FEATURE ARTICLE

Overwinter habitat selection by Antarctic krill under varying sea-ice conditions: implications for top predators and fishery management

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ABSTRACT: Climate change will affect Antarctic krill Euphausia superba, krill-dependent predators, and fisheries in the Southern Ocean as areas typically covered by sea ice become ice-free in some winters. Research cruises conducted around the South Shetland Islands of the Antarctic Peninsula during winters with contrasting ice conditions provide the first acoustic estimates of krill biomass, habitat use, and association with top predators to examine potential interactions with the krill fishery. Krill abundance was very low in offshore waters during all winters. In Bransfield Strait, median krill abundance was an order of magnitude higher (8 krill m⁻²) compared to summer (0.25 krill m⁻²), and this pattern was observed in all winters regardless of ice cover. Acoustic estimates of krill biomass were also an order of magnitude higher (~5 500 000 metric tons [t] in 2014) than a 15 yr summer average (520 000 t). Looking at krill-dependent predators, during winter, crabeater seals Lobodon carcinophagus were concentrated in Bransfield Strait where ice provided habitat, while Antarctic fur seals Arctocephalus gazella were more broadly distributed. Krill overwinter in coastal basin environments independent of ice and primary production and in an area that is becoming more frequently ice-free. While long-term projections of climate change have focused on changing krill habitat and productivity declines, more immediate impacts of ongoing climate change include increased risks of negative fishery–krill–predator interactions, alteration of upper trophic level community structure, and changes in the pelagic ecology of this system. Development of management strategies to mitigate the increased risk to krill populations and their dependent predators over management timescales will be necessary to minimize the impacts of long-term climate change.

KEY WORDS: Southern Ocean · Antarctic krill · Fishery interactions · Sea ice · Commission for the Conservation of Antarctic Marine Living Resources · CCAMLR · Ocean warming

Changes in sea-ice coverage will impact trophic interactions during the Antarctic winter.

Photos: Jennifer Walsh and Abraham Borker

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INTRODUCTION

Antarctic krill *Euphausia superba* is a key species in the Southern Ocean and is a principal link between phytoplankton production and upper trophic levels (Atkinson et al. 2008). They are important for nutrient cycling and carbon flux through the ecosystem (Laws 1977, Smetacek & Nicol 2005), and their massive biomass supports an important commercial fishery (Nicol et al. 2012, Nicol & Foster 2016). A number of studies have shown a strong relationship between seasonal sea-ice extent and recruitment success of krill, emphasizing the importance of physical processes in winter (Loeb et al. 1997, Atkinson et al. 2004, Ross et al. 2014, Saba et al. 2014). In the last quarter of the 20th century, krill densities across the Southwest Atlantic sector of the Southern Ocean, near the Antarctic Peninsula, have reportedly declined between 36 and 80%, concomitant with declines in sea-ice extent and duration (Atkinson et al. 2004). The decline in sea ice, driven by climatic changes in atmospheric heat transport, has presumably led to concomitant changes in quality and quantity of over-winter habitat for krill at all life stages (but see Melbourne-Thomas et al. 2016). Such changes in the physical environment are likely to continue in the future and result in an increased number of low-ice and ice-free years (Stammerjohn et al. 2008b). This may have negative impacts on the food web (Flores et al. 2012a,b) and ice-dependent krill predators (Siniff et al. 2008, Dunn et al. 2011, Santora 2014), and could increase fishery interactions (Nicol et al. 2012).

Along the Antarctic Peninsula, krill are distributed on the shelf and into oceanic waters during summer, where larger krill spawn (Siegel et al. 2013) and where the fishery historically exploited large temporally stable aggregations (Nicol & Foster 2016). At-sea observations and surveys have related patterns of habitat use by birds and mammals to the distribution of krill hotspots and krill size distribution (Santora et al. 2012, Santora & Veit 2013), while studies focused on land-based predators have examined summer reproductive success of birds and mammals, including Antarctic fur seals *Arctocephalus gazella* (Croll & Tershy 1998) and penguins (Trivelpiece et al. 2011), in relation to krill abundance and demographic patterns. These studies have shown that the spatial distribution of krill during summer is a major factor influencing the distribution of predators and thus the spatial structure of ecological interactions.

There are considerable gaps in understanding how the pelagic ecosystem of the Southern Ocean is restructured from summer to winter, especially now that climate change is impacting the initialization of ice formation and duration (Stammerjohn et al. 2008b). The seasonal abandonment by krill of off-shore waters (Siegel 1988), where 90% of the krill biomass reside during austral summer (Atkinson et al. 2009), and the hypothesized shelfward migration during autumn and winter (Mackintosh 1972, Siegel 1988), suggest that krill density should increase greatly in coastal and near-shore environments during winter. However, contradictory evidence exists for the magnitude of the concentration of krill in coastal areas (Mackintosh 1972, Stepnik 1982, Siegel 1988, 2005, Zhou et al. 1994, Lascara et al. 1999, Zhou & Dorland 2004, Lawson et al. 2008a,b, Atkinson et al. 2009, Nowacek et al. 2011), with different results based on timing, sampling gear, and conditions. Additionally, recent effort has focused on the importance of deep-ocean (>1500 m) benthic environments as overwintering habitats (Atkinson et al. 2009, Schmidt et al. 2011) that might harbor a large fraction of krill during winter. Clearly distinguishing the magnitude of coastal concentration is necessary to understand krill population dynamics and life history tradeoffs, and also to understand availability of krill for krill-dependent predators (Ribic et al. 2008) as well as the fishery.

