + Fe + C$
Kerguelen Plateau (KEOPS)	4.4–6.2	Sarthou et al. (2005)
Kerguelen Plateau (KEOPS2)	19–39.8	Fourquez et al. (2015)
South of Australia (SOIREE)	3.07-11.9	Bowie et al. (2001)
Southeast of New Zealand (FeCycle)	26.2–101	Strzepek et al. (2005)

^{*}Heterotrophic bacteria only.

and phytoplankton related to Fe-speciation might act as a selection process (Hopkinson and Barbeau, 2012; Hogle et al., 2016; Debeljak et al., 2019).

Over the course of a phytoplankton bloom there is a transition from the utilization of new Fe (i.e., winter reserve Fe stocks) to regenerated Fe (Boyd et al., 2012) which maintains primary productivity. During this transition, rapidly growing heterotrophic bacteria may quickly shift to Fe limitation if phytoplankton-derived organic carbon is available, resulting in their enhanced ability to compete for Fe. As the bloom status moves toward senescence and cells exude DOC, competition between pico- and nanoplankton and heterotrophic bacteria may result in different amounts of Fe regenerated. Significantly, In the present study we calculated a Fe regeneration rate nearly twofold larger for the $<1~\mu m$ (0.92 nmolFe $L^{-1}~d^{-1}$) versus $<20~\mu m$ (0.53 nmolFe L^{-1} d⁻¹) or in unfiltered seawater (0.48 nmolFe L⁻¹ d⁻¹). It is reported that organic Fe complexes are available to few phytoplankton species (Kranzler et al., 2011; Shaked and Lis, 2012; Lis et al., 2015). This could suggest that the bloom duration is primarily set by DOC availability and competition for DFe between heterotrophic bacteria and phytoplankton.

Synechococcus Like It Organic

Many studies make operational distinctions based on sizefractionated samples. In the study of Strzepek et al., 2005 (FeCycle), flow cytometric analyses revealed that both picophytoplankton and eukaryotic phytoplankton were $> 1 \mu m$, and heterotrophic bacteria were submicron in size. We used the same size cutoffs of 1 µm to separate phytoplankton and heterotrophic bacteria in this study. It remains, however, difficult to draw clear distinctions between the size-fractionation and assigning microorganisms to eukaryotes or prokaryotes. At SOTS, most pico- and nanoplankton (70%) were cyanobacteria (most likely Synechococcus). Synechococcus is one of the most prominent genera of picoplanktonic marine cyanobacteria (Buitenhuis et al., 2012) that have particularly high Fe demands relative to heterotrophic bacteria and eukaryotic phytoplankton (Raven, 1990; Morrissey and Bowler, 2012; Lis et al., 2015). There is growing evidence of the ability of the genera of Synechococcus to assimilate organic nutrients (Yelton et al., 2016), and more broadly there are reports that suggest some of the photosynthetic picoeukaryotes are mixotrophs (Farnelid et al., 2016). Cyanobacteria may be responsible for an important part of the total Fe taken up by the phytoplankton fraction (>0.8 µm), and that due to their ability to take up organic compounds for their metabolism they may also have

benefited from the +Fe+C treatment, as did their heterotrophic counterparts. In our experiment, cyanobacteria, but none of the pico- and nanoeukaryotes had significantly increased in cell abundance in the +Fe+C treatment (**Supplementary Figure 4**). This incubation was performed under very low light intensities which suggests a complementary mechanism such as mixotrophy. This raises the question of whether a mixotrophic capacity can become an advantageous for these microorganisms, and if they could become also competitors for C availability if this is proven.

The Fate and Duration of Phytoplankton Blooms in the Southern Ocean Driven by Interspecific Relationships

Interactions between autotrophic and heterotrophic microbes could affect the dynamics of nutrient-limited phytoplankton blooms. This hypothesis originates from an investigation of what was initially perceived as an isolated event in the vicinity of the Kerguelen plateau (Fourquez et al., 2015). As we reached similar conclusions in the present study, these joint findings raise the issue of whether such interactions are widespread across the Southern Ocean?

