Modelled estimates of spatial variability of iron stress in the Atlantic sector of the Southern Ocean

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Abstract. The Atlantic sector of the Southern Ocean is characterized by markedly different frontal zones with specific seasonal and sub-seasonal dynamics. Demonstrated here is the effect of iron on the potential maximum productivity rates of the phytoplankton community. A series of iron addition productivity versus irradiance (PE) experiments utilizing a unique experimental design that allowed for 24 h incubations were performed within the austral summer of 2015/16 to determine the photosynthetic parameters αB, PmaxB, and Ek. Mean values for each photosynthetic parameter under iron-replete conditions were 1.46 ± 0.55 (µg (µg Chl a)−1 h−1 (µM photons m−2 s−1)−1) for αB, 72.55 ± 27.97 (µg (µg Chl a)−1 h−1) for PmaxB and 50.84 ± 11.89 (µM photons m−2 s−1) for Ek, whereas mean values under the control conditions were 1.25 ± 0.92 (µg (µg Chl a)−1 h−1 (µM photons m−2 s−1)−1) for αB, 62.44 ± 36.96 (µg (µg Chl a)−1 h−1) for PmaxB and 55.81 ± 19.60 (µM photons m−2 s−1) for Ek. There were no clear spatial patterns in either the absolute values or the absolute differences between the treatments at the experimental locations. When these parameters are integrated into a standard depth-integrated primary production model across a latitudinal transect, the effect of iron addition shows higher levels of primary production south of 50° S, with very little difference observed in the subantarctic and polar frontal zone. These results emphasize the need for better parameterization of photosynthetic parameters in biogeochemical models around sensitivities in their response to iron supply. Future biogeochemical models will need to consider the combined and individual effects of iron and light to better resolve the natural background in primary production and predict its response under a changing climate.

1 Introduction

Phytoplankton primary production (PP) in the Southern Ocean is a key contributor to global atmospheric CO2 drawdown, responsible for 30–40% of global anthropogenic carbon uptake (Khatiwala et al., 2009; Mikaloff Fletcher et al., 2006; Schlitzer, 2002). High nutrient availability fuels this phytoplankton production, but growth is ultimately constrained by the lack of availability of the micronutrient iron (Fe) (de Baar et al., 1990; Martin et al., 1990). This leads to high levels of macronutrients that remain unutilized by phytoplankton growth in what is known as a high-nutrient, low-chlorophyll (HNLC) conditions. Maximum primary productivity rates of the Southern Ocean are also limited by light availability due to low incident solar angles, persistent cloud cover and deep mixed layers that curtail production and subsequently affect the efficiency of the biological carbon pump. Under future climate change scenarios, altered upwelling and mixed layer stratification (Boyd et al., 2001; Boyd and Doney, 2002), changes in sea ice cover (Close and Goosse, 2013; de Lavergne et al., 2014; Montes-Hugo et al., 2008; Zhang, 2007) and food-web dynamics (Dubischar and Bathmann, 1997; Moore et al., 2013; Pakhomov and Frone- man, 2004; Smetacek et al., 2004) will alter both the nutrient and light supply, strongly impacting primary production rates. As such, it is important that we understand the sensitivity of phytoplankton production to light and micronutrient...
availability so that we may improve our predictive capability of the response of the Southern Ocean carbon pump to a changing climate.

Iron plays a critical role in modulating PP due to the high requirements of the photosynthetic apparatus, photosystems I and II (Quigg et al., 2003; Raven, 1990; Strzepek and Harrison, 2004; Twining and Baines, 2013). Light availability can further increase the demand for iron, as low irradiance levels increase requirements associated with the synthesis of additional photosynthetic units to increase potential light absorption (Maldonado et al., 1999; Raven, 1990; Strzepek et al., 2012; Sunda and Huntsman, 1997). Iron is also required to activate both nitrate and nitrite reductase (de Baar et al., 2005), which facilitate the assimilation of nitrate and nitrite and their subsequent intracellular reduction to ammonium. In HNLC regions, such as the Southern Ocean, nitrate uptake rates ($\rho_{\text{NO}_3}$) have also frequently been reported as becoming iron-limited (Cochlan, 2008; Lucas et al., 2007; Moore et al., 2013; Price et al., 1994). However, it has also been demonstrated that iron limitation rather than inhibiting nitrate reductase activity results in a bottleneck further downstream due to a reduction in photosynthetically derived reductant (Milligan and Harrison, 2000). This would lead to an excretion of excess nitrate back into the water column that would further contribute to HNLC conditions such as those present in the Southern Ocean.

Estimating PP in the oceans towards an improved understanding of the effects of iron and light limitation requires an understanding of the relationship between photosynthesis ($P$) and irradiance ($E$) (Behrenfeld and Falkowski, 1997b; Dower and Lucas, 1993; Platt et al., 2007). $PE$ responses are derived from an equation by Platt et al. (1980), where the responses are parameterized as a function of irradiance. The parameters derived include $P_{\text{max}}^B$, the biomass-specific rate of photosynthesis at saturating irradiances; $a^B$, the irradiance-limited biomass-specific initial slope; and $E_k$, the irradiance at which saturation is initiated. The response of these parameters can be not only a function of temperature (Behrenfeld and Falkowski, 1997b) but also as a change in the quantum efficiency of photosynthesis, usually as the result of changes in iron availability. In previous iron fertilization experiments a doubling of $a^B$ has been reported (Hiscock et al., 2008), yet this response is not consistent across Southern Ocean waters (Feng et al., 2010; Hopkinson et al., 2007; Moore et al., 2007; Smith Jr. and Donaldson, 2015). Given their relative importance within PP models (Behrenfeld and Falkowski, 1997a, b; Sathyendranath and Platt, 2007), a greater understanding of the drivers of the variability within these photosynthetic parameters is therefore required, particularly if we are to accurately quantify and constrain PP in the Southern Ocean to examine seasonal and interannual variability and trends.