There is a critical need to better understand the winter structure of these ecologically-important and climatically-vulnerable ecosystems, especially given the observed declines in sea-ice extent and duration (Stammerjohn et al. 2008a), estimated declines in krill abundance (Atkinson et al. 2004), increased localized fishing pressure during autumn and winter (Nicol & Foster 2016), and potential interaction with non-monitored species. Here we use a multidisciplinary data set from 3 winter surveys to examine the relationships among the spatial distribution of predators, the abundance and biomass of krill, sea ice, and primary production around the northern Antarctic Peninsula. Specifically, we test whether: (1) the relative importance to krill of areas in the Antarctic Peninsula varies from summer to winter (i.e. seasonal migration hypothesis), and (2) krill-predator habitat use is dependent on sea-ice concentration and krill biomass. We provide the first-ever winter acoustic estimates of krill biomass in this region during a relatively ice-free season using the latest algorithms (CCAMLR 2010, Fielding et al. 2011) and compare these estimates with historical data collected during summer.
MATERIALS AND METHODS

Study area

The northern Antarctic Peninsula region of the Southern Ocean is a bathymetrically-complex area with a variety of water currents that converge and mix around the South Shetland Islands, the tip of the peninsula at Joinville Island (JI), and within Bранs-field Strait (BS; our Fig. 1; Jiang et al. 2013). In the BS, cold salty Weddell-shelf water enters near JI and flows south along the northern coast of the Antarctic Peninsula (von Gyldenfeldt et al. 2002, Thompson et al. 2009, Thompson & Youngs 2013). Water from the Antarctic Circumpolar Current (ACC; Bellingshausen surface water, and Upper Circumpolar Deep Water) enters the BS from the southwest at the surface and at mid-depths, and flows northeast along the southern coast of the South Shetland Islands (Gordon & Nowlin 1978, Gordon et al. 2000). Additional inflows of coastal water from the western Antarctic Peninsula through Gerlache Strait (GS) can enter into the BS from the southwest. The mixing and advection of these waters in the BS, coupled with the steep bathymetry and deep basins, create a cyclonic circulation within the BS that has a strong coastal boundary current on the north side and eddies over the basins (Zhou et al. 2002, 2006).

We adapted a survey design employed during austral summer to study the distribution and abundance of krill and predators during the austral winter, thereby providing direct comparisons of the spatial distribution, abundance, and biomass of krill over approximately 125 000 km² of the Southern Ocean. From 1996 to 2011, the US Antarctic Marine Living Resources (AMLR) Program annually conducted summer surveys (January–March) around the South Shetland Islands and sampled from a grid of up to 110 fixed stations allocated at 15–20 nautical mile (nmi) intervals once or twice each summer (Reiss et al. 2008). The survey area is divided into 4 strata: the west shelf stratum (WA), north of Livingston and King George Islands; the BS stratum, between the South Shetland Islands and the Antarctic Peninsula; the Elephant Island stratum (EI), an oceanographically-dynamic area and historic center of the summer krill fishery; and the coastal waters at the tip of the Antarctic Peninsula near JI.

Winter surveys were conducted aboard the research vessel/icebreaker (RVIB) ‘Nathaniel B. Palmer’ in August through September, 2012–2014. Winter stations were selected from the historic survey grid based on the amount of time available during the cruise and ice conditions (Fig. 1). During winter 2012 (August 1–17), a smaller number of stations was sampled owing to limited ship time. In that year, 37 stations were sampled mostly in the EI stratum, with some samples in the BS and JI strata. No samples were collected in the WA stratum. In 2013 (August 9 to September 8) and 2014 (August 19 to September 18), 88 and 114 stations were sampled throughout the South Shetland Islands area, respectively. During the 2 years (2012 and 2013) when ice was extensive, we sampled as far as 55 km into the ice in the JI and BS strata. In ice-free areas offshore, we sampled most of the US AMLR survey stations during 2013, and in 2014, the lack of significant pack ice enabled us to sample the entire survey grid.

Sea-ice concentration data

Monthly sea-ice concentration data for June, July, and August of each year were extracted from the National Snow and Ice Data Center website (www.nsidc.org), and the sea-ice area (km²) within the 15 and 50% ice concentration isopleths in each month was calculated for the area between 58° S and 48° W, and 68° S and 65° W (see Figs. 2 & 3).

Integrated chlorophyll a determination

At each station, a conductivity, temperature, and depth (CTD) cast was made to within 10 m of the bottom or 750 m depth. The CTD, an SBE9/11 (SBE Inc.), was equipped with 10 l bottles for water sampling, and bottles were closed on the upcast at 750, 200, 100, 75, 50, 40, 30, 20, 15, and 5 m. Chlorophyll a (chl a) concentrations were determined fluorometrically following Holm-Hansen et al. (1965). Integrated chl a (to 100 m depth; mg chl a m⁻²) was calculated for each station (Reiss et al. 2009) and averaged for summer (15 yr) and winter (3 yr) surveys by survey stratum (EI, JI, BS, WA).

Zooplankton sampling

Krill and zooplankton were sampled using a 1.8 m (2.54 m² mouth area) Isaacs-Kidd midwater trawl (IKMT) fitted with a 505 µm mesh net. Volume filtered during trawls was determined using a calibrated General Oceanics flow meter (model 2030R) mounted on the depressor frame in front of the net.
All tows were double-oblique to 170 m or to ca. 10 m above bottom in shallower waters. Real-time tow depths were derived from a pressure sensor mounted on the trawl bridle. Tow speeds were ~2 knots, with volumes filtered averaging ~3621 m³ (SE 21.16). Zooplankton abundance, including that of krill, was standardized to no. m⁻² based on volume of water filtered multiplied by the depth of the tow.

