Three-dimensional observations of swarms of Antarctic krill (Euphausia superba) made using a multi-beam echosounder

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A B S T R A C T

Antarctic krill (Euphausia superba) aggregate in dense swarms. Previous investigations of krill swarms have used conventional single- or split-beam echosounders that, with post-processing, provide a two-dimensional (2-D) view of the water column, leaving the third dimension to be inferred. We used a multi-beam echosounder system (SM20, 200 kHz, Kongsberg Mesotech Ltd, Canada) from an inflatable boat (length=5.5 m) to sample water-column backscatter, particularly krill swarms, directly in 2-D and, with post-processing, to provide a three-dimensional (3-D) view of entire krill swarms. The study took place over six days (2–8 February 2006) in the vicinity of Livingston Island, South Shetland Islands, Antarctica (62.4 S, 60.7 W). An automatic 3-D aggregation detection algorithm resolved 1006 krill swarms from the survey data. Principal component analyses indicated that swarm morphology metrics such as length, surface area and volume accounted for the largest between swarm variance, followed by echo energy, and finally swarm geographic location. Swarms did not form basic cylindrical or spherical shapes, but had quite consistent surface area to volume ratios of 3.3 m–1. Swarms were spatially segregated, with larger sizes (mean north-south length=276 m, at least double that of two other swarm classifications), found to the northwest of the survey area. The apparent clustering of swarm types suggests that krill biomass surveys and ecosystem investigations may require stratified survey design, in response to varying 3-D swarm morphology, variation that may be driven in turn by environmental characteristics such as bathymetry.

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1. Introduction

Many pelagic marine organisms exhibit patchy spatial distributions that are driven by a variety of biotic and abiotic factors (Genin, 2004). Aggregation appears to be a fundamental component of the behaviour of Antarctic krill (Euphausia superba), and indeed the swarm has been referred to as the “fundamental unit of krill ecology” (Murphy et al., 1988; Hofmann et al., 2004). Aggregations of Antarctic krill may form as a consequence of individuals seeking to reduce the chance of predation (O’Brien, 1987; Hammer and Hammer, 2000; Szulkin and Dawidowicz, 2006), to facilitate mating and/or feeding (Watkins et al., 1992; Miller et al., 1993; Hofmann et al., 2004), or to convey locomotive energetic advantage (Ritz, 1994, 2000).

Previous investigations suggest there are large variations in krill swarm shape (Miller et al., 1993; Wood-Walker et al., 2003) and packing density (Barange et al., 1993). Swarm dimensions and density have been used to classify swarms into types, and previous studies have implied spatial clustering of swarms by type (Miller and Hampton, 1989; Watkins and Murray, 1998). For example, Mauchline (1980) classified three types of aggregation based on numerical density: the densest swarm type contained from 1000 to 100,000 krill m–3, and was followed by logarithmically decreasing class densities of 1 to 100 and 0.1 to 1 krill m–3.

It is believed that some of the observed variation in swarm density and shape occurs in response to environmental conditions (Barange et al., 1993; Alonzo and Mangel, 2001; Hofmann et al., 2004), such as upwelling or localised water currents that occur in the vicinity of rapid changes in bathymetry (Trathan et al., 2003). It will be important to understand how krill aggregate to elucidate possible relationships between various potential physical or biological forcing mechanisms and swarm shape. This in turn would enable the extent to which krill demographics (e.g., age, sex, and maturity) are important to swarm formation (Watkins et al., 1986, 1992; Tarling et al., 2007). Understanding the mechanisms of krill swarm formation, and potential spatial variation in these mechanisms, may also be vital for unbiased
estimates of biomass (Gerlotto et al., 1999), and for addressing ecological issues such as predator–prey interactions (Zamon et al., 1996) and responses by krill to variation in their abiotic environment such as water depth (Hewitt and Demer, 2000; Trathan et al., 2003).

Most observations of krill swarms have been made acoustically with scientific echosounders, as per biomass surveys (e.g., Everson and Miller, 1994). Acoustic observations of krill swarms made using conventional vertically-downward looking single- or split-beam echosounders (SBE) are limited by the small conical sampling volumes (typically 7°) inherent with these instruments. Following each SBE acoustic transmission (ping), samples of volume backscattering strength (\(S_v\)) are recorded versus sound-propagation time, or water depth. Sequential recordings combine to build up a two-dimensional (2-D) matrix of water-column observations (Reid and Simmonds, 1993). The narrow acoustic beam effectively samples only a 2-D slice through the water column and any krill swarm in it. The three-dimensional (3-D) krill swarm shape cannot be estimated directly from 2-D observations without making assumptions about the swarm shape (e.g., it is cylindrical or spherical; Simmonds and MacLennan, 2005). Consequently, the volume of a krill swarm that falls outside of the narrow SBE beam cannot be determined. Krill biomass estimation techniques and investigations into krill ecology would both benefit from improved 3-D observations of individual krill swarms.

