Role of krill versus bottom-up factors in controlling phytoplankton biomass in the northern Antarctic waters of South Georgia

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ABSTRACT: The extent to which Antarctic phytoplankton stocks are controlled by ‘bottom-up’ and/or ‘top-down’ factors is highly variable. Here we consider data collected at South Georgia during 3 summer surveys that recorded substantial hydrographic variability. A suite of bottom-up and top-down controlling factors were measured simultaneously at the mesoscale. Sea surface temperature varied by >2°C, macronutrients ranged from near-winter concentrations to near-depleted, while mean densities of a major grazer, krill Euphausia superba, varied between near-zero and >400 g wet mass m−2. A general linear model was used to identify the main factors implicated in the observed differences in phytoplankton biomass. Despite east-to-west and on- to off-shelf temperature gradients, temperature per se was not implicated in phytoplankton variability. Also, while there was an abundance of NO3-N in surface waters, NH4-N was the key nutrient throughout. A domed relationship between phytoplankton and krill peaked between 2 and 4 mg chlorophyll a m−3 and 6 and 30 g krill m−2. The positive side of this dome was represented by the west off-shelf region downstream of South Georgia. Here, an ample supply of micro- and macronutrients promoted high primary production, and low densities of krill presumably had little grazing effect. This positive relationship between krill and phytoplankton biomasses was interpreted as krill accumulating in areas of good feeding conditions. The negative side of the dome was typified by the east off-shelf region, where macronutrients remained high, primary production rates were low, and krill densities were very high. The grazing rates calculated here suggested that krill affect their food stocks severely, and the negative krill–phytoplankton relationship in this region may reflect locally high krill densities driving down their food supply.

KEY WORDS: Antarctic phytoplankton · Macronutrients · NH4 · Temperature · Grazing effect

INTRODUCTION

The extent to which Antarctic phytoplankton stocks are controlled by ‘bottom-up’ physical and chemical factors or by ‘top-down’ grazing has been debated for decades (e.g. Hardy & Gunther 1935, Hart 1942, Chisholm & Morel 1991, Atkinson et al. 2001, Ward et al. 2002, 2005, Smetacek et al. 2004, Smith & Lancelot 2004, Harris 2005, Murphy et al. 2007). To date, bottom-up factors have probably received the most attention, doubtless because it is easier to measure temperature, mixed layer depths and nutrients than to quantify grazing by a plethora of patchily distributed grazer species. Furthermore, the simultaneous measurement of bottom-up and top-down control is fraught with difficulty (Martin 2003), especially in the Southern Ocean where sampling opportunities are infrequent.

Iron enrichment experiments have demonstrated unequivocally that, over large parts of the Southern Ocean, phytoplankton are under bottom-up control due to a chronic lack of iron in surface waters (e.g. Boyd et al. 2007). Smith & Lancelot (2004) argued further that bottom-up control was strongest for the large, bloom-forming diatoms, since their large size offered a
refuge from predation from the protozoans that gener-
ally dominate the grazer assemblage (Calbet & Landry
2004). In contrast, the meso- and macroplankton that
can deal with the large or chain-forming diatoms have
longer generation times, and thus cannot increase their
populations fast enough to keep a bloom in check (Smetacek et al. 2004).

While these generalities may hold, local or regional
exceptions may exist. One that has received attention is the
potential for Antarctic krill Euphausia superba
(hereafter krill) to substantially reduce their food sup-
ply. Krill are important grazers in large, productive,
mid-latitude subsystems of the Southern Ocean, often
with ‘key species’ status due to their large biomass,
high energy demands and prominent role in predator
species (Atkinson et al. 2001). While the production and
grazing effect of krill is minor compared to protozoans
and mesozooplankton over the Southern Ocean as a
whole (Voronina 1998, Pakhomov et al. 2002, Shreeve
et al. 2005), their very uneven distribution is key, as
compared to the other grazers. Thus, in areas where
krill swarms are locally numerous they have often been
suggested to have a major grazing effect, with biogeo-
chemical ramifications (reviewed by Smetacek et al.
2004). However, much of the evidence for this has been
anecdotal, and only rarely has it been possible to moni-
tor suitable indices of phytoplankton, krill, and the po-
tential bottom-up control factors simultaneously.

Where such studies have been conducted (e.g.
Hardy & Gunther 1935, Weber & El-Sayed 1985,
Weber et al. 1986, Lawson et al. 2008), they have ar-
rived at a variety of conclusions. Both positive and neg-
ative relationships between phytoplankton and krill
have been found, with the former interpreted as the
krill aggregating in good feeding grounds and the lat-
ter implying that krill have grazed down their food
(Weber et al. 1986). The sign of this relationship has
been suggested to depend partly on the scale of analy-
sis, and various positive, negative and dome-shaped
relationships have been found across a wide range of
scales (e.g. Weber et al. 1986, Atkinson et al. 2008,
Lawson et al. 2008).

In this study, we selected South Georgia as a study
site to examine the control on phytoplankton via bot-
tom-up factors and krill grazing. While such productiv-
ity hotspots comprise only a fraction of the Southern
Ocean, they are important biogeochemically in terms of
total carbon flows and for commercially important
food webs (Atkinson et al. 2001, Schlitzer 2002, Duck-
low et al. 2006). South Georgia is also particularly well
suited to such a study. First, during summer it is prob-
bly iron replete (Holeton et al. 2005, Korb et al. 2008,
Whitehouse et al. 2008), making it easier to consist-
tently assess the effects of other bottom-up factors (e.g.
macronutrients, temperature, mixed-layer depths)
across multiple cruises. Second, there is good back-
ground knowledge of this area, with extensive datasets
on physics, nutrients, phytoplankton and krill (e.g.
Whitehouse et al. 1996, Brierley et al. 1997, Trathan et

We used data from 3 summer surveys to the north of
South Georgia to examine the controls on phytoplank-
ton biomass. The surveys were conducted in consecu-
tive years during which there was substantial hydro-
graphic variability within the region (Meredith et al.
2005). We examined the interactions observable at the
mesoscale (10s of km) by means of regressions, gener-
alised additive models (GAMs) and general linear
models (GLMs). These in effect use the variation in
bottom-up factors and at the mesoscale krill biomass as
a kind of ‘natural experiment’ to identify the main fac-
tors implicated in the observed differences in phyto-
plankton biomass.