**Kril measurement**

Immediately after each IKMT trawl, all zooplankton were sorted from the fresh samples. Krill were removed first, and a subsample of up to 100 post-larval krill was measured. Total body length (mm) was determined by measuring the distance from the tip of the rostrum to the posterior tip of the uropods (Standard 1 as described by Mauchline 1980). Length–frequency distributions for krill were derived for each stratum and season surveyed and used to convert the acoustic data into biomass.

**Acoustic data and biomass determination**

Acoustic data were collected in all years; however, acoustic data were only useful for estimating the biomass of krill in 2014, given the low ice concentrations and preponderance of open water in that year. A Simrad EK60 echosounder was used to collect acoustic data at 3 frequencies (38, 120, and 200 kHz). All 3 frequencies were calibrated using the standard sphere technique (Foote 1990).

Krill were delineated from other acoustic scatterers using the 3-frequency stochastic distorted-wave Born approximation (SDWBA) model, using 95% of the total krill length–frequencies (measured from net tows) for each stratum to define dB-difference windows (Demer & Conti 2005, Reiss et al. 2008, CCAMLR 2010). Only daytime acoustic data were used for biomass estimation to minimize potential bias caused by the diel vertical migration of krill (Demer & Hewitt 1995). Acoustic data were processed using Echoview (ver. 4.9), with additional manual removal of noise created by ice scraping the hull. Data were integrated over the upper 250 m of the water column and into 1 nmi bins. In general, where krill targets could be identified within the echogram but could not be isolated from the surrounding noise, those noisy data were excluded from the integration, thereby minimizing potential bias of including noise, but potentially biasing our biomass estimates downwards. Similarly, as we integrated the acoustic energy from 250 m to the surface, this approach would also exclude any biomass that was present deeper in the water column.

**Predator and sea-ice surveys**

Continuous observations of seabirds and marine mammals were conducted from the bridge (Santora 2014). Observations were collected simultaneously by 2 observers during daylight hours; one observer recorded all seabirds, while the other observer used 20 × 60 binoculars (20× magnification, 60 mm objective lens diameter) to scan for marine mammals. As ship survey speed varied according to sea-ice concentration and the icebreaker followed an irregular path through ice leads, predator observations and environmental data were binned into 1 nmi intervals. In 2012, the sea ice was monitored during daylight hours, and scored on the percent cover, noting the
general type of ice present. In 2013 and 2014, sea ice was continuously monitored during predator surveys and was classified according to a standardized sea-ice classification protocol (Scientific Committee on Antarctic Research [SCAR] Antarctic Sea Ice Processes and Climate [ASPeCt] program; Worby et al. 1999). Sea-ice conditions were logged automatically every 15 s and included estimates of percent ice coverage (0–10 scale, with 0 being open water and 10 being fully consolidated ice with no leads) and ice type (e.g. 1st and 2nd year, nilas, thin grey ice) (Worby et al. 1999, Worby & Comiso 2004).

**Statistical analysis**

We examined stratum- and season-specific variability in krill abundance from net tows to assess if regional krill abundance varied from summer to winter. Our focus was on broad changes in the structure of the pelagic ecosystem and changes in krill and predator distribution patterns between summer and winter, not on the inter-annual variability in summer or winter abundance and distributions. We averaged krill abundance from net tows over the 15 summer surveys (1996–2011) and also for 3 winter surveys (2012–2014) among strata and used 2-way ANOVA to compare krill abundance and water-column phytoplankton biomass among strata (EI, WA, JI, SA) and between seasons (summer and winter). We tested the data for normality and applied various transformations. In general, however, all data from each factor could not be normalized using a single transformation. So we ln(x + 1)-transformed the data to reduce the variances between groups and use the 2-way ANOVA acknowledging these limitations. We used Tukey-Kramer multiple comparisons to examine interactions and main effects. Alpha level = 0.05.

Although summer and winter surveys were not consecutive, we assume the overall spatial pattern of krill abundance during past summers is comparable to the summers between winter surveys. Indeed, recent work (Siegel 2013) has shown that in the summer of 2013, krill distributions around the South Shetland Islands were similar to the pattern of distribution observed for the summer periods between 1996 and 2011, indicating that historical summer distributions are likely representative of patterns during the study period.

We used generalized additive models (GAMs) to assess the relationships between pinnipeds, sea-ice concentration, and integrated krill biomass. However, acoustic estimates of krill biomass were only available in 2014, so models were limited to this low-ice year. Counts of pinnipeds (crabeater seals *Lobodon carcinophagus* and Antarctic fur seals *Arctocephalus gazella*), estimates of sea-ice concentration (percentage), and krill biomass were grouped into 1 nmi intervals (n = 629). The GAMs were specified as: Seal (count per nmi) = s(sea-ice concentration) + s(krill biomass) + te(Lon, Lat), where s is a smooth regression spline and te is a smoothed spatial interaction term between longitude and latitude. GAMs were implemented using the mgcv package in the R statistical program (R Development Core Team 2016), and we used generalized cross-validation to estimate smoothness parameters (Zuur et al. 2009). Adjusted pseudo-R² and percent deviance explained were used to evaluate model performance.