The Atlantic sector of the Southern Ocean is composed of a series of circumpolar fronts that are characterized by large geostrophic velocities (Nowlin and Klinck, 1986; Orsi et al., 1995). The fronts constrain water masses with distinct physical and chemical properties that define different oceanographic zones. These spatial zones display not only zonal variability with the fronts but also display important seasonal contrasts (Thomalla et al., 2011), with differing bloom initiation dates and temporal extent of bloom duration. Whilst the bloom initiation dates can in part be explained by day length and sea ice cover further polewards, the differences in the extent and duration of blooms between the zones requires an alternative and more nuanced explanation. One theory that has been postulated is that the supply mechanisms of iron to the mixed layer following the spring bloom vary between zones (Thomalla et al., 2011). Weak diapycnal inputs and a heavy reliance on iron recycling was suggested by Tagliabue et al. (2014) to match approximate phytoplankton utilization within the pelagic zones. An alternative theory that postulates the importance of summer storms may also be pivotal in understanding the seasonal dynamics of phytoplankton primary productivity (Nicholson et al., 2016; Swart et al., 2015; Thomalla et al., 2015), with respect to the sustained bloom observed in the sub-Antarctic Zone (SAZ). Here, summer storms are said to periodically deepen the mixed layer to below the ferricline followed by rapid shoaling during quiescent periods that balances the supply of light and iron in the upper oceans favouring phytoplankton growth that culminates in a sustained summer bloom (Swart et al., 2015). Regardless of the mechanisms at play, an understanding of when and where iron concentrations and supply mechanisms limits potential phytoplankton growth and productivity is needed to better understand the drivers that determine the characteristics of the Southern Ocean seasonal cycle.

To this end, a research cruise was conducted in the austral summer of 2015/16 as part of the third multidisciplinary Southern Ocean Seasonal Cycle Experiment (SOSCEX III), which aimed to identify and understand the physical and chemical controls on the seasonal cycle of the biological carbon pump. As part of this study, shipboard nutrient addition $PE$ experiments were performed to determine the extent of iron limitation upon phytoplankton primary production.

2 Materials and methods

2.1 Oceanographic sampling

The samples and data presented here were obtained during the 55th South African National Antarctic Expedition (3 December 2015 to 11 February 2016) on board the S.A. Agulhas II to the Atlantic sector of the Southern Ocean as part of SOSCEX III (Swart et al., 2012). During the cruise, six nutrient addition $PE$ long-term experiments were performed within the Atlantic sector of the Southern Ocean (Fig. 1) to determine the extent to which relief from iron limitation could alter the maximal primary productivity rates of the phytoplankton community. Uncontaminated whole seawater was collected from 30 to 50 m depth using Teflon-lined,
external closure 12 L Go-Flo samplers deployed on a trace metal clean CTD rosette system.

### 2.2 PE experimental setup

Phytoplankton productivity was measured by the incorporation of $^{13}$C stable isotopes in response to an increasing light gradient. Inside a trace metal clean laboratory class-100 container, bulk trace metal clean seawater was decanted unscreened into an acid-washed 50 L LDPE carboy (Thermo Scientific) to ensure homogenization; this was then redistributed into acid-cleaned 1.0 L polycarbonate bottles (Nalgene). All experimental conditions were conducted and carried out following trace metal clean standards and conditions. Sample manipulations were conducted under a laminar flow hood. All bottles were inoculated with $^{13}$C (10 µM NaH$_2$C$^{13}$O$_3$ 100 mL$^{-1}$) spikes to achieve an enrichment of $\sim$5%; 11 bottles received the addition of FeCl$_3$ (+2.0 nM, “Fe”), whereas 11 bottles received the $^{13}$C spikes alone (“Control”). The bottles were incubated in screened (LEE Filters) LDPE boxes within light- and temperature-controlled incubators. Experimental temperature was set to mimic the in situ sample collection temperature. Irradiances were measured within the screened boxes using a handheld 4$\pi$ PAR sensor (Biospherical Instruments) and ranged from 0–400 µM photons m$^{-2}$s$^{-1}$. Bottles tops were covered with parafilm and double-bagged with clear polyethylene bags to minimize contamination risks during the incubation. Due to physical constraints, the experiments were not conducted as triplicates, and as such evaluation of the precision/error within experiments is not possible.

Experiments were incubated for 24 h, after which the samples were vacuum filtered through a pre-combusted (400 °C for 24 h) GF/F filter. Samples were acid-fumed with concentrated HCl for 24 h to remove inorganic carbon before being dried in an oven at 40 °C for 24 h. The isotopic composition of all samples were determined by mass spectrometry on a Flash EA 1112 series elemental analyser (Thermo Finnigan). Carbon uptake rates (µM C h$^{-1}$) were calculated from the equation of Dugdale and Wilkerson (1986), utilizing in situ determinations of dissolved inorganic carbon (DIC). The uptake rates normalized to the chlorophyll a (Chl) concentration, were used to calculate the maximal light-saturated Chl-specific photosynthetic fixation rates ($P_B^{max}$), the light-limited slope ($\alpha^B$) and the photoacclimation parameter ($E_k$). The curves and parameters were generated using a non-linear least squares fit to the equation of Platt et al. (1980).
2.3 Chlorophyll a and nutrient analysis