MATERIALS AND METHODS

Survey methods. Underway oceanographic and bio-
acoustic data were collected simultaneously within 2
survey grids: 1 to the northwest and another to the north-
east of South Georgia (Fig. 1). Survey grids comprised
10×80 km long transects, roughly perpendicular to the
island’s shelf-break. While surveying, the nominal ship
speed was 10 knots. We surveyed 2 transects d−1, and
each survey grid took 5 d to complete. Data were col-
clected during 3 consecutive years: 1996/97 (cruise JR17,
December 1996 and January 1997); 1997/98 (cruise
JR28, January and February 1998); 1998/99 (cruise JR38,
December 1998 and January 1999). In addition, on- and
off-shelf CTD casts were made during most survey days
within each surveyed region.

Krill acoustics data. The transects were surveyed
primarily to acoustically estimate the biomass of krill,
as part of a long-term monitoring programme (Brierley
et al. 1999, Murphy et al. 2007). Acoustic data were
collected solely during daylight hours to reduce the
possibility of bias due to diurnal migration (Demer &
Hewitt 1995) or change in target orientation (Everson
1982). Volume backscattering strength (Sv, dB re 1 m−1)
was collected using a Simrad EK500 (38, 120 kHz) cal-
ibrated using the standard sphere method (Foote et al.
1987). The ping rate varied between 1.5 and 2.5 s, but
was constant within a survey.

All acoustic data were processed using Echoview
software (ver. 4.20.59.8698); relevant values for the
speed of sound and absorption coefficients were input
derived from concurrent CTD data); surface noise,
false bottom echoes and interference were identified
and removed; and time varied gain amplified back-
ground noise was subtracted (Watkins & Brierley
Cleaned data for each frequency were averaged into 5 m (vertical) by 500 m (horizontal) integration cells and imported into Matlab where any further manipulation or analysis was undertaken.

$S_v$ at 120 kHz ($S_{120\text{kHz}}$) were apportioned to krill using a variable dB window differencing identification technique using the 120 and 38 kHz ($S_{120-38}$) frequencies (Method 2, Reiss et al. 2008). Window ranges for different size ranges of krill (identified from the net samples in each survey region in each year) were taken from the current agreed Commission for the Conservation of Antarctic Marine Living Resources (CCAMLR) protocol for krill $B_0$ assessment (CCAMLR 2009).

The $S_{120\text{kHz}}$ attributable to krill were integrated from 10 m below the surface to either a maximum of 250 m or 2 m above the seabed, resulting in nautical area scattering coefficient (NASC; m$^2$ nautical mile$^{-2}$) values. NASC were converted to krill biomass (g m$^{-2}$) using the simplified stochastic distorted-wave Born approximation (SDWBA) target strength model, which was applied using fixed values for orientation (11, 4) and material properties based on those calculated/inferred for the CCAMLR synoptic survey in the Scotia Sea region at the same time of year as the present study (Conti & Demer 2006, CCAMLR 2009), and weighted mean masses per individual, using the mass to length relationship also collected during the CCAMLR Scotia Sea survey (Hewitt et al. 2004).

**Oceanographic, nutrient and chlorophyll data.** The underway sample source for chlorophyll (chl) and nutrient samples was the ship’s non-toxic supply at ~7 m below sea surface. A Turner Model 10 through-flow fluorometer was used to measure $in\ vivo$ fluorescence, while discrete samples, taken hourly from the same source, were extracted with acetone and analysed with a Sequoia-Turner Model 112 benchtop fluorometer. The benchtop fluorometer was calibrated against a standard prepared from chl extracted from the cyanobacterium *Anacystis nidulans* (Sigma Chemicals; Parsons et al. 1984). Underway chl was estimated from $in\ vivo$ fluorescence with a model that used simultaneous measurements of incident photosynthetically active radiation and the high frequency discrete samples to allow for anomalies such as those caused by light-quenching and variations in phytoplankton community composition. The fitted model accounted for >96% of the variance in the underway measurements (A. W. A. Murray & J. Priddle, British Antarctic Survey [BAS], unpubl. report). Macronutrients were monitored continuously by feeding a filtered sample stream to a Technicon autoanalyser (Whitehouse 1997, Woodward et al. 1999). The same analytical methods were used to measure chl and macronutrients in discrete CTD water bottle samples. Underway temperature was measured with a sensor at

Fig. 1. (a) Location of South Georgia (SG) relative to the mean positions of the Antarctic Polar Front (APF) and the Southern Antarctic Circumpolar Current Front (SACCF). Pale and dark grey shading are delineated by the 1000 and the 3000 m isobaths, respectively. (b) West (WS) and east (ES) survey transects along with CTD station locations (o) in relation to South Georgia, the 1000 m isobath, the mean position of the SACCF and the 0.1° latitude by 0.2° longitude grid by which samples were grouped. (c) General current flow in relation to South Georgia, the 1000 m isobath, and the mean positions of the APF and the SACCF (from Korb et al. 2004)
the non-toxic supply's intake on the ship's hull, and was periodically checked against CTD measurements.

Calculation of nutrient depletion. To enable regional comparisons to be made, nutrient deficits were calculated by subtracting measured surface values from an estimated generic pre-bloom concentration for South Georgia of 35, 2.1 and 30 mmol m\(^{-3}\) for Si(OH)\(_4\), PO\(_4\) and NO\(_3\), respectively (Whitehouse et al. 1996, 2000, 2008, BAS unpubl. data). These calculated deficits should be regarded only as an index of nutrient use, as upwelling and re-supply doubtless occur to some extent at the island's shelf-break. To identify preferential nutrient use, e.g. greater nitrate utilisation, we then calculated ratios of nutrient deficits between pairs of nutrients.