In this investigation, we used a multi-beam echosounder system (MBE) to observe Antarctic krill swarms in 2-D in the field, and extended these 2-D MBE observations into 3-D visualisations of swarms during post-processing. The purpose of this investigation was three fold. Firstly we sought to determine if krill could be observed in the nearshore environment using an MBE deployed from a small boat. Secondly, we wanted to investigate if the 3-D acoustic reconstruction of krill swarms could be used to improve understanding of the variability of swarm shape and density. Thirdly, we sought, using multivariate analyses, to examine the scale of variation of various swarm metrics. We hoped that, if successful, combining these elements could lead to future incorporation of MBEs into studies to improve understanding of krill biology and ecology.

2. Materials and methods

Two inflatable boats (Mark V Zodiacs, length=5.5 m) were deployed in the vicinity of Cape Sheriff, from 2 to 9 February, 2006. Operational constraints meant that a different number of transects was sampled each day (Fig. 1). The survey-site seabed depth ranged from 20 to 140 m. Seabed depth is an important consideration because it influences the MBE sampling volume and, due to side-lobe interference, the maximum observable across-track swarm width.

One inflatable boat, R/V Roald, was equipped with an MBE (200 kHz SM20, Kongsberg Mesotech Ltd, Canada) and conducted a high-resolution seabed-bathymetry survey (100% seabed coverage, resolution=1 m) with simultaneous water-column sampling to observe krill swarms acoustically. The second inflatable boat, R/V Ernests, was equipped with 38 and 200 kHz SBEs (calibrated single-beam Simrad ES60s). The MBE survey comprised 35 2.5-km transects and four 3.5-km transects, each with a 120 m inter-transect spacing (Fig. 1). On the final day of surveying, two ‘tie lines’ of length=5.2 km with spacing=1.2 km were run perpendicular to the main transects, the purpose of which was to assess any day affect (i.e. possible day-to-day variation) in the MBE krill swarm data. In addition to acoustic observations, visual observations of air-breathing krill predators were collected from both Zodiacs (Cox et al., 2009).

2.1. Multibeam equipment and data description

The MBE had a total swath width of 120°, made up of 128 receive beams each with a 1.5° across-track and 20° along-track beam width. The MBE head was mounted facing vertically downwards, along the centreline of the boat, so ensoniﬁed a 60° swath either side of the track line. An orthogonally-mounted (with respect to the MBE head) external proﬁling transmitter was used and reduced the along-track beam width from 20° to 1.5°. This improved the precision of water-column target sampling and reduced between-pong along-track sampling volume overlap. Acoustic pulses were transmitted every 1.5 to 3 s (this varied because of computer processing limitations). Time varied gain was 20 \(\log_{10}(\text{range})\), pulse length was 825 μs, and the transmission power was set to ‘medium’. MBE detections throughout the ﬁxed 200-m observation range had 0.5 m resolution to standardise sampling and were logged continuously to the control computer. Recorded MBE data were converted to the SM2000 data format using Kongsberg Mesotech MsToSm (v1.0), and resulting \(S_v\) data were processed using Echoview v3.5 (Myriax, Hobart, Australia).

2.2. Scaling uncalibrated MBE \(S_v\) observations

The MBE cannot be calibrated easily in the field using the conventional standard reference sphere techniques (Foote et al., 1987) that would typically be applied to SBEs. Consequently, the uncalibrated data from the MBE were calibrated by comparison to the \(S_v\) observations calibrated by the standard sphere method and collected by the ES60 along the tie lines (two lines run...
perpendicularly to the 41 main MBE survey transects, Fig. 1). The tie lines were surveyed with the SBE-equipped R/V Ernest following the MBE-equipped R/V Roald. The Zodiacs were separated by approximately 200 m to avoid acoustic cross-talk between the 200 kHz systems, but essentially sampled the same bodies of water. Thus, it was assumed that both instruments sampled the same distribution of water-column backscatter.