**RESULTS**

**Sea-ice extent and distribution**

Sea-ice coverage in the study area varied among years and over the seasons (Figs. 2 & 3). In 2012, the area within the 15% concentration isopleth was 116 132 km² in June, increased slightly (133 377 km²) in July, and then increased to more than 195 000 km² in August. The area within the 50% sea-ice concentration isopleth ranged from 64 to 69% of that within the 15% sea-ice concentration isopleth during June and July 2012, declining to 47% by August as the ice extent continued to expand across the areas. In 2013, the 15% sea-ice area increased linearly from June to August, from 97 310 to 341 257 km², respectively. At the same time, the 50% sea-ice area increased from 68 368 to 226 913 km². In 2014, sea-ice extent was extremely variable; the area within the 15% sea-ice concentration isopleth declined between June and July 2012, declining to 47% by August as the ice extent continued to expand across the areas. In 2013, the 15% sea-ice area increased linearly from June to August, from 97 310 to 341 257 km², respectively. At the same time, the 50% sea-ice area increased from 68 368 to 226 913 km². In 2014, sea-ice extent was extremely variable; the area within the 15% sea-ice concentration isopleth declined between June and July, from 102 027 to 100 158 km² in July, then increased to 218 557 km² in August. The area encompassed by the 50% sea-ice concentration isopleth followed a different trend. In June 2014, the 50% sea-ice concentration isopleth covered 70% of the area of the 15% isopleth. The 50% isopleth declined to 34% of the 15% sea-ice concentration area in July and August, indicating that most of the spatial coverage of ice consisted of loose and unconsolidated ice.

**Seasonal variability in water column productivity**

Depth-integrated chl a biomass varied significantly between summer and winter and among survey
strata but showed a similar pattern among winters (Fig. 4a). In winter, integrated chl a ranged from a low of 5.4 mg m$^{-2}$ in BS during 2012 to a high of 18.4 mg m$^{-2}$ in EI and WA during 2014. In general, the EI and WA had the highest integrated chl a biomass in winter, while BS and JI had consistently lower integrated chl a biomass. In summer, over the 15 yr time-series, the patterns were reversed; EI and WA strata had the lowest average chl a (54.85 and 41.8 mg m$^{-2}$, respectively), while the JI and BS strata had higher integrated chl a biomass (58.9 and 68.4 mg m$^{-2}$, respectively) (Fig. 4a). Overall, integrated chl a biomass was 4.7 times greater in summer compared to winter. Results of a 2-way ANOVA (season and stratum as factors) showed a significant inter-
action between season and stratum \((F_{(3,2451)} = 14.8, p < 0.0001\)\), and significant difference between seasons \((F_{(1,2451)} = 952.3, p < 0.00001\)\) and also strata \((F_{(3,2451)} = 5.01, p < 0.005\)\). Therefore, while phytoplankton biomass was highest during summer as expected, the interaction indicates that the spatial distribution of chl \(a\) biomass differs between seasons, with relatively higher chl \(a\) biomass offshore within the ACC (EI and WA) strata and lower within the coastal BS and JI strata during winter.

### Seasonal changes in net-based krill abundance and distribution

The spatial distribution and abundance of krill also changed between summer and winter (Table 1, Fig. 4b). Median abundances in the WA and EI strata during summer ranged between 0.32 and 0.77 krill m\(^{-2}\), respectively. During winter, median krill abundance in these 2 strata declined by an order of magnitude to 0.04 krill m\(^{-2}\) in the EI stratum during both winters, and to 0.05 krill m\(^{-2}\) in the WA stratum in 2014. In the JI stratum, median krill abundance during summer was 0.3 krill m\(^{-2}\), and ranged between 0.33 and 1.81 krill m\(^{-2}\) in winter 2013 and 2014, respectively. For the BS stratum during summer, median krill abundance was 0.25 krill m\(^{-2}\), and increased more than an order of magnitude to a median abundance of 8.07 krill m\(^{-2}\) during winter.

Results of a 2-way ANOVA (season and stratum as factors) on ln\((x + 1)\) krill abundance showed that krill were significantly more abundant in winter compared to summer (Fig. 4b; ANOVA, \(F_{(3,2648)} = 44.8, p < 0.0001\)).

Significant interaction between season and stratum \((F_{(3,2648)} = 44.8, p < 0.0001)\) further emphasizes the magnitude of the seasonal change in distribution from offshore spawning areas (WA and EI) during summer to coastal habitats in the JI and BS strata during winter.

### Distribution of acoustic krill biomass

Acoustic estimates of krill biomass during summer among the 4 strata ranged almost an order of magnitude and showed considerable seasonal variability (Table 2, Fig. 5). Mean biomass density of krill during winter 2014 in the BS was 228 g m\(^{-2}\), an order of magnitude greater than the average biomass density for this stratum during summer. Additionally, this high biomass density was between 4 and 12 times greater than the biomass densities in the other strata during summer. Within the BS stratum, krill biomass ranged from 14,695 metric tons (t) in summer 2005 to more than 1.6 million t in summer 2011, and averaged 590,000 t. In contrast, summer biomass in the WA and EI strata averaged 1.87 and 2.19 million t, respectively.