Samples for Chl analysis, 250 mL, were filtered onto GF/F filters and then extracted into 90% acetone for 24 h in the dark at −20°C, followed by analysis with a fluorometer (TD70; Turner Designs) (Welschmeyer, 1994). Macronutrient samples were drawn into 50 mL diluivials and stored at −20°C until analysis on land. Nitrate + nitrite and silicate were measured using a Lachat flow injection analyser (Egan, 2008; Wolters, 2002), whilst nitrite and phosphate were determined manually by colorimetric method as specified by Grasshoff et al. (1983). Dissolved iron samples (DFe) were carefully collected in acid-washed 125 mL LDPE bottles, acidified with 30% HCl Suprapur to pH ∼ 1.7 (using 2 mL L⁻¹ criteria) and stored at room temperature until analysis on land. pigment (DFe) were determined analytically by Platt et al. (1980). Rates were calculated for daily primary productivity was solved PP₀ = Pmax × (1 − e⁻¹/(Pmax × E0)). (1) PP₀ (mg C m⁻² d⁻¹) is the primary production at the surface, Pmax the maximal light-saturated photosynthetic fixation rate, α the light-limited slope and Em₀ is daily PAR at the surface, calculated by assuming maximum PAR at midday, zero PAR at sunrise and sunset, a constant gradient of light between time steps and extrapolating the measured PAR (from an above-water Biospherical 4π PAR sensor at the time of the station into an isosceles triangle; see also Thomalla et al., 2015).

2.4 Phytoplankton photosynthetic physiology

Variable Chl fluorescence was measured using a Chelsea Scientific Instruments FastOcean fast repetition rate fluorometer (FRRf) integrated with a FastAct laboratory system. Samples were acclimated in dark bottles at in situ temperatures, and FRRf measurements were blank-corrected using carefully prepared 0.2 µm filtrates for all samples (Cullen and Davis, 2003). Protocols for FRRf measurements consisted of the following: 100 × 2 µs saturation flashlets with a 2 µs interval, followed by 25 × 1 µs relaxation flashlets with an interval of 84 µs with a sequence interval of 100 ms. Sequences were repeated 32 times, resulting in an acquisition length of 3.2 s. The power of the excitation LED (4.450) was adjusted between samples to saturate the observed fluorescence transients within a given range of R <> R₂. R <> R₂, the probability of a reaction centre being closed during the first flashlet, is optimized between 0.042 and 0.064 per the manufacturer specifications. By adopting this approach, it ensures the best signal-to-noise ratio in the recovered parameters whilst accommodating significant variations in the photophysiology of the phytoplankton community without having to adjust the protocol. Data from the FRRf were analysed to derive fluorescence parameters as defined in Baker et al. (2001) and Roháček (2002) by fitting transients to the model of Kolber et al. (1998).

2.5 Pigment analysis and CHEMTAX

Pigment samples were collected by filtering 0.5–2.0 L of water onto GF/F filters. Filters were frozen and stored at −80°C until analysis in Villefranche, France, on an Agilent Technologies HPLC 1200. Filters were extracted in 100% methanol, disrupted by sonication, clarified by filtration and analysed by HPLC following the methods of Ras et al. (2008). Limits of detection were on the order of 0.1 ng L⁻¹. Pigment composition data were standardized through root square transformation before cluster analysis utilizing multi-dimensional scaling where similar samples appear together and dissimilar samples do not. Samples were grouped and analysed in CHEMTAX (Mackey et al., 1996) using the pigment ratios from Gibberd et al. (2013). Multiple iterations of pigment ratios were used to reduce uncertainty in the taxonomic abundance as described in Gibberd et al. (2013), with the solution that had the smallest residual used for the estimated taxonomic abundance.

2.6 Particle size analysis

The size distribution of the particle population was measured by running 40 mL of water sample through a 100 µm aperture on a Beckman Coulter multizer (20 runs at 2.0 mL per run), binning the size counts into 400 bins between 2 and 60 µm. Data were subsequently analysed utilizing custom MATLAB scripts to calculate the effective diameter of particles within the sample following Hansen and Travis (1974).

2.7 Depth-integrated production

Water column primary production rates were calculated according to Platt et al. (1980) and Platt and Sathyendranath (1993) as in Thomalla et al. (2015), where

\[ PP_0 = P_{\text{max}} \times (1 - e^{-\frac{\alpha \times E_{\text{m}_0}}{P_{\text{max}}}}) \]  

(1)

\[ PP_0 \text{ (mg C m}^{-2} \text{ d}^{-1}) \text{ is the primary production at the surface, } P_{\text{max}} \text{ the maximal light-saturated photosynthetic fixation rate, } \alpha \text{ the light-limited slope and } E_{\text{m}_0} \text{ is daily PAR at the surface, calculated by assuming maximum PAR at midday, zero PAR at sunrise and sunset, a constant gradient of light between time steps and extrapolating the measured PAR (from an above-water Biospherical 4} \pi \text{ PAR sensor at the time of the station into an isosceles triangle; see also Thomalla et al., 2015).} \]

\[ E_{\text{m}_0} = \frac{E_{\text{m}_0}}{E_k} \]  

(2)

The results were generalized by calculating \( E_{\text{m}_0}(2) \), the dimensionless daily surface irradiance, while primary productivity over the entire water column \( PP_{\text{wc}} \) (mg C m⁻² d⁻¹) was calculated with the following Eq. (3). The dimensionless function \( f(E_{\text{m}_0}) \) for daily primary productivity was solved analytically by Platt et al. (1980). Rates were calculated for both the iron addition and control treatments, allowing the difference between the integrated rates to be solved.

\[ PP_{\text{wc}} = PP_0 \times \frac{f(E_{\text{m}_0})}{k_d} \]  

(3)
Table 1. Locations for PE experiments conducted during the cruise along with details for the initial chemical, physiological and physical setup conditions.