Scale of data analysis. The underway data were collected at various frequencies: once per minute for temperature and chl \(a\), once per 10 s for macronutrients, while acoustic records were averaged into 0.5 km lengths of transect. Therefore, a common scale for all of the data needed to be ascertained to allow further analysis, with the finest data resolution imposed by the acoustic measurements: 0.5 km section of transect. Each survey grid took 5 d to complete, and transects were spaced ~7 to 10 km apart. Current velocities to the north of South Georgia (particularly off-shelf) can be as high as 50 cm s\(^{-1}\) (Brandon et al. 2000, Meredith et al. 2003a,b, Whitehouse et al. 2008), so water could potentially be transported across a survey grid in ~2 d. Given the high current velocities, passive nature of phytoplankton and nutrients in contrast with mobile krill, and the heavily skewed and aggregated krill distribution we found, the finest data resolution was inappropriate to resolve relationships that may have resulted through advection. Instead, a coarser scale was required. Given the influence of physical processes (e.g. Huntley & Niiler 1995) on the distribution of phytoplankton, krill, and nutrients, we were guided by the horizontal scale of eddies as quantified by internal Rossby radius, which is ~10 km at the latitude of South Georgia (Houry et al. 1987). With transects spaced ~7 to 10 km apart, this is the finest resolution we could use that might allow for effects brought about by advection downstream. A 0.1° latitude by 0.2° longitude grid around South Georgia conveniently produces a cell size of approximately 11 x 13 km, so we made our comparisons at this scale with mean values calculated for each grid cell (Fig. 1b).

We treated the west and east surveys separately for many of our comparisons, since their conditions vary substantially (e.g. Brierley et al. 1999, Korb & Whitehouse 2004, Meredith et al. 2005). For the same reason, we divided the surveys into shelf (<1000 m) and oceanic (>1000 m) sub-regions. Most of the island's shelf is <500 m depth, but the 1000 m isobath provides a much better delineation of the shelf break.

Data analysis was carried out with analyses of variance (ANOVAs), GAMs and GLMs in the Minitab program. The degrees of freedom of the fitted GAMs were chosen by cross-validation, i.e. minimising the sums of squares of deviations from the fitted values excluding 1 data point at a time.

To further elucidate inter- and intra-survey box differences, longitude was also included as an independent variable in the GLMs, despite the different orientation of the 2 transect grids. Given the variability in the predominantly east-to-west current flow and the counterclockwise rotation of 'north-south' Antarctic Circumpolar Current (ACC) characteristics to the north of South Georgia, we considered a simple unweighted east–west measurement to be a suitable spatial factor.

RESULTS

Surface data compared to the top 50 m of the water column

As our only sample source during underway surveying was the ship's non-toxic supply (~7 m below sea surface) and primary production measurements would usually be made over at least the euphotic layer, we needed to assess how representative our surface temperature and concentrations of chl \(a\) and macronutrients were of the water column below. We examined the relationship between mean surface (0 to 7 m) CTD-derived chl \(a\), temperature and nutrient data and those averaged over 0 to 50 m at the CTD stations within the surveyed areas (Table 1). Regressions between all parameters were highly significant (p < 0.001) with R\(^2\) values ≥87%. This relationship concurs with previous measurements in this area (see Gilpin et al. 2002, Korb et al. 2004, Whitehouse et al. 2008) in showing a good correlation between surface data and those collected within the top 50 m of the water column.

Table 1. Relationship between mean surface (0 to 7 m) CTD-derived chl \(a\), temperature and nutrient data and those averaged over 0 to 50 m. All available data from the 3 cruises were used, and station localities are indicated in Fig. 1b. Surface values = \(x + (y \times \text{near-surface values})\), and all regressions were highly significant (p < 0.001)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>n</th>
<th>R(^2)</th>
<th>(x)</th>
<th>(y)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Log(_10) chl (a)</td>
<td>16</td>
<td>96</td>
<td>0.0328</td>
<td>1.087</td>
</tr>
<tr>
<td>Temperature</td>
<td>44</td>
<td>88</td>
<td>0.2891</td>
<td>0.9966</td>
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<tr>
<td>Si(OH)(_4)</td>
<td>29</td>
<td>98</td>
<td>-0.1938</td>
<td>0.9848</td>
</tr>
<tr>
<td>PO(_4)</td>
<td>31</td>
<td>94</td>
<td>0.0005</td>
<td>0.9748</td>
</tr>
<tr>
<td>NO(_3)</td>
<td>31</td>
<td>96</td>
<td>-0.0541</td>
<td>0.9868</td>
</tr>
<tr>
<td>NH(_4)</td>
<td>31</td>
<td>87</td>
<td>-0.1245</td>
<td>0.9687</td>
</tr>
</tbody>
</table>
Spatial and temporal variability

All parameters showed variability from year to year, from west to east, and from on-shelf to off-shelf. In the first 2 surveys, chl \(a\) concentrations were significantly higher in the west than the east \((p < 0.001)\) but differed less during the third survey \((\text{Fig. 2a})\). They differed less from on-shelf to off-shelf waters than they did either between west and east or between years. Highest values \((\text{mean } \pm \text{SD}: 6 \pm 2.3 \text{ mg m}^{-3})\) were found in west off-shelf waters during 1996/97 and the lowest \((0.7 \pm 0.14 \text{ mg m}^{-3})\) in east on-shelf waters during 1997/98.