**Table 1.** Mean (±SD) and median Antarctic krill abundance (no. m\(^{-2}\)) from net tows across 4 sampling strata around Antarctic Peninsula between 15 summers and 3 winters. N: number of years surveyed between 1996 and 2014 (not all areas were sampled in all years); BS: Bransfield Strait; EI: Elephant Island; JI: Joinville Island; WA: west shelf

<table>
<thead>
<tr>
<th>Stratum</th>
<th>N</th>
<th>No. of tows</th>
<th>Summer</th>
<th>Winter</th>
</tr>
</thead>
<tbody>
<tr>
<td>EI</td>
<td>15</td>
<td>1273</td>
<td>8.7 (57.25)</td>
<td>0.77</td>
</tr>
<tr>
<td>JI</td>
<td>15</td>
<td>188</td>
<td>12.78 (34.88)</td>
<td>0.30</td>
</tr>
<tr>
<td>BS</td>
<td>15</td>
<td>387</td>
<td>19.85 (92.38)</td>
<td>0.25</td>
</tr>
<tr>
<td>WA</td>
<td>15</td>
<td>561</td>
<td>5.75 (45.75)</td>
<td>0.32</td>
</tr>
</tbody>
</table>

**Table 2.** Mean biomass (millions of tons) and mean density (g m\(^{-2}\)) of Antarctic krill in 3 strata around South Shetland Islands. Winter acoustic biomass estimates are from the single acoustic survey conducted in austral winter 2014 in the same strata. Coefficient of variation (CV) of survey biomass calculated using the Jolly & Hampton (1990) method. N: number of years surveyed between 1996 and 2011 (not all areas were sampled in all years); BS: Bransfield Strait; EI: Elephant Island; WA: west shelf

<table>
<thead>
<tr>
<th>Stratum</th>
<th>Summer</th>
<th>Winter</th>
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<tbody>
<tr>
<td></td>
<td>Mean (CV)</td>
<td>Mean krill abundance</td>
</tr>
<tr>
<td>EI</td>
<td>15</td>
<td>2.19 (27.5)</td>
</tr>
<tr>
<td>WA</td>
<td>15</td>
<td>1.87 (49.5)</td>
</tr>
<tr>
<td>BS</td>
<td>15</td>
<td>0.47 (37.8)</td>
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</table>
During winter 2014, acoustic estimates of krill biomass in the WA and EI strata were much lower than summer estimates (200 000 and 76 000 t, respectively). The acoustic estimate of krill biomass in the BS stratum during winter 2014 was 5.5 million t, about 3.5 times the maximum summer biomass, and nearly an order of magnitude greater than the average summer biomass recorded for this stratum (Table 2, Fig. 5). In fact, the biomass in BS (20% of the survey area) represented about 92% of the average total summer biomass for the entire US AMLR study area, and 8% of the standing stock of the southwestern Atlantic estimated during summer 2000 (Hewitt et al. 2004, CCAMLR 2010).

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**Pinniped distribution and relationship to sea ice and krill biomass**

The spatial distributions and relative abundances of crabeater and Antarctic fur seals further emphasize the importance of the seasonal shift in krill distribution into the BS during winter (Fig. 6). Regardless of year, crabeater seals were almost entirely associated with high krill biomass areas within BS during winter (Figs. 5 & 6a,b). Crabeater seals were virtually absent from areas with low sea-ice concentrations like the WA or EI strata, but were also absent from the JI stratum where first-year ice occurred but krill biomass was low. The relative abundance of crabeater seals peaked when sea-ice concentration was greater than 70% within BS (Fig. 6c,d). In comparison, both years, Antarctic fur seals (Figs. 5 & 6a,b) were more broadly distributed than crabeater seals, and occurred throughout the survey area, including in areas with low to moderate sea-ice concentrations (e.g. 20 to 40%; Fig. 6c,d). Both crabeater and Antarctic fur seals were more abundant in 2014 than 2013 (Mann-Whitney U, p < 0.001), with larger aggregations of animals (repeated sightings of 150 to 400 ind. nmi⁻¹) concentrated on the lower number of ice floes suitable for hauling out in 2014. The GAMs performed better for
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... which we attributed to the broader distribution of fur seals across a variety of sea-ice habitats (Table 3, Figs. 6 & 7). GAMs indicated that crabeater and fur seals were positively related to sea-ice concentration and krill biomass during 2014, but displayed different functional relationships (Fig. 7). The GAM for crabeater seal abundance showed a threshold response to sea-ice concentration, and high abundance of crabeater seals was always associated with high krill biomass. In contrast, the GAM for fur seals indicates they used a variety of sea-ice concentrations, and their abundance increased monotonically with krill biomass.

Table 3. Results of generalized additive models for assessing spatial distribution pattern and response of crabeater seals and Antarctic fur seals to sea-ice concentration and Antarctic krill biomass; \( te(\text{Lon}, \text{Lat}) \) is spatial component that includes a smoothed interaction term between longitude and latitude. Edf: estimated degrees of freedom, Res.df: residual degrees of freedom, %Dev: percent deviance explained. \( R^2 = \text{adjusted pseudo-R}^2 \)