<table>
<thead>
<tr>
<th>Experiment</th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
<th>5</th>
<th>6</th>
</tr>
</thead>
<tbody>
<tr>
<td>Initiation date</td>
<td>08/12/2015</td>
<td>05/01/2016</td>
<td>07/01/2016</td>
<td>08/01/2016</td>
<td>09/01/2016</td>
<td>26/01/2016</td>
</tr>
<tr>
<td>Latitude (° S)</td>
<td>-42.69</td>
<td>-42.69</td>
<td>-45.99</td>
<td>-50.45</td>
<td>-55.70</td>
<td>-70.44</td>
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<tr>
<td>Longitude (° E/W)</td>
<td>08.74</td>
<td>08.74</td>
<td>05.93</td>
<td>01.04</td>
<td>-00.00</td>
<td>-07.82</td>
</tr>
<tr>
<td>Collection depth (m)</td>
<td>30</td>
<td>35</td>
<td>35</td>
<td>35</td>
<td>35</td>
<td>35</td>
</tr>
<tr>
<td>Sunrise–sunset</td>
<td>03:30–18:30</td>
<td>04:00–19:00</td>
<td>04:00–19:00</td>
<td>04:00–19:00</td>
<td>04:00–19:00</td>
<td>00:00–00:00</td>
</tr>
<tr>
<td>Chl (µg L$^{-1}$)</td>
<td>0.97</td>
<td>0.84</td>
<td>0.89</td>
<td>2.30</td>
<td>1.15</td>
<td>1.49</td>
</tr>
<tr>
<td>Nitrates (µM)</td>
<td>7.21</td>
<td>10.20</td>
<td>15.83</td>
<td>21.07</td>
<td>17.02</td>
<td>23.81</td>
</tr>
<tr>
<td>Silicates (µM)</td>
<td>0.86</td>
<td>0.72</td>
<td>0.09</td>
<td>3.76</td>
<td>30.83</td>
<td>48.81</td>
</tr>
<tr>
<td>Phosphates (µM)</td>
<td>0.88</td>
<td>0.76</td>
<td>0.95</td>
<td>1.28</td>
<td>1.15</td>
<td>0.94</td>
</tr>
<tr>
<td>DFe (nM)</td>
<td>0.16</td>
<td>0.17</td>
<td>0.07</td>
<td>0.03</td>
<td>0.05</td>
<td>0.10</td>
</tr>
<tr>
<td>$F_{v}/F_{m}$</td>
<td>0.19</td>
<td>0.30</td>
<td>0.35</td>
<td>0.30</td>
<td>0.35</td>
<td>0.37</td>
</tr>
<tr>
<td>$\sigma_{PSII}$ (nm$^{-2}$)</td>
<td>14.79</td>
<td>6.45</td>
<td>5.50</td>
<td>5.59</td>
<td>5.37</td>
<td>3.89</td>
</tr>
<tr>
<td>MLD (m)</td>
<td>33.77</td>
<td>56.96</td>
<td>108.42</td>
<td>70.11</td>
<td>42.89</td>
<td>40.80</td>
</tr>
<tr>
<td>Salinity</td>
<td>33.87</td>
<td>33.70</td>
<td>33.88</td>
<td>33.80</td>
<td>33.73</td>
<td>33.72</td>
</tr>
<tr>
<td>Temp. (°C)</td>
<td>10.80</td>
<td>10.44</td>
<td>6.72</td>
<td>3.17</td>
<td>-1.42</td>
<td>-1.51</td>
</tr>
<tr>
<td>Average daytime PAR (µM photons m$^{-2}$ s$^{-1}$)</td>
<td>1055.31</td>
<td>787.35</td>
<td>289.18</td>
<td>524.41</td>
<td>769.87</td>
<td>673.62</td>
</tr>
<tr>
<td>Euphotic depth (m)</td>
<td>72.79</td>
<td>75.10</td>
<td>52.95</td>
<td>47.92</td>
<td>69.13</td>
<td>78.07</td>
</tr>
</tbody>
</table>

*24 h day length; b see Sect. 2.7 for details.

$K_d$ was initially calculated as the slope of the natural log of in situ PAR with depth from CTD profiles. When in situ PAR with depth was not available, $K_d$ was also calculated from in situ surface Chl concentrations with the following Eq. (4) (Morel, 1988; Morel et al., 2007). Co-located calculations utilizing in situ PAR versus chlorophyll-derived $K_d$ demonstrated on average a 40% higher $K_d$ when calculated with chlorophyll.

$$K_d = 0.0166 + 0.0773 \times [\text{Chl}]^{0.6715} \quad (4)$$

2.8 Ancillary physical data

Temperature and salinity profiles were obtained from a Sea-Bird CTD mounted on the rosette system. The mixed layer depth (MLD) was calculated following de Boyer Montégut et al. (2004), which identifies the MLD as the depth where the temperature differs from the temperature at 10 m by more than 0.2 °C ($\Delta T_{10m} = 0.2 °C$). The position of the fronts was determined using sea surface height (SSH) data from maps of absolute dynamic topography (MADT) according to Swart et al. (2010).

3 Results

3.1 Oceanographic context

The experimental setup locations covered a wide range of pelagic zones from the SAZ to the marginal ice zone (MIZ), each with different physical, chemical and biological properties (see Table 1). Chl concentrations between experiment initiation locations varied between 0.84 and 2.30 µg L$^{-1}$, peaking just south of the polar front at ∼ 50° S. Initial temperatures displayed a characteristic decrease from 10.80 °C at the most northerly location to −1.51 °C at the MIZ, whereas there were no distinct differences in salinity ranging from 33.70 to 33.88. Macronutrient concentrations all increased polewards, with peaks of 28.15, 1.34 and 48.81 µM for nitrate, phosphate and silicate respectively. Dissolved iron concentrations decreased polewards from a maximum of 0.17 nM in the SAZ to minimum values of 0.03 and 0.05 nM at 50 and 55° S respectively, before increasing again in the MIZ to 0.10 nM.