In the first 2 surveys, krill densities were significantly higher on the east but were more uniformly distributed during 1998/99 \((\text{Fig. 2b})\). Within survey boxes, densities were significantly higher on-shelf during the first 2 surveys but again were more uniformly distributed during 1998/99. Highest values \((409 \pm 416 \text{ g wet mass m}^{-2})\) were found in east on-shelf waters during 1997/98 and the lowest \((5.1 \pm 7.8 \text{ g m}^{-2})\) in west off-shelf waters during the same season.

Ocean temperature within the sub-regions varied by \(-2^\circ\text{C}\) \((\text{mean } 1.5 \text{ to } 3.5^\circ\text{C})\) over the 3 surveys \((\text{Fig. 2c})\). During the 2 earlier surveys, temperatures were significantly lower in the west \((p < 0.001)\) but were more uniform during 1998/99. In 1996/97, the greatest on- to off-shelf temperature difference \((-0.95^\circ\text{C})\) was found in the east, while overall, the lowest temperatures were encountered during 1997/98.

\(\text{Si(OH)}_4\) concentrations were particularly high during 1997/98 \((>20 \text{ mmol m}^{-3}\) in most regions), while the lowest were found in 1998/99 \((<10 \text{ mmol m}^{-3}\) in most regions; \(\text{Fig. 2d})\). In 1996/97, particularly low values \((5.2 \pm 6.2 \text{ mmol m}^{-3})\) in west off-shelf waters contrasted with all other regions where mean concentrations were \(>17 \text{ mmol m}^{-3}\). \(\text{PO}_4\) and \(\text{NO}_3\) concentrations were closely related, and the lowest values were always found in the west \((\text{Fig. 2e,f})\). During 1997/98, particularly low values \((-0.5 \text{ and } 12 \text{ mmol m}^{-3},\) respectively\) were found in marked contrast to ambient \(\text{Si(OH)}_4\) concentrations. These lowest ambient \(\text{NO}_3\) values in the west were coincidental with high \(\text{NO}_3:\text{PO}_4\) deficit ratios of \(-16:1\) and low \(\text{Si(OH)}_4:\text{NO}_3\) deficit ratios of \(-1\) to \(2\) \((\text{Fig. 3})\). Overall, \(\text{NH}_4\) concentrations were highest \((>0.5 \text{ mmol m}^{-3}\) in most regions\) during the 1997/98 survey with lower values \(<0.5 \text{ mmol m}^{-3}\) in most regions\) evident during 1998/99 \((\text{Fig. 2g})\). During 1996/97, the greatest west-to-east difference \((0.74 \text{ mmol m}^{-3}\) in on-shelf waters\) was found with the lowest concentrations \((0.18 \pm 0.08 \text{ mmol m}^{-3})\) throughout the 3 surveys in western waters.

![Fig. 2. Mean (+SD) values for the west and east (key in b), on- and off-shelf sub-regions during the 3 sampling years. (a) Chl a, (b) Euphausia superba density, (c) temperature, (d) Si(OH)_4, (e) PO_4, (f) NO_3, (g) NH_4. (b) Particularly large SD of krill density are noted on the plot as opposed to being graphically represented.](image-url)
The relationship between chl a and krill is complex, and a year-by-year comparison indicated a highly variable and non-linear relationship (Fig. 4a). In summers with abundant krill, 1996/97 and 1997/98, the GAMs showed domed relationships with the respective log10 chl a peaks of 0.62 and 0.35 mg m\(^{-3}\) at log10 krill of 0.81 and 1.0 g m\(^{-2}\). There was a relatively flatter response for 1998/99 with a 0.52 mg m\(^{-3}\) log10 chl a peak at a log10 krill density of 1.47 g m\(^{-2}\). To summarise, in real terms, the GAMs indicated chl a peaks between 2.2 and 4.2 mg m\(^{-3}\) at krill densities between 6.5 and 29.5 g m\(^{-2}\).

If we consider data from within our sub-regions, then some distinct chl a/krill trends emerge. The west off-shelf sub-region is atypical with chl a concentrations positively related to krill abundance (Fig. 5a). Overall, and for the 1998/1999 summer, this relationship is highly significant (p < 0.001). In all other sub-regions, chl a concentrations are negatively related to krill abundance. Overall, the relationship is highly significant (p < 0.001) in both east off- and on-shelf waters where krill are most abundant, but it is also negative in west on-shelf waters (p < 0.01).

During the first 2 surveys, chl a related positively to temperature. A smaller range of values appeared to produce a slightly negative relationship in 1998/99 (Fig. 4b). Highest concentrations of chl a were generally confined to waters >2°C.

There was a consistently negative relationship between chl a and Si(OH)\(_4\), PO\(_4\) and NO\(_3\) (Fig. 4c,d,e), but note the smaller range of data measured during 1998/99. High chl a values during 1996/97 coincided with very low Si(OH)\(_4\) concentrations. By contrast, a large part of the 1997/98 survey was conducted in low chl a waters where Si(OH)\(_4\) remained close to pre-bloom, winter concentrations (Whitehouse et al. 1996, 2000). However, this apparent under-utilisation of Si(OH)\(_4\) in 1997/98 was in stark contrast with PO\(_4\) and NO\(_3\) use, when some of the lowest concentrations for these nutrients were measured. A negative relationship between chl a and NH\(_4\) appeared to be consistent for the 2 earlier surveys, but a small NH\(_4\) range in 1998/99 resulted in virtually no relationship with chl a (Fig. 4f).