<table>
<thead>
<tr>
<th></th>
<th>Edf</th>
<th>Res.df</th>
<th>( F )</th>
<th>p</th>
<th>%Dev</th>
<th>( R^2 )</th>
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<tr>
<td><strong>Crabeater seal</strong></td>
<td></td>
<td></td>
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<tr>
<td>Sea ice</td>
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<td>2.81</td>
<td>16.45</td>
<td>&lt;0.0001</td>
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<td>67.26</td>
<td>&lt;0.0001</td>
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<tr>
<td>( te(\text{Lon}, \text{Lat}) )</td>
<td>13.94</td>
<td>15.06</td>
<td>7.35</td>
<td>&lt;0.0001</td>
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<tr>
<td><strong>Antarctic fur seal</strong></td>
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<tr>
<td>Sea ice</td>
<td>2.75</td>
<td>2.95</td>
<td>12.50</td>
<td>&lt;0.0001</td>
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<td>0.31</td>
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<tr>
<td>Krill biomass</td>
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<td>2.56</td>
<td>28.75</td>
<td>&lt;0.0001</td>
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<tr>
<td>( te(\text{Lon}, \text{Lat}) )</td>
<td>22.08</td>
<td>22.79</td>
<td>17.34</td>
<td>&lt;0.0001</td>
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Fig. 6. Relative abundance (ind. nmi\(^{-1}\)) of crabeater seals (CRSE; red circles) and Antarctic fur seals (FUSE; blue circles) in the study area during winter (a) 2013 and (b) 2014. Note difference in scales. Average relative abundance (ind. nmi\(^{-1}\)) \( \pm 1 \) SD of crabeater (red bars) and Antarctic fur seals (blue bars) by ice concentration (%) during daytime visual surveys along acoustic transect during (c) 2013 and (d) 2014.
DISCUSSION

Krill biomass in winter and summer

There is considerable discussion in the literature (Siegel 1988, Lascara et al. 1999, Siegel et al. 2013, Cleary et al. 2016) regarding the importance of the seasonal shoreward migration of krill to coastal waters. Yet, there is less information on the magnitude of this difference and the resulting changes in ecosystem structure (Atkinson et al. 2008). Our data show that the coastal waters of the BS have been an important overwinter habitat for krill within the northern Antarctic Peninsula ecosystem. Krill biomass in the BS increased by more than an order of magnitude between summer and winter, while offshore areas have a similar biomass decline. These findings support the seasonal shoreward migration hypothesis (Siegel 1988, 1989). This order of magnitude seasonal change in distribution and biomass occurs regardless of the concurrent sea-ice conditions (i.e. concentration and area). This spatial re-organization in winter krill biomass has major consequences for understanding the ecology of krill, inferring the impacts of climate change on krill habitat during winter, trophic interactions, and fisheries management.

The use of acoustics in winter is often limited by the presence of ice. As the ship moves through even thin ice, the noise of the ice against the hull contaminates the acoustic returns that might be attributed to krill and other scatterers. This was the case in 2012 and 2013, when the ship routinely broke solid ice, obviating the ability to derive areal estimates of krill biomass in most areas in those years. However, estimates of krill biomass in ice-free areas in both 2012 and 2013 were also very low (~1 g m⁻²; US AMLR Program unpubl. data). In contrast, during 2014, the ice concentration and distribution were sufficiently low that the entire northern Antarctic Peninsula was sampled, and high-quality acoustic data were collected throughout (Fig. 6). The lack of ice meant that there was little physical habitat for the development of ice-algal communities that are often hypothesized to be critical to overwinter survival (Meyer 2012). This observation is strong evidence that the concentration of krill post-larvae in the BS was independent of sea ice or its algal community in 2014.

In contrast to the noise that limits the use of acoustics to largely ice-free areas and winters, nets were deployed regardless of the ice concentration and estimates of krill abundance from net tows were made in each stratum in all years. The ability to tow the nets in the wake of the ship regardless of ice thickness also meant that net-based abundance estimates of krill could be used to corroborate the overall pattern of krill abundance and distribution in the high- (2012 and 2013) and low- (2014) ice years. Over the 3 winters, net-based krill densities were more than an order of magnitude higher within the BS and JI strata compared to summer and showed that krill were much less abundant in WA and EI strata (Fig. 4b) during winter compared to summer. Although net-based krill densities are often underestimates of the true density of krill in a sample, owing to net avoidance and vertical migration of krill below the maximum tow depth during daytime (a significantly reduced problem in winter when daylight is just 6–8 h compared to summer), the overall spatial patterns of relative krill abundance and biomass are

![Graphs showing results of generalized additive models for assessing functional relationships between krill biomass and sea-ice concentration.](image-url)
that over 5 million t were present suggests that our times (US AMLR Program unpubl. data). Yet, the fact were observed on the bottom as deep as 500 m at 2011, 2014). Indeed, krill were observed to vertically feeding are all likely possibilities (Schmidt et al. 2012). Benthic feeding, cannibalism, and omnivorous 2014) despite the need to feed during winter (Meyer 2012). Furthermore, krill post-larvae cannot survive winter without feeding (Meyer 2012), and while post-larvae may not require the sea ice to avoid advective loss, the productivity of sea-ice communities may enhance krill survival and future production, as winter conditions can impact krill body condition the following summer (Reiss et al. 2014). Yet, we found little evidence that during winter, krill were concentrated in areas of higher primary production (e.g. offshore in the ACC, or in the WA stratum) where chl a biomass was high. Krill were also not constrained only to areas with high sea-ice concentrations. Instead, krill were concentrated in the BS regardless of the physical or biological conditions we observed. The flexibility of krill diets may explain this apparent independence (Schmidt et al. 2014) despite the need to feed during winter (Meyer 2012). Benthic feeding, cannibalism, and omnivorous feeding are all likely possibilities (Schmidt et al. 2011, 2014). Indeed, krill were observed to vertically migrate within the BS between night and day, and were observed on the bottom as deep as 500 m at times (US AMLR Program unpubl. data). Yet, the fact that over 5 million t were present suggests that our understanding of krill overwinter survival strategies is still very limited.