Phytoplankton photophysiology, $F_{v}/F_{m}$, increased polewards from a minimum of 0.19 to a maximum of 0.37, whereas $\sigma_{PSII}$, the effective absorption cross section of PSII, decreased polewards from 14.79 to 3.89 nm$^{-2}$. The effective diameter of the phytoplankton population, a relative measure of size, increased polewards from a minimum of 4.29 ± 0.35 µm in the SAZ to a maximum of 8.59 ± 0.68 µm in the MIZ. Estimated taxonomic abundance through HPLC analysis and CHEMTAX determined that the dominant groups at all stations were either diatoms, haptophytes or a mix of the two. Haptophytes were the dominant group (>68 % of total Chl) in the SAZ during experiments 1 and 2, with diatoms becoming dominant (>70 % of total Chl) from experiment 4 onwards.

MLDs were highly variable and ranged from ∼34 m at experiment 1 to ∼108 m at experiment 3. The MLD was typically deeper than the experimental setup depth (average difference of ∼15 m) at all experiments except for experiment...
5, where the collection depth was 7 m below the MLD. The CTD density profile at experiment 5 was indicative of two mixed layers present, with the experiment performed above the deeper of the mixed layers (≈56 m). Experiments 1 and 2 that were set up in the same location in the SAZ but 28 days apart had markedly different setup conditions: a 41 % increase in the nitrate concentration from 7.21 to 10.20 µM, a 2-fold increase in $F_v/F_m$ from 0.19 to 0.35 with a concurrent 56 % decrease in $\sigma_{PSII}$ from 14.79 to 6.45 nm$^{-2}$ and a deepening of the MLD from ≈34 to ≈57 m.

The light environment within the water column at each location was determined by calculating the percentage light depth as a function of the vertical attenuation coefficient of irradiance ($K_d$). The percentage light depths of the experiments ranged between 3.46 and 14.78 %. The 1 % light depth, which typically coincides with the compensation light depth i.e. the depth where rates of production equate to rates of respiration, is consistently below the MLD, except for experiment 4, where it was 22 m above the mixed layer.

### 3.2 PE parameters

$PE$ curves for carbon uptake ($\rho C$) (Figs. 2 and S1 in the Supplement), summarized in Table 2, display consistent results with greater values of $\alpha^B$ and $P^B_{\text{max}}$ with the addition of iron compared to the control treatments (Figs. S2–S3). The $PE$ curves for the control treatments did not display any significant outliers ($r^2 = >95\%$), we can assume that contamination levels were minimal, as no measurements of DFe in the sample bottles were collected. The values derived here fall within the range previously reported for iron addition experiments in the Southern Ocean (Hiscock et al., 2008; Hopkinson et al., 2007; Moore et al., 2007; Smith Jr. and Donaldson, 2015). Maximum values of $\alpha^B$ (mg (mg Chl $a$)$^{-1}$ h$^{-1}$ (µmol photons m$^{-2}$ s$^{-1}$)$^{-1}$) for $\rho C$ were $2.23 \times 10^{-3}$ from experiment 2 Fe treatment and $2.43 \times 10^{-3}$ from experiment 1 control treatment, with minimum values of $0.13 \times 10^{-3}$ from experiment 5 control treatment and $0.56 \times 10^{-3}$ from experiment 6 Fe treatment. $P^B_{\text{max}}$ (mg (mg Chl $a$)$^{-1}$ h$^{-1}$ (µmol photons m$^{-2}$ s$^{-1}$)$^{-1}$) peaked in experiment 1 Fe treatment, with a minimum value of $1.06 \times 10^{-2}$ in experiment 5 control treatment. $E_k$ ($\mu$ mol photons m$^{-2}$ s$^{-1}$) peaked at 79.77, with minimum values in experiment 1 control treatment. Despite the substantial differences in setup conditions for experiments 1 and 2 in the SAZ, occupied twice over the space of 28 days, there were no significant differences in the responses of the $PE$ parameters to Fe. Due to constraints in light levels for the incubator setup, light levels that may result in photoinhibition...
Maximum percentage differences of $-12.10$ and $82.52\%$; the greatest value for $\Delta E_k$ was $-40.92$ for experiment 5. Maximal values of all differences were consistently found in experiment 5, which was set up just south of the Southern Boundary Front (Fig. 3).

Potential drivers of variability within the photosynthetic parameters were determined through a Pearson’s linear correlation coefficient matrix (Fig. 4), revealing significant negative and positive relationships with sea surface temperature (SST), salinity, nitrate and silicate concentrations; photosynthetic physiology parameters ($F_v/F_m$ and $\alpha_{PSII}$); and measures of the community structure, effective diameter and ratio of diatoms to haptophytes. There were no significant relationships with either dissolved iron concentrations or chlorophyll concentrations. Other parameters that did not show any relationships were excluded from the matrix include MLD, the light environment (in situ PAR and 1% light depth) and phosphate concentrations. $\alpha^B$ for the control treatments displayed the greatest number of relationships with SST, nitrate concentrations, community structure variables and $F_v/F_m$. The relative differences in all the parameters showed strong positive correlations with SST and salinity ($p<0.05$). A principal component analysis (PCA) was carried out on the data with the variables’ PCA projection on the factor plane represented in Fig. S4 in the Supplement. The sum of the first two PCs explained 76.74% of the total variance. The factor plane representation splits the variables, both experimental and initial conditions, into the four different quadrants. The grouping of the variables within each quadrant agree with the positive correlations determined within the correlation coefficient matrix, whereas variables in opposite quadrants agree with the negative correlations.