Multivariate examination of chl a variability

Given the hydrological dynamics around South Georgia with rapid current flow in off-shelf waters (primarily east to west; Fig. 1c) and a varying degree of water retention over its shelf (Meredith et al. 2005), we assessed chl a variability separately for the on- and off-shelf regions. We modelled log10 chl a concentration variability with year, longitude, log10 krill density, temperature and nutrient concentrations as explanatory factors (Table 2). Longitude was a significant (p < 0.001) explanatory factor in both on-shelf and off-shelf waters. Also, for on-shelf waters, krill, temperature, Si(OH)\(_4\) and NH\(_4\) were highly significant predictors, and the model explained ~83% of chl a variability. For off-shelf waters, with the exception of krill and Si(OH)\(_4\), all predictors were highly significant (p < 0.001), and the model explained ~80% of chl a variability. Significantly higher concentrations of chl a occurred in the west, as has been documented previously (e.g. Atkinson et al. 2001, Korb et al. 2004), so to further explore the relative roles of the predictor variables, it was necessary to consider the on- and off-shelf sub-regions separately in the west and east surveys.
For the west on-shelf sub-region, NH$_4$ was the most significant predictor variable for chl $a$ concentration, while Si(OH)$_4$ ($F = 3.7, p = 0.06$) and krill ($F = 3.4, p = 0.07$) were near-significant (Table 2). These variables were all negatively related and together with the minor factors explained ~57 % of chl $a$ variability. For the east on-shelf sub-region, longitude remained a significant predictor followed by Si(OH)$_4$ and to a lesser degree NH$_4$ and krill. Both nutrients and krill were negatively related and together with the minor factors explained ~57 % of chl $a$ variability. For the west on-shelf sub-region, NH$_4$ was the most significant predictor variable for chl $a$ concentration, while Si(OH)$_4$ ($F = 3.7, p = 0.06$) and krill ($F = 3.4, p = 0.07$) were near-significant (Table 2). These variables were all negatively related and together with the minor factors explained ~57 % of chl $a$ variability. For the east on-shelf sub-region, longitude remained a significant predictor followed by Si(OH)$_4$ and to a lesser degree NH$_4$ and krill. Both nutrients and krill were negatively related and together with the minor factors explained ~57 % of chl $a$ variability. For the west on-shelf sub-region, NH$_4$ was the most significant predictor variable for chl $a$ concentration, while Si(OH)$_4$ ($F = 3.7, p = 0.06$) and krill ($F = 3.4, p = 0.07$) were near-significant (Table 2). These variables were all negatively related and together with the minor factors explained ~57 % of chl $a$ variability. For the east on-shelf sub-region, longitude remained a significant predictor followed by Si(OH)$_4$ and to a lesser degree NH$_4$ and krill. Both nutrients and krill were negatively related and together with the minor factors explained ~57 % of chl $a$ variability.

Fig. 4. Relationship during the 3 sampling years (identified in f) of log$_{10}$ chl $a$ with (a) log$_{10}$ Euphausia superba density, (b) temperature, (c) Si(OH)$_4$, (d) PO$_4$, (e) NO$_3$, (f) NH$_4$. For the log$_{10}$ krill density comparisons, generalised additive models were used and the degrees of freedom were 5 for years 1996/97 and 1997/98 and 3 for 1998/99. The other relationships are simple linear regression lines to indicate trends.
Fig. 5. Relationship between log$_{10}$ chl $a$ and log$_{10}$ *Euphausia superba* density during the 3 sampling years and overall (identified in c) in the sub-regions (a) west off-shelf, (b) east off-shelf, (c) west on-shelf, (d) east on-shelf. Simple linear regression lines are used to indicate trends. Note the different x-axis scales used for each panel.

Table 2. Summary of general linear model relating log$_{10}$ chlorophyll to longitude, log$_{10}$ *Euphausia superba*, temperature, Si(OH)$_4$, PO$_4$, NO$_3$, NH$_4$, and year (as a categorical variable). Regional coefficients (standard error) are detailed and significant slopes are indicated thus: ***$p < 0.001$, **$p < 0.01$, *$p < 0.05$. The year effects are deviations around zero.

<table>
<thead>
<tr>
<th></th>
<th>West and East</th>
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<td>R$^2$</td>
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<td>57 %</td>
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<td>$-0.03$ (0.05)</td>
<td>$-0.31$ (0.04)***</td>
<td>$-0.08$ (0.01)***</td>
<td>$-0.06$ (0.05)</td>
<td>$-0.30$ (0.07)***</td>
<td></td>
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<tr>
<td>Log$_{10}$ krill</td>
<td>$-0.07$ (0.02)***</td>
<td>$-0.05$ (0.03)</td>
<td>$-0.04$ (0.02)*</td>
<td>$-0.01$ (0.02)</td>
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<td>Temperature</td>
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<td>0.08 (0.08)</td>
<td>0.05 (0.05)</td>
<td>$-0.22$ (0.04)***</td>
<td>$-0.36$ (0.08)***</td>
<td>$-0.09$ (0.06)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Si(OH)$_4$</td>
<td>$-0.01$ (0.003)***</td>
<td>$-0.01$ (0.003)</td>
<td>$-0.02$ (0.01)***</td>
<td>$-0.01$ (0.004)*</td>
<td>$-0.01$ (0.01)*</td>
<td>$-0.02$ (0.01)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>PO$_4$</td>
<td>$-0.05$ (0.13)</td>
<td>0.03 (0.17)</td>
<td>$-0.15$ (0.19)</td>
<td>$-0.51$ (0.15)***</td>
<td>$-0.15$ (0.20)</td>
<td>$-1.23$ (0.26)***</td>
<td></td>
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</tr>
<tr>
<td>NO$_3$</td>
<td>$-0.004$ (0.01)</td>
<td>$-0.01$ (0.01)</td>
<td>0.0003 (0.01)</td>
<td>$-0.03$ (0.01)***</td>
<td>$-0.04$ (0.01)***</td>
<td>$-0.02$ (0.01)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>NH$_4$</td>
<td>$-0.17$ (0.03)***</td>
<td>$-0.29$ (0.09)***</td>
<td>$-0.09$ (0.04)*</td>
<td>$-0.40$ (0.07)***</td>
<td>$-0.50$ (0.10)***</td>
<td>$-0.21$ (0.09)*</td>
<td></td>
<td></td>
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<tr>
<td>1996/97</td>
<td>0.025 (0.04)</td>
<td>0.04 (0.06)</td>
<td>0.07 (0.06)</td>
<td>0.29 (0.04)***</td>
<td>0.37 (0.07)***</td>
<td>0.33 (0.09)***</td>
<td></td>
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<tr>
<td>1997/98</td>
<td>$-0.05$ (0.04)</td>
<td>$-0.04$ (0.06)</td>
<td>$-0.05$ (0.09)</td>
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<td>$-0.38$ (0.09)***</td>
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<td>1998/99</td>
<td>0.025 (0.03)</td>
<td>0.001 (0.04)</td>
<td>$-0.02$ (0.08)</td>
<td>0.06 (0.03)*</td>
<td>0.01 (0.05)</td>
<td>$-0.12$ (0.07)*</td>
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explained ~87% of chl a variability. For the west off-shelf sub-region, temperature was the most significant explanatory factor followed by NH4, then NO3 and to a lesser degree Si(OH)4. The nutrients were negatively related to chl a, as was temperature, and with the minor factors they explained ~78% of chl a variability. For the east off-shelf sub-region, PO4 was the most significant explanatory factor along with longitude followed by krill and NH4. PO4 and krill were negatively related to chl a, and together with the minor factors explained ~77% of its variability. When considering these 4 sub-regions, NH4 was the most important predictor and was significant in all of them. Longitude, Si(OH)4 and krill were the next most important predictors, followed by NO3, PO4 and temperature.