The $S_v$ data from the SBE and the nadir beam (number 63) of the MBE were compared. Krill swarm boundaries were determined using the SHAPES algorithm implemented in the Echoview schools detection module (see Barange, 1994; Coetzee, 2000). Then, the data were spatially aligned and averaged in 2 ping by 3 m vertical cells. Echoes from krill were identified in the gridded SBE data using the dB difference technique (Brierley et al., 1998; $-0.7 < S_v_{200 kHz} - S_v_{38 kHz} < 13.3$ dB). Gridded mean $S_v$ values from both instruments for grid cells lying within the SBE detected swarms were used to determine the cumulative density functions (CDFs) for the mean $S_v$ from both the SBE ($CDF_{SBE}$) and the MBE ($CDF_{MBE}$). Both CDFs were calculated using

$$p(t) = \frac{1}{n} \sum_{i=1}^{n} I(x_i)$$

where $I(x_i)$ is an indexing function:

$$I(x_i) = \begin{cases} 1 & : x_i \leq t \\ 0 & : \text{otherwise} \end{cases}$$

$t$ is the required resolution for the CDF mapping, $i$ is $S_v$ observation number and $n$ is the total number of $S_v$ observations. The two CDFs were mapped by determining which gridded $S_v$ values from both acoustic instruments occurred at equal proportions $p(t)$ along their respective CDFs (left panel, Fig. 2). This between-instrument map was used as a look-up table to rescale the MBE $S_v$ observations to equivalent ES60 $S_v$.

2.3. Automated 3-D krill swarm detection and sensitivity analysis

Krill swarms were identified from the MBE observations using a 3-D school detection algorithm (Echoview v 3.5 multibeam module, Myriax). The algorithm identifies contiguous groups of acoustic returns in each beam and bounds with prisms the extremities of each acoustic return group. These prisms are triangulated, reducing each prism to two triangles. The perimeter of the 3-D school is generated by retaining the visible vertices of the triangles, which are used to create a 3-D bounding surface around the contiguous acoustic return. At this point, the user-defined size parameters, minimum longest dimension, minimum middle dimension and minimum shortest dimension, are used to eliminate detected swarms with dimensions smaller than these minimum parameters. In addition, the minimum $S_v$ threshold (dB re m$^{-1}$, $S_v$) defines the minimum density of acoustic returns that are transferred to the 3-D detection algorithm, hence defining the krill swarm boundary. Acoustic observations of ranges less than 5 m were ignored due to sea-surface noise and near-field effects (Melvin et al., 2003). For the purposes of 3-D target detection, the search volume for the 3-D algorithm was constrained to water column targets by referencing 0.5 m above the seabed as defined by the MBE sounder detected bottom identification algorithm. For each identified swarm, various metrics were extracted (see Table 1 for description of quantitative swarm descriptor variables).

The sensitivity of perceived swarm metrics (Table 1, shown in italics) to variation of the 3-D detection algorithm parameters was investigated (Table 2) and used to select an optimum set of parameters as follows.

Four transects, (numbers 2, 17, 22 and 33), were selected at random from the total of 41 transects run to furnish data for the sensitivity analysis. The sensitivity of the 3-D school detection parameters was investigated in two stages. Firstly, the three minimum dimension parameters (minimum longest, middle and

![Fig. 2. SM20 calibration. Left panel: The cumulative distribution function (CDF) mapping of the uncalibrated SM20 multi-beam echosounder $S_v$ observations and calibrated ES60 single-beam echosounder $S_v$ observations (Section 2.2). Right panel: The linear relationship between the CDF of the SM20 and ES60 in the linear domain ($S_v$). The CDF linear relationship occurs in the range $-70$ dB $< ES60 S_v < -50$ dB. The linear regression ($r^2 = 0.96$, $p < 2.2e^{-16}$, black line) and 95% prediction intervals (dotted black line) are shown.](image)
shortest aggregation dimensions) were varied. In the absence of any prior information about the 3-D shape of krill swarms in the nearshore study region, the same length was used for each of the 3-D detection length parameters (Table 2), with the minimum MBE detection threshold fixed at $S_{\text{v}} = 51.8$ dB re 1 m$^{-1}$ or approximately 166 krill m$^{-3}$ using a $TS = 74$ dB re m$^2$. The second part of the sensitivity analysis investigated the effect of varying the minimum threshold, with the minimum school dimensions set to 5 m.

2.4. 3-D krill swarm descriptors

The 2-D nearest-neighbour-distance (NND) was calculated using the great circle distance between the geocentric latitude and longitude of each detected krill swarm (Table 1). To determine the krill swarm metrics that accounted for the largest inter-swarm differences, a principal component analysis (PCA) was performed. Because some of the 3-D swarm metrics had different units and scales (e.g., swarm geometric latitude and swarm volume) the PCA was carried out on a matrix of normalised individual principal components than average, and thus made the PCA eigen vectors, individual eigen vector elements ($u_{ij}$) greater than a scaling factor ($u_{ij} > 0.7 \cdot \text{max} \{u_{ij}\}$) (Mardia et al., 1979) was used to identify which swarm metrics exerted more influence on individual principal components than average, and thus made the greatest contribution to inter-swarm variation. A three-type partition analysis was then performed in the reduced factorial space provided by the PCA results to determine if swarms formed distinct types. A three-type partition was selected based on the classification schemes of Mauchline (1980). The appropriateness of the three-type analysis was validated by the results of a pseudo $F$-test for analyses with different numbers of clusters (Johnson, 1998), that determined differences in the sums of squares of within cluster differences.