In the BS, the order of magnitude increase in biomass observed in winter must have resulted from active horizontal migration from offshore feeding and spawning areas occupied during summer, rather than by passive transport by currents from those areas. This is because the circulation patterns are largely controlled by the bathymetric contours of the region, and seasonal differences in the strength of currents are not substantially different (Savidge & Amft 2009, Jiang et al. 2013). It is known that hydrographic features can affect the distribution of krill at macro- (1000s km; Nicol et al. 2000) and meso-scales (10s to 100s km; Allen et al. 2001, Simard & Lavoie 1999, Santora et al. 2012), and can result in consistently high krill concentrations if krill behavior also contributes. At fine scales, behavior, such as diel vertical migration, is also known to exert strong control on the formation of krill aggregations (Dorman et al. 2015), and it is generally agreed that aggregation and accumulation of krill and zooplankton may result from interactions among bathymetry, circulation, and behavior (e.g. diel vertical migration, swarming, and the need to continually swim) (Simard & Lavoie 1999, Allen et al. 2001, Cotté & Simard 2005, Santora & Reiss 2011). However, it is clear from the magnitude of the biomass accumulation in the BS that krill behavior (which can change seasonally) is responsible for the accumulation of krill in this region. While sea ice is often regarded as an important component of krill habitat, less emphasis has been placed on understanding other physical features of the environment that could provide organizational cues for krill aggregating over winter.

Acoustic biomass densities observed within the BS during winter 2014 averaged 228 g m⁻², much higher than any biomass density observed during summer in this region (Cossio & Reiss 2007, Reiss et al. 2008), and higher than elsewhere along the peninsula where acoustic measurements have been made during autumn or winter. For example, Lawson et al. (2008a,b) reported that during the Southern Ocean Global Ocean Ecosystem Dynamics (GLOBEC) program, acoustic biomass ranged from 1.3 to 77 g m⁻² and was greatest below 100 m in Marguerite Bay during autumn and winter 2001 and 2002, and that krill were also found on the bottom (~500 m). This pattern of acoustic biomass density (higher concentrations nearshore) indicates that Marguerite Bay is an important coastal overwintering area. In the BS, krill density was concentrated in the upper 200 m, although krill were visible on the 38 and 120 kHz echosounders down to 500 m.

Our acoustic estimates of krill biomass are likely underestimates because the signal to noise ratio of the 200 kHz echosounder limits the integration depth to 250 m. Despite the potential limitation of integrating over just 250 m, our estimates of krill biomass are substantially higher than the biomass observed in Marguerite Bay, and the krill were concentrated in the water column, further highlighting the importance of the basins in the BS in winter. Overall, it is clear that other areas (e.g. Marguerite Bay, and the straits and passages around islands along the peninsula) with similar hydrographic features that might concentrate krill near the coast during winter are important and, with decreasing sea ice due to climate change, such areas may become accessible and subject to higher exploitation rates by the krill fishery in the future.
Sea ice, krill, and predators

Climate-induced variability in sea-ice extent, duration, and composition (size and floe types) will impact upper trophic level predators in polar ecosystems (Siniff et al. 2008, Forcada et al. 2012). For example, sea ice provides important foraging habitat for krill-dependent predators, and is required by some predators to haul out and reproduce (Ribic et al. 1991, Siniff et al. 2008). Our data suggest that crabeater seal habitat is associated with sea-ice concentrations greater than 70%, while Antarctic fur seals were associated with a broader range of sea-ice conditions. However, both species were clearly associated with areas of increased krill biomass, indicating they are closely linked to the krill biomass during winter. Crabeater seals are ice-dependent and must therefore find areas of reliable prey concentration associated with pack ice or in the vicinity of polynyas and other oceanographic features that concentrate krill and predators alike (Burns et al. 2004, 2008). The consistent overlap between crabeater seals and krill biomass within coastal waters of the BS indicates that further research with respect to the sensitivity of crabeater seals to changes in winter sea-ice conditions and potential interactions with the krill fishery is warranted (Siniff et al. 2008).

The at-sea abundance of Antarctic fur seals in our study area increases by an order of magnitude from summer to early autumn (Santora 2013), and the high abundances reported in the present study indicate this region is an important overwintering area (Santora 2014). Likewise, the relatively high abundance of Antarctic fur seals, composed of sub-adult and adult males (J. Santora pers. obs.) within the BS, indicates this species is using the same krill biomass hotspots as crabeater seals. Estimates of krill consumption by Antarctic fur seals rely on summer monitoring data (Hill et al. 2007), yet the large numbers of these predators and concentrated food resources in the BS during winter suggest that this area may be important for understanding trophodynamics and population-wide consumption estimates.

Future declines in winter sea-ice conditions may impact ice-dependent seals, especially if sea-ice extent, duration, or concentrations decrease in areas where krill biomass is concentrated. Our observations indicate a significant increase in both species during 2014 when sea ice was less prevalent and floes were smaller, suggesting that haul-out habitat in krill-rich areas could be limited. Additionally, there is a potential for increased predation by killer whales *Orcinus orca* on crabeater seals during these low-ice years as killer whales gain greater access to areas where flows may be smaller, potentially making hunting easier. Moreover, in years with less sea ice, the potential interaction between the krill fishery and seals may be intensified as the seals are constrained to smaller or more fragmented ice habitat within areas where the fishery may focus future effort (Nicol & Foster 2016).