The year coefficients give an indication of temporal variability in the system (Table 2). On-shelf measurements were consistent throughout, whereas most of those off-shelf contained significant year-to-year variability.

**DISCUSSION**

These 3 surveys illustrate the variability that exists between macronutrients, phytoplankton abundance and krill density within a relatively small area, roughly 330 km wide. Most of this variability is due to the influence of the extensive shelf of South Georgia and its location relative to the major fronts of the ACC. The Scotia Arc steers the generally eastward flowing ACC northwards past South Georgia so that the usual north-south temperature gradients associated with the ACC are rotated counter-clockwise as they pass the island. If our chl a and krill data are considered in relation to this local west-east temperature gradient, it is apparent that the warmer western waters hold a greater standing stock of phytoplankton while the colder waters to the east hold greater densities of krill (Fig. 6). The present dataset allows us to make observations over multiple trophic levels. Below we examine the evidence for bottom-up and top-down controls on phytoplankton stocks at the mesoscale.

**Bottom-up controls on phytoplankton**

Over large scales, bottom-up controls on phytoplankton appear to be dominant. In addition to the major west–east gradient across the ACC, the presence of the South Georgia shelf had a major influence. The retention of water over the island’s shelf has major physical and chemical implications for the region’s primary production (e.g. Meredith et al. 2005, Whitehouse et al. 2008). Our model indicated year-to-year consistency over the island shelf, whereas there was significant inter-annual variability in off-shelf waters, particularly in 1996/97 and 1997/98. During these 2 earlier surveys, on- and off-shelf waters lay either side of a distinct shelf-break front, whereas during 1998/99 there was little difference between on- and off-shelf water mass densities (Meredith et al. 2005). This may account for the consistency found throughout the sub-regions during 1998/99 and infers exchange between on- and off-shelf waters during that season. Conversely, retention over the island shelf during the earlier surveys might buffer year-to-year variability, while rapidly flowing off-shelf waters are more susceptible to remote change (Meredith et al. 2005).

In both on- and off-shelf regions, chl a concentrations in the west were greater than those in the east (Fig. 2), and this was paralleled by 5- to 9-fold greater primary production rates (Table 3). This could reflect higher water temperatures in the west, which might promote NO3 uptake (Reay et al. 2001, Shreeve et al. 2002). Indeed, our analyses of all on-shelf and all off-shelf regions included temperature as a significant
Table 3. Sub-region mean (plus range where available) primary production rates (g C m\(^{-2}\) d\(^{-1}\)) integrated over the euphotic zone of 0.8 to 1.8 g C m\(^{-2}\) d\(^{-1}\) (Table 3) would total 96 to 216 g color.gsfc.nasa.gov/). Typical primary production rates

<table>
<thead>
<tr>
<th>Season</th>
<th>On-shelf (&lt;1000 m)</th>
<th>Off-shelf (&gt;1000 m)</th>
<th>Source</th>
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</thead>
<tbody>
<tr>
<td></td>
<td>West</td>
<td>East</td>
<td>West</td>
</tr>
<tr>
<td>1993/94</td>
<td>1.48 (0.84–2.48)</td>
<td>0.07</td>
<td>1.66</td>
</tr>
<tr>
<td>1995/96</td>
<td>0.78 (0.58–1.15)</td>
<td>0.14 (0.12–0.14)</td>
<td>1.18 (0.15–2.50)</td>
</tr>
<tr>
<td>2001/02</td>
<td>2.10 (0.84–3.34)</td>
<td>–</td>
<td>1.97 (1.25–2.55)</td>
</tr>
<tr>
<td>2002/03</td>
<td>0.50 (0.36–0.64)</td>
<td>–</td>
<td>0.81 (0.57–1.38)</td>
</tr>
<tr>
<td>Mean</td>
<td>1.22</td>
<td>0.24</td>
<td>1.81</td>
</tr>
</tbody>
</table>

predictor. However, at a smaller scale, temperature remained a significant predictor in one sub-region only, the west off-shelf, where it varied negatively with chl \(a\). This suggests that temperature per se is not the main predictor of chl \(a\) variability, and instead, elevated phytoplankton growth is facilitated by other factors and happens to occur in warm waters to the west.