### 3.3 Primary production

Depth-integrated primary production (PP$_{wc}$) was calculated at each experimental location and displayed a wide range of variability with and without iron (Fig. 5). On average PP$_{wc}$ was higher in the iron addition treatments (Fig. 5a), with an average of $387.32 \pm 207.18$ (mg C m$^{-2}$ d$^{-1}$) for iron addition and an average of $315.37 \pm 229.37$ (mg C m$^{-2}$ d$^{-1}$) for the control. The maximum absolute differences in PP$_{wc}$ ($\Delta$PP$_{wc}$, Fig. 5b) of $228.82$ mc C m$^{-2}$ s$^{-1}$ was found in experiments 5 at $\sim 55^\circ$S near the Southern Boundary Front, with very little difference observed in $\Delta$PP$_{wc}$ at experiments 3 and 4.

The responses of Fe addition to primary production from the six experiments were extrapolated onto broader spatial and temporal scales, whereby underway measurements of Chl were converted into $K_d$ using Eq. (4). This, when combined with underway measurements of surface PAR, allowed us to look at latitudinal gradients in primary production (as per Eqs. 1, 2 and 3). As the $PE$ parameters displayed strong linear correlations with latitude ($\alpha R^2 = 0.73$ and 0.66, $P_{max}$ $R^2 = 0.91$ and 0.68 for Fe and Control respectively), a linear
Figure 4. Matrix of Pearson’s linear correlation coefficients between the photosynthetic parameters determined experimentally and in situ variables measured, including $\alpha_B^B$, $P_{\text{max}}^B$ and $E_k$ from the both Fe and control treatments; the relative differences; sea surface temperature (SST); salinity, nitrate, silicate and dissolved iron concentration; Chl concentration; effective diameter; $F_{v}/F_{m}$; $\sigma_{PSII}$; and community composition (ratio of diatoms to haptophytes). The strength of the linear relationship associated between each pair of variables is indicated by the colour of the square, with the negative and positive correlations denoted by “−” and “+” within all squares where significant ($p < 0.05$).

Figure 5. Modelled outputs of primary production utilizing experimentally derived photosynthetic parameters. (a) Depth-integrated primary production ($PP_{wc}$) (mg C m$^{-2}$ d$^{-1}$) and (b) $\Delta PP_{wc}$ (mg C m$^{-2}$ d$^{-1}$). Ocean fronts, indicated by grey lines, are displayed as in Fig. 3.
interpolation was applied to $P_{\text{max}}$ and $\alpha$, extrapolating the values from six points to a 0.1° resolution along the cruise track. The interpolated values of $P_{\text{max}}$ and $\alpha$ were combined with underway measurements of $K_d$ and PAR to calculate $PP_{\text{wc}}$ with and without Fe addition for the three different occupations of the same transect line (Fig. 6a). A high degree of variability was revealed between occupations in the SAZ and polar frontal zone (PFZ) but no clear differences between the iron and control treatments. Variability in the SAZ and PFZ appears to be temporally driven, with higher values of $PP_{\text{wc}}$ found in the third occupation of the transect line later in the summer season. Differences in $PP_{\text{wc}}$ between the two treatments become evident south of 50° S (Fig. 6a and b), with all three iron treatment occupations being $\sim 0.5 \text{ g C m}^{-2} \text{ d}^{-1}$ higher than their control treatment counterparts. The differences between the control and Fe treatments were calculated for each transect, which when combined allowed for the calculation of an average absolute difference in primary productivity ($\Delta PP_{\text{wc}}$, Fig. 6c). $\Delta PP_{\text{wc}}$ is slightly negative within the SAZ and PFZ, before sharply increasing to a maximum difference of $0.85 \text{ g C m}^{-2} \text{ d}^{-1}$ at 58° S. $\Delta PP_{\text{wc}}$ begins to decrease with increasing latitude before reaching an average difference of $0.11 \text{ g C m}^{-2} \text{ d}^{-1}$ in the MIZ. Representing these differences in $PP_{\text{wc}}$ as a percentage difference (Fig. 6d) shows that within the SAZ, PFZ and MIZ the differences are $\pm 10–20 \%$, whereas within the Antarctic zone (55–65° S) the differences between the treatments can be as much as 80%. Given the limitations of our data set (which requires the use of interpolated values of $P_{\text{max}}$ and $\alpha$) together with the weight we place on the conversion of these parameters to PP (with chlorophyll and PAR), it is important that we understand the sensitivity of the PP model to variability in the different input parameters. To test this, we performed a series of sensitivity tests to determine which components present the greatest influence on the final PP values. The sensitivity tests were divided into the three components of the equation: $K_d$ derived from chlorophyll (Fig. S5), surface PAR (Fig. S6) and the photosynthetic parameters ($P_{\text{max}}$ and $\alpha$) (Fig. S7). For consistency, the range of variation for each parameter was calculated and used as a factor to alter each component.
The mean range of variability for $K_d$ was 84.33 %, surface PAR was 68.73 %, and $\alpha$ and $P_{\text{max}}$ were 82.85 and 83.01 % respectively. If $K_d$ values are increased by 84.33 % this results in a 29.61 % decrease in $\Delta P_{\text{wc}}$, whereas a decrease of $K_d$ results in an increase in $\Delta P_{\text{wc}}$ of 59.17 %. Increasing surface PAR resulted in an increase in $\Delta P_{\text{wc}}$ of 3.50 %, whilst decreasing PAR corresponded to a decrease of 8.06 %. The largest differences in $\Delta P_{\text{wc}}$ were generated when $P_{\text{max}}$ was altered by 83.01 %, in accordance with the range of variability, resulting in an increase of 42.97 % and a decrease of 80.92 % in $\Delta P_{\text{wc}}$ (for an increase and decrease in $P_{\text{max}}$ respectively). The other PE parameter, $\alpha$, did not result in the same level of changes in $\Delta P_{\text{wc}}$ and only increased by 4.01 % and decreased by 12.22 % for an increase and decrease in $\alpha$ by 82.85 % respectively.