Climate and management implications

The response of the Southern Ocean to global climate change and warming is projected to result in changes throughout the physical and biological components of the ecosystem (Constable et al. 2014, Gutt et al. 2015). The Intergovernmental Panel on Climate Change (IPCC) projects that physical changes owing to increasing water and air temperatures over the next 85 yr (IPCC 2007) will continue to accelerate and broaden changes in ecosystem structure and function (Gutt et al. 2015). Thus, there is considerable interest in projecting the impacts of climate change over the long term. For example, habitat models built from climate change projections indicate shifts in spatial habitat for krill by 2100 because of changing water temperature and primary production (Hill et al. 2013), changes in larval production (Piñones & Fedorov 2016), impacts on larval survival as pH declines (Kawaguchi et al. 2013), and changes in the distribution of sea ice and coincident larval habitat (Melbourne-Thomas et al. 2016) emphasizing the importance of long-term effects. However, over the last 30 to 50 yr, climate-related changes in the primary atmospheric climate mode, the Southern Annular Mode (SAM), have strengthened the warm westerly winds and driven declines in sea-ice extent and duration around the Antarctic Peninsula (Stammerjohn et al. 2008b, Yuan & Li. 2008). As a result, annual sea-ice duration has declined by more than 90 d, with later ice formation and earlier melting. Thus, contemporary climate-induced changes will have more immediate ecological and management consequences before longer-term outcomes may come to fruition.

The declines in sea-ice extent and duration within the Antarctic Peninsula ecosystem will potentially increase access by the krill fishery to areas that were historically ice-covered, requiring appropriate management actions. As part of the Commission for the Conservation of Antarctic Marine Living Resources (CCAMLR) Ecosystem Monitoring Program (CEMP), risks to krill-dependent predators from fishing ac-
tivity are largely evaluated based on demographic monitoring studies conducted during the austral summer, when land-based predators (e.g., penguins and pinnipeds) are reproducitively active (Agnew 1997, Siniff et al. 2008). Comparatively less monitoring of predator populations has occurred during post-breeding periods, especially in winter. Many species that are monitored within the CEMP undergo late-summer migrations to overwinter in other areas of the Antarctic or sub-Antarctic (Stevick et al. 2004, Lea et al. 2008, Hinke et al. 2015). It was assumed that spatial overlap and negative interactions between these krill predators and the fishery were minimal during winter in the Antarctic Peninsula. The data here show a substantial abundance of pinnipeds, including Antarctic fur seals, which are monitored within the CEMP, and crabeater seals (an unmonitored species), coincident with regionally-confined, dense, krill aggregations in the BS. Recent tagging data on a number of krill predators (penguins and pinnipeds; Hinke et al. 2017; https://swfsc.noaa.gov/ AntarcticPredators/) also show that the BS is an important habitat during winter. This enormously concentrated and predictable food source provides a simple answer for the importance of this area to these predators.

Over the last decade, the krill fishery has increased its catch, shifted its main period of fishing from midsummer towards autumn and winter (in part owing to open water during autumn), and has become more spatio-temporally concentrated (CCAMLR 2014, Nicol & Foster 2016). In 2009, CCAMLR instituted interim catch limits for this area (Food and Agriculture Organization of the United Nations [FAO] Subarea 48.1; 155 000 t). Much of the catch has recently been taken in the BS, and in some years the fishery has operated into August. More recently, catch limits have been reached by mid-April and mid-May, resulting in early closures of the krill fishery in Subarea 48.1 in those years (Nicol et al. 2012). While the current catch is low relative to the estimated regional biomass for this area during winter (Table 2), there is considerable interest in expanding the fishery above the current 155 000 t limit within this area, which would allow continued fishing into winter. Thus, new data are needed to understand the potential effects of high local exploitation rates in winter, in addition to effects of the larger overall catch limits.

The current catch limits for krill apply to areas (e.g. all of Subarea 48.1) that are much larger than the area of the BS (CCAMLR 2014). It is unclear whether the current local exploitation rates within the BS increase the risk to meeting the objectives in Article II of the Convention on the Conservation of Antarctic Marine Living Resources, which requires that fishing impacts be reversible within 20 to 30 yr and that impacts on krill-dependent and associated predators be considered. Developing strategies to mitigate the increased risk to krill populations and their dependent predators in the face of changing ecosystem structure will likely require monitoring the winter distribution of krill and krill predators (Hinke et al. 2017), examining the effects of climate change over both the short and long term, and developing harvest-control rules to ensure the krill fishery is managed in an ecosystem-based context.

CONCLUSIONS

Here we have been able to capitalize on the fact that sea-ice extent and duration have declined and revealed the magnitude of the seasonal redistribution of krill from offshore to onshore, the location of a major krill overwintering ground, and the link to winter-habitat use for upper trophic level predators in the northern Antarctic Peninsula region. Yet, as climate change continues around the Southern Ocean, the structure of its pelagic ecosystems will change (Schofield et al. 2010). Over the long term, projected effects on the pelagic environment will include changes to krill habitat (Hill et al. 2013, Melbourne-Thomas et al. 2016) and krill productivity (Kawaguchi et al. 2013), suggesting the potential for a long-term change to the functional links within the Southern Ocean. Understanding the details of these changes will require studies throughout the ecosystem (Smetacek & Nicol 2005), including across seasons. Winter studies, especially studies that focus on the transition from winter to spring, are critically important to developing a better and more quantitative understanding of the structure and function of the Southern Ocean ecosystem. Such studies are also important to develop effective, precautionary management strategies that can allow for the rational development of the krill fishery while protecting krill-dependent predators.

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