One stand-out result of our study is the amount of Fe taken up by heterotrophic bacteria in the absence of competition. Our reported bacterial Fe uptake rates are well beyond the range of those previously reported for Southern Ocean microbes. For example, in comparison to the Fe uptake by the entire microbial community, the rate of heterotrophic bacteria alone is by 12-fold higher than that measured during FeCycle (HNLC waters southeast of New Zealand) and more than 100-fold higher than that measured above the Kerguelen plateau during summer (KEOPS2; **Table 4**). The observation of the present study underlines the potential of heterotrophic bacteria to control the decline of the bloom in the absence of competition with pico-and nanoplankton.

It is only recently that the potential influence of the composition of phytoplankton community and its interactions with heterotrophic microbes has been taken into consideration (Lima-Mendez et al., 2015; Bunse et al., 2016; Farnelid et al., 2016; Zhou et al., 2018; Liu et al., 2019). For instance, Liu et al. (2019) showed a pronounced association between assemblages of diatoms and heterotrophic microbes at the onset of spring phytoplankton blooms occurring in the region off Kerguelen Island. The quality and quantity of DOC derived from phytoplankton exudates (Landa et al., 2015) and resource

competition for Fe (Fourquez et al., 2015) are the two explanations put forward to explain how diatom assemblages shape the habitat type for their heterotrophic counterparts (Liu et al., 2019).

Uncertainties remain on the effects of climate change on the composition of phytoplankton assemblage; but there is compelling evidence that all regions of the Southern Ocean will encounter changes in phytoplankton community composition (Hays et al., 2005; Deppeler and Davidson, 2017). Models project that waters of the Southern Ocean will become warmer, and that rising temperatures will cause rates of grazing to increase more rapidly than rates of phytoplankton growth (Sarmento et al., 2010; Evans et al., 2011; Caron and Hutchins, 2012; Behrenfeld, 2014; Cael and Follows, 2016). Increasing temperature is also expected to increase bacterial respiration rates (Vázquez-Domínguez et al., 2007). Thus, phytoplankton standing stocks are likely to decline and the proportion of primary production respired in near-surface waters by heterotrophic bacteria will increase (Deppeler and Davidson, 2017; Cavan and Boyd, 2018; Cavan et al., 2019). The study of Cavan and Boyd (2018), which have predicted an increase in POC-normalized respiration, estimates that the biological pump efficiency (POC export scaled to primary production) would decrease by $17 \pm 7\%$ (SE) by 2100 for the subantarctic site SOTS. Such reports of increased rates in bacterial respiration are enzymatic reactions as the temperature increases, but are also supported by the enhanced release DOC-derived from phytoplankton at higher temperatures to support heterotrophic bacteria (Hutchins et al., 2019).

For microbial ecologists, the existence of interactions between primary producers and bacteria that shape the activity and the diversity of both partners is well recognized (Amin et al., 2015), but the mechanisms of such interactions remain mostly unknown. The term "interactive co-limitation" was first proposed by Bertrand et al. (2015) to describe scenarios in which at least two limiting "nutrients cycle are affected by one another through interactions among different microbial functional groups" (Bertrand et al., 2015). The findings of our study appear to be an example of interactive colimitation. Moreover, they expand on the emerging recognition that interaction between microorganisms is an ecological trait to be considered in the study of Fe and C biogeochemistry. As Fe sources - including the inputs, amounts and the nature of Fe - are often considered to be the primary drivers of Southern Ocean productivity, our study teaches us that Fe bioavailability for microorganisms is not a simple matter of chemistry.

DATA AVAILABILITY STATEMENT

All datasets generated for this study are included in the article/Supplementary Material.

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AUTHOR CONTRIBUTIONS

MF conceived and conducted the study, and wrote the manuscript. MB contributed to the design of the study. MB, ME, and PB contributed to data collection at sea. SD contributed to the sample analysis. IO and TT contributed to data analysis and interpretation. PB acquired funding. All co-authors reviewed and edited this or previous versions of the manuscript and gave final approval for publication.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fmars. 2019.00776/full#supplementary-material

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Conflict of Interest: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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