4 Discussion

Phytoplankton biomass in the Southern Ocean is potentially limited in their extent and magnitude predominantly by the availability of the micronutrient iron (Blain et al., 2007; Boyd et al., 2000; Pollard et al., 2009). This conclusion is based on the combination of two factors: the high iron requirements for photosynthetic proteins (Quigg et al., 2003; Raven, 1990; Strzepek and Harrison, 2004; Tving and Baines, 2013) and the lack of supply sources of iron to the Southern Ocean (Duce and Tindale, 1991; Tagliabue et al., 2014). The result of this is an environment that displays high degrees of spatial and temporal variability in primary production in response to highly variable iron supply mechanisms that result in chlorophyll patchiness (Fig. 1) and a complex seasonality (Thomalla et al., 2011). Iron limitation is potentially strongest during the summer months, when light levels are not considered limiting (Boyd et al., 2010) and the spring bloom is expected to have utilized the bulk of the winter iron resupply. In the austral summer of 2015/2016 a series of iron addition photosynthesis versus irradiance experiments were performed in the Atlantic sector of the Southern Ocean to determine the extent to which iron availability was limiting maximal rates of primary productivity.

The addition of iron appeared to stimulate increased productivity to varying degrees (Figs. 2, 3b, and S1–S3) with average $P_{\text{max}}$ and $\alpha$ values being higher for an iron-replete system ($12.75 \pm 6.95$ and $0.25 \pm 0.14$) compared to a control system ($11.17 \pm 8.23$ and $0.22 \pm 0.19$), suggestive that iron is indeed a micronutrient-limiting phytoplankton production in this region. Similar responses have been reported by Hiscock et al. (2008) under conditions of sub-saturating light conditions, where the addition of iron can result in a doubling of photosynthetic rates. However, a nutrient addition PE experiment in the Ross Sea demonstrated no significant increases in $\alpha$ or $P_{\text{max}}^B$ (Smith Jr. and Donaldson, 2015). One potential reason for this is the length of their incubation period, which was only 2 h and may not have been enough for the phytoplankton to incorporate the iron into their photosynthetic proteins and produce higher productivity rates. Indeed, nutrient addition experiments performed under similar conditions were shown to require 24 h to see any significant differences in initial changes in photophysiology (Brown- ing et al., 2014; Ryan-Keogh et al., 2017; Ryan-Keogh et al., 2013) with changes in biomass only being reported after 48 h. This shortcoming highlights the attraction of the unique experimental design utilized here, which allows for 24 h Fe addition and control incubations at varying light levels and constant temperature. However, it should be noted that a time length of 24 h may not be sufficient to complete alleviate the iron-mediated photosynthetic response and as such these results may only reflect initial responses rather than longer-term community-level responses to relief from iron limitation. It should be noted, however, that light acclimation can between 2 and 6 h and as such be reflected in the potential iron demand, a lower demand at higher irradiances (Strzepek et al., 2012). Such incidences would impact the observed differences between PE parameters in control versus Fe addition experiments. However, since the light range of the experiments (0–400) fall below the maximum light intensities measured in situ (Table 1), acclimation responses are unlikely to dominate and, if occurring, would indeed result in an underestimation of the differences between control and addition experiments. The experimental design of 24 h, whilst suitable for investigating iron limitation, means that results are not truly representative of in situ photosynthetic parameters and should not be interpreted as such.

Potential factors that are known to be associated with iron-induced enhanced primary productivity include temperature, macronutrient concentrations, Chl, MLD, light history and community composition. A Pearson’s linear correlation matrix (Fig. 4) was carried out on an array of variables to examine the influence of key physical, chemical and biological factors on the variability of photosynthetic parameters in this study. Significant relationships were found with SST; salinity and macronutrient concentrations, which show strong latitudinal gradients. A proxy for the community structure that utilized the ratio of the two dominant groupings (diatoms and haptophytes) also indicated strong significant relationships with the PE parameters, which is potentially driven by Si availability controlling community structure. Indeed, it has been demonstrated that in the SAZ, where haptophytes dominated during this study, there is evidence for Fe-Si co-limitation. In a study by Hutchins et al. (2001) it was demonstrated that the addition of both Fe and Si resulted in the greatest responses in chlorophyll and the photosynthetic parameters. The relationship here may not be driven by Fe availability on the PE parameters but rather community-level limitation. No significant relationships were, however, found between PE parameters and iron or Chl concentrations. The lack of significant relationships could be due to the small range of variability observed in these parameters; for example, Chl concentrations at all stations were typically low (0.84–2.30 µg L$^{-1}$) when compared to the range of chloro-
phyll concentrations measured throughout the entire cruise (0.01–11.25 µg L\(^{-1}\)). The lack of a relationship with dissolved iron concentrations highlights how this proxy is not necessarily a good indicator of iron stress, as any limiting nutrient would be expected to be severely depleted by biological uptake with a resultant ambient concentration that would remain close to zero despite possible event scale supply (Ryan-Keogh et al., 2017).

The photosynthetic parameters derived here are important components in a suite of models that derive estimates of phytoplankton primary production (Behrenfeld and Falkowski, 1997a, b; Sathyendranath and Platt, 2007). Different primary production models inherently consist of certain biases towards modelling the photosynthetic parameters, whereas others have excluded them entirely from the computation of primary productivity rates. Hiscock et al. (2008) demonstrated that the variables in the Behrenfeld and Falkowski (1997b) standard depth-integrated model (DIM) exerted considerably different forcing mechanisms on the final primary productivity rates. In the case of this DIM, phytoplankton biomass was the dominant variable that could result in 3 orders of magnitude changes in primary production, compared to only a 40-fold change when altering the photosynthetic parameter \(P_{\text{opt}}^B\) (i.e. \(b_{\text{max}}^B\)). This highlights the need to understand the sensitivity of different PP models to variability within their input parameters.