Our data show a close relationship between phytoplankton standing stock and nutrient use with lowest ambient nutrient concentrations evident in the west. High phytoplankton standing stock and growth rates in waters northwest and downstream of South Georgia have been attributed to relief from Fe stress (e.g. Korb & Whitehouse 2004), brought about by sedimentary flux from the island’s extensive shelf and/or upwelling of deep nutrient-rich waters. Relief from Fe stress would allow greater utilisation of NO\(_3\), and indeed, our GLM indicates that NO\(_3\) is a significant predictor of chl \(a\) in west off-shelf waters, downstream of the island mass (Table 2). However, NH\(_4\) was also a significant factor in this sub-region, indicating that both N sources were important downstream of South Georgia. In all other sub-regions, recycled NH\(_4\) was the most important N source.

Temperature’s negative relationship with chl \(a\) in west off-shelf waters might be related to the upwelling of cool, nutrient-rich water (Meredith et al. 2005). The magnitude and longevity of blooms to the west are dependent on a persistent supply of macro- and micronutrients from upstream or via upwelling or rapid recycling. Mean chl \(a\) concentration was 3.9 mg m\(^{-3}\) (0.4 to 12.0 mg m\(^{-3}\)), and satellite images indicated that blooms of this magnitude persisted between December and March in all 3 yr of this study (see Korb et al. 2004 and ocean color and temperature scanner [OCTS] data on the NASA Ocean Color website: http://oceancolor.gsfc.nasa.gov/). Typical primary production rates of 0.8 to 1.8 g C m\(^{-2}\) d\(^{-1}\) (Table 3) would total 96 to 216 g C m\(^{-2}\) 120 d\(^{-1}\) (4 mo). Assuming Redfield uptake ratios, this would equate to 2.5 to 5.7 mmol PO\(_4\)-P and 40 to 91 mmol NO\(_3\)-N over a typical euphotic depth of ~30 m (Korb & Whitehouse 2004). With typical winter concentrations of 2.1 and 30 mmol m\(^{-3}\) for PO\(_4\) and NO\(_3\), respectively (BAS unpubl. data), re-supply of nutrients through, for instance, upwelling or recycling as indicated by NH\(_4\) use (Table 2), must occur.

Another aspect to the N preference of phytoplankton is evidenced by both NO\(_3\) and NH\(_4\) being significant predictors in west off-shelf waters. High NH\(_4\) concentrations and utilisation would usually inhibit nitrate uptake (Armstrong 1999). Given that simultaneous NO\(_3\) and NH\(_4\) use is implausible, perhaps patchiness in Fe availability (facilitating NO\(_3\) use) or grazer distribution (resulting in available NH\(_4\)) enables the alternating use of both N sources. This may be an important contributory factor for higher primary production rates in the west as well as the exceptional longevity of the region’s bloom.

Si(OH)\(_4\) was a highly significant predictor of chl \(a\) in east, on-shelf regions, with lower significant values evident in all other regions, which is due to the usual dominance of diatoms in the system. However, 1997/98 was atypical in 2 respects. For this year, Meredith et al. (2005) described the passage of a cold, large-scale ocean anomaly. Unusually, water density was higher over the island’s shelf relative to off-shelf waters and the net baroclinic flow was cyclonic or eastwards along the northern shelf-break. Cyclonic flow around South Georgia is the opposite direction to the general circulation offshore and would lead to stronger lateral shear and greater barotropic frontal instability (Meredith et al. 2005). This increased potential for the transfer of water between the shelf and the open ocean may have resulted in a greater availability of micronutrients from deep water and/or shelf sediment. In turn, this facilitated more efficient NO\(_3\) utilisation than is usually the case, as indicated by near-Redfield depletion ratios of ~16:1 NO\(_3\):PO\(_4\) and ~1:1 Si(OH)\(_4\):NO\(_3\) (Fig. 3). Secondly, the greatest apparent NO\(_3\) and PO\(_4\) depletions occurred during this season while Si(OH)\(_4\) concentrations remained relatively high. Conditions appear to have favoured non-diatom phytoplankton growth that is atypical for South Georgia (see Korb & Whitehouse 2004).
Top-down control on phytoplankton

Our GLM identified krill as a significant predictor of chl a in the east (Table 2), and our examination of sub-regions (Fig. 5) confirmed that they were negatively related in eastern waters. A negative relationship might be expected if grazing pressure has an important control on phytoplankton. Our model indicated that krill was not a significant predictor of chl a in western waters but did indicate a positive relationship in the west off-shelf sub-region, which was supported by our examination of sub-regions.

If we consider krill ingestion estimates in east off-shelf waters for 1997/98 when krill density was high (mean 260 g m$^{-2}$) and phytoplankton abundance was low (mean 0.8 mg chl a m$^{-3}$), then top-down control on phytoplankton appears inevitable. The 260 g wet mass m$^{-2}$ is equivalent to 52 g dry mass m$^{-2}$ or 20.8 g C m$^{-2}$ (using relationships in Morris et al. 1988 and Schnack 1985). With a C ration of ~5% d$^{-1}$ measured at South Georgia (Pakhomov et al. 1997a, b), krill would consume ~1.0 g C m$^{-2}$ d$^{-1}$. The 0.8 mg chl a m$^{-3}$ is equivalent to 40 mg C m$^{-3}$ (using a C:chl a ratio of 50), which, over a typical 75 m surface mixed layer (Korb & Whitehouse 2004), equals 3 g C m$^{-2}$. So, notwithstanding other grazers, the ingestion by krill alone would remove 33% of phytoplankton per day. Similarly, if we compare krill ingestion to primary production (mean 0.2 g C m$^{-2}$ d$^{-1}$; Table 3), then krill would appear to consume on average 5 times the daily primary production.

The krill daily carbon ration used for this calculation (5%) is not excessive given that their maximum daily ration is probably about 13 to 27% (Clarke et al. 1988, Pakhomov et al. 1997b). An independent grazing rate can be derived from gross growth rates of 1% of krill mass d$^{-1}$ measured for northern South Georgia (Atkinson et al. 2006). Based on a gross growth efficiency of 30% (Straile 1997), the daily ration would be 3.3%. Whether based on a ration of 3.3 or 5%, high concentrations of krill within the east off-shelf region are likely to have a substantial grazing effect on phytoplankton.