### Table 1

<table>
<thead>
<tr>
<th>Metric name</th>
<th>Abbreviation</th>
<th>Units</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Surface area</td>
<td>$S$</td>
<td>m$^2$</td>
<td>Surface area of the 3-D krill swarm</td>
</tr>
<tr>
<td>North-south length</td>
<td>$l_{NS}$</td>
<td>m</td>
<td>Maximum length of the 3-D swarm along a north-south</td>
</tr>
<tr>
<td>East-west length</td>
<td>$l_{EW}$</td>
<td>m</td>
<td>Maximum length of the 3-D swarm along an east-west</td>
</tr>
<tr>
<td>Height</td>
<td>$h$</td>
<td>m</td>
<td>Maximum swarm height ($z_{\text{max}}-z_{\text{min}}$). $z_{\text{max}}=$maximum swarm depth, $z_{\text{min}}=$minimum swarm depth.</td>
</tr>
<tr>
<td>Volume</td>
<td>$V$</td>
<td>m$^3$</td>
<td>3-D swarm volume</td>
</tr>
<tr>
<td>Roughness</td>
<td>$R$</td>
<td>m$^{-1}$</td>
<td>$R=S/V$</td>
</tr>
<tr>
<td>Length/height ratio</td>
<td>$l/h$</td>
<td></td>
<td>length/height</td>
</tr>
<tr>
<td>$S_{\text{v}}$–mean</td>
<td>$S_{\text{v}}$</td>
<td>dB re 1 m$^{-1}$</td>
<td>Mean volume backscattering strength within a swarm</td>
</tr>
<tr>
<td>$S_{\text{v}}$–min</td>
<td>$S_{\text{v}}$</td>
<td>dB re 1 m$^{-1}$</td>
<td>Minimum $S_{\text{v}}$</td>
</tr>
<tr>
<td>$S_{\text{v}}$–max</td>
<td>$S_{\text{v}}$</td>
<td>dB re 1 m$^{-1}$</td>
<td>Maximum $S_{\text{v}}$</td>
</tr>
<tr>
<td>Seabed depth under swarm*a</td>
<td>$z$</td>
<td>m</td>
<td>Seabed depth under geometric centre of swarm</td>
</tr>
<tr>
<td>Position in water column*a</td>
<td>$pos_{\text{mc}}$</td>
<td></td>
<td>$pos_{\text{mc}}=1-(z_{\text{max}}-z_{\text{o}})$ giving 0 &lt; $pos_{\text{mc}}$ &lt; 1</td>
</tr>
<tr>
<td>geometric centre position</td>
<td>$(\phi, \lambda)$</td>
<td>deg</td>
<td>Latitude and longitude of geometric centre of 3-D krill swarm.</td>
</tr>
<tr>
<td>geometric centre depth</td>
<td>$Z$</td>
<td>m</td>
<td>Depth at the geometric centre of the 3-D krill swarm.</td>
</tr>
<tr>
<td>Nearest neighbour distance</td>
<td>NND</td>
<td>m</td>
<td>Radial distance calculated using great circle distance</td>
</tr>
<tr>
<td>Time of day (GMT)</td>
<td>$t$</td>
<td></td>
<td>Time of swarm observation</td>
</tr>
<tr>
<td>Cross track distance*a</td>
<td>$x$</td>
<td>m</td>
<td>Cross track distance from geometric centre of swarm to MBE</td>
</tr>
</tbody>
</table>

*a Not calculated using Echoview.

3. Results

Using 'optimal' detection parameters (see Sections 2.3 and 3.1), a total of 1084 krill swarms were observed using the MBE, of which 1006 were determined to be entirely within the MBE sampling volume (i.e. not extending into side lobes or beyond the outermost beams).

3.1. 3-D detection algorithm sensitivity analysis

Sensitivity analysis of the minimum school dimension showed that the number of detected swarms decreased with increasing minimum swarm dimensions (Fig. 3). Median swarm height, north-south and east-west length, and swarm volume all increased with increasing minimum swarm dimensions. Median minimum NND increased with increasing minimum swarm size, partly since fewer swarms were detected. The NND results at larger minimum swarm sizes indicated that the few remaining large, high-density ($S_{\text{v}} > S_{\text{v}}$) krill swarms were not spatially clustered.

While the sensitivity plots (Fig. 3) were informative, there was insufficient information to determine analytically the minimum dimension swarm detection parameters. Visual inspection of the 3-D krill swarm boundaries generated using the detection algorithm showed that the 2-m minimum school dimension (Table 2) caused splits in larger swarms and an increase in the number of seabed side-