Results from the production model applied here (Eqs. 1–3) show a general decrease with latitude in depth-integrated primary production \((PP_{\text{wc}})\), with significant differences between treatments \((t\ test, p<0.05)\). One station near the Southern Boundary exhibited the greatest differences in \(\Delta PP_{\text{wc}}\) with a value of 0.89 g C m\(^{-2}\) d\(^{-1}\) (Fig. 5b), with the lowest observed \(\Delta PP_{\text{wc}}\) of 0.11 g C m\(^{-2}\) d\(^{-1}\) south of the polar front. The low sampling frequency of the experiments both spatially and temporally (six experiments spanning 2 months and the entire latitudinal extent of the Southern Ocean) together with the diverse range of initial setup conditions (Table 1) make it difficult to interpret the causal relationships observed within each experiment with any certainty. Instead, the information from these experiments was maximized through an alternate approach that utilized the range of variability in \(PE\) parameters in control versus iron addition experiments to gain a broader spatial interpretation of the response of phytoplankton production to iron addition.

A linear interpolation of the \(PE\) parameters \((P_{\text{max}}\) and \(\alpha\)) with latitude, together with underway measurements of PAR and \(K_d\) (derived from surface Chl) allow for the generation of high-resolution rates of \(PP_{\text{wc}}\) with and without Fe addition for three occupations of the cruise transect (Fig. 6a). Within the SAZ and PFZ there was a high degree of variability between the three occupations, with higher \(PP_{\text{wc}}\) values later in the growing season (Fig. 6a). However, there were no clear differences between the iron and control treatments in any of the occupations. This may not reflect a lack of iron limitation in the SAZ, as it has been demonstrated previously that there is ecological and physiological iron limitation (Coale et al., 2004), with longer experiments demonstrating increases in \(P_{\text{max}}\) and \(\alpha\) following iron addition (Hutchins et al., 2001). However, south of 50°S there were no differences observed as the growing season progressed with similar PPwc values across the three occupations of the cruise transect, but there was a clear difference between the iron and control treatments (Fig. 6b and c). Here, a maximum percentage difference of ~80 % (Fig. 6d) was observed between control and iron-replete conditions, with \(\Delta PP_{\text{wc}}\) peaking at 0.85 g C m\(^{-2}\) yr\(^{-1}\) at 55°S. Differences between iron addition and control systems begin to decline within the MIZ (Fig. 6c). These results suggest that there are potential differences in iron availability and supply within different zones of the Southern Ocean, which agrees with previous studies which postulated that the bloom extent and duration within the SAZ could potentially be driven by enhanced iron supply through storm–eddy interaction (Nicholson et al., 2016) while in the MIZ addition iron is supplied through melting ice (Gao et al., 2003; Grotti et al., 2005; Sedwick and DiTulio, 1997). The Fe addition test performed here demonstrates the sensitivity of waters south of 50°S to Fe availability. If models do not consider this sensitivity then the degree of error for PP models can be as high as 80 %. It must be noted that the transects will not only reflect latitudinal gradients but also contain a seasonal signal as the cruise spanned 2 months across the austral summer. A seasonal shift in community structure of haptophytes increasing their dominance beyond the SAZ into the PFZ was evident from underway measurements of community structure (data not shown), indicative of seasonal Si limitation for this region (Boyd et al., 2010). Moreover, the complex seasonality of this region represents shifts between varying co-limitations that will be represented not only in the \(PE\) parameters measured but also in the additional components utilized to calculate \(PP_{\text{wc}}\).

From these results, it became clear that higher values of \(P_{\text{max}}\) and \(\alpha\) because of iron addition were significantly influencing the model outputs of primary production. However, the extent to which changes in the \(PE\) parameters were responsible for the latitudinal trend in \(\Delta PP_{\text{wc}}\) versus changes in ancillary parameters (e.g. Chl, PAR) is unclear. To test our interpretation of the variability in \(PP_{\text{wc}}\) being a direct response to Fe availability through changes in the \(PE\) parameters, a series of sensitivity analyses were performed which showed that PAR and \(\alpha\) exerted very little influence (Figs. S6 and S7). Biomass (Chl), as represented through \(K_d\), did exert a large influence on \(PP_{\text{wc}}\) (up to 59 %, Fig. S5), but this influence could be overestimated due to potential errors in the calculation of \(K_d\) (Morel et al., 2007). However the greatest influence was \(P_{\text{max}}\) (up to 81 %, Fig. S7). As such, we can conclude that the primary driver of the latitudinal trend in \(\Delta PP_{\text{wc}}\) is the result of changes in the maximum photosynthetic capacity \((P_{\text{max}})\) to iron addition; however, regions along the transect may be experiencing seasonal co-limitation of Fe.
and Si, particularly during the third transect conducted during late summer.

The photosynthetic parameters $P_{\text{max}}$ and $\alpha$ remain difficult to fully parameterize due to interacting effects of iron, light availability, temperature and community structure, yet these parameters remain critical components of different biogeochemical models. Our results show that if models fail to capture the interacting effects of iron and other parameters on primary productivity, then the degree of error across vast extents of the Southern Ocean can be significant (as much as 80%). On the other hand, any model that can correctly account for variability in these parameters will better reproduce the natural background levels of primary productivity and the seasonal cycle for application to iron-limited areas of the ocean including the Subarctic Pacific and the Southern Ocean.

Data availability. Data used in this article can be found in the Supplement.

The Supplement related to this article is available online at https://doi.org/10.5194/bg-14-3883-2017-supplement.

Competing interests. The authors declare that they have no conflict of interest.

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3895


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