Over the South Georgia shelf, krill are also likely to have a substantial grazing effect, especially in the east on-shelf sub-region where their densities can be high. Here, however, our GLM showed a weaker negative relationship between krill density and chl a compared to east off-shelf waters. How dense krill swarms interact with their food sources in shallow shelf regions is poorly known and beyond the scope of the present dataset. However, a possible explanation for our weaker on-shelf GLM relationships is that the krill here had switched to alternative food sources such as copepods or heterotrophic flagellates (Atkinson & Snýder 1997, Pakhomov et al. 1997a, Schmidt et al. 2006). Indeed, in any area with several hundred g of krill under 1 m$^{2}$, our calculated removal of about one-third of phytoplankton stocks per day is clearly unsustainable (Atkinson et al. 2001). Additional mitigating factors in these regions may include decreased daily rations, feeding on competing grazers or on detritus, and high excretion of NH$_4$ to promote plant growth.

South Georgia contains some of the highest krill densities anywhere in the Southern Ocean (Atkinson et al. 2008), but densities can also be very low here. If we consider krill ingestion estimates in west off-shelf waters for 1997/98 when krill density was low (mean 5.1 g m$^{-2}$) and phytoplankton abundance was high (mean 2.2 mg chl a m$^{-3}$), then krill appear to have little grazing effect. The 5.1 g wet mass m$^{-2}$ equates to ~0.41 g C m$^{-2}$ using the same relationships as above. With a C ration of 5% d$^{-1}$, krill would consume 0.02 g C m$^{-2}$ d$^{-1}$. The 2.2 mg chl a m$^{-3}$ over a typical 50 m surface mixed layer (Korb & Whitehouse 2004) equates to ~5.5 g C m$^{-2}$. Therefore, krill ingestion would remove 0.4% of phytoplankton per day. As a proportion of primary production (mean = 1.81 g C m$^{-2}$ d$^{-1}$; Table 3), krill would remove ~1% d$^{-1}$.

Overall perspective

A previous station-based analysis of nutrients, phytoplankton and grazers (Shreeve et al. 2002) concluded that the colder waters to the east of South Georgia contained more krill because of their more recent advection from farther south, while the warmer waters to the west were more favourable for phytoplankton growth. Our larger dataset does not contradict this view but does implicate nutrients as the key factors facilitating phytoplankton growth in the west and suggests a greater interaction between krill and phytoplankton.

Throughout the region, NH$_4$ appears to be a key N source facilitating primary production. However, the addition of NO$_3$ as a significant predictor of chl a in west off-shelf waters may be critical in facilitating the major blooms that occur in this region (Korb et al. 2004), and Fe availability rather than temperature may be the primary control here. The data also suggest a link between NH$_4$ availability and krill distribution; note the concurrent reduction of krill abundance and NH$_4$ concentrations in 1998/99 (Figs. 2 & 4). Experiments suggested that krill excretion in the eastern area could supply 16 to 50% of the NH$_4$ demand here (Atkinson & Whitehouse 2000), while other metazoans (copepods and small euphausiids) would contribute a similar amount (Atkinson & Whitehouse 2001). However, given the likely short residence times and reac-
activity of NH$_4$, these dynamics and feedbacks would be difficult to investigate in a field study such as ours.

While the dynamics of controls on phytoplankton in areas of exceptionally high krill biomass remain speculative, our data allow some generalisations for South Georgia (Fig. 7). The data show a contrasting relationship between krill and phytoplankton in the west offshore region (positive) compared to the rest of the surveyed area (negative). Thus the positive and negative relationships for the various sub-regions (Fig. 5) seem to represent the opposing slopes on either side of this dome. The re-supply of micro- and macronutrients in the west is sufficient to promote high rates of primary production in an area of low krill density, and the krill are located where their food is. This convergence may be due to physical forcing, or perhaps krill are able to maintain their position where grazing opportunities are best. In the east and over parts of the shelf, macronutrient concentrations remain high, primary production rates are low and grazing pressure is implicated.

The actual mechanisms of the krill–food interaction would probably occur at a smaller spatial scale, for example, at the scales of mobile krill swarms that contain actively feeding individuals (Antezana & Ray 1984, Schmidt et al. 2006). However, logic would suggest that the mechanism of the various bottom-up and top-down controls varies greatly in time and space. This would explain the range of positive and negative krill–food relationships reported in the literature at different scales (e.g. Weber et al. 1986, Atkinson et al. 2008). Overall, the relative importance of top-down and bottom-up controls varies both with scale of observation and with local predator densities.

In high latitude environments as elsewhere in the world ocean, protozoans tend to be the main grazers of phytoplankton (Calbet & Landry 2004). This also holds generally in the Southern Ocean, where the grazing effect of metazoans is often found to be small (e.g. Urban-Rich 2001, Dubischar & Bathmann 1997). South Georgia can provide an exception to this rule, where metazooplankton as well as krill biomass is high, and their grazing effect can be significant (Pakhomov et al. 1997a,b, Atkinson et al. 2001). However, at the mesoscale it is likely that the distribution of protozoans and copepods is much more even than that of krill, so their grazing can be envisaged as a ‘background’ on which the krill are superimposed. Krill density varies greatly at the mesoscale and can be measured acoustically, allowing us glimpses of the simultaneous top-down and bottom-up controls on phytoplankton.

Acknowledgements. We thank the officers, crew and scientific parties involved with data collection during cruises JR17, JR28 and JR38. We particularly thank J. Watkins for the collection of krill density data, J. Priddle for chlorophyll analysis, A. Rees for assistance with nutrient analysis and A. Wood for help with data analysis. In addition, we acknowledge 4 anonymous referees whose constructive comments enabled us to improve our analysis.

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Submitted: December 5, 2008; Accepted: August 27, 2009
Proofs received from author(s): October 14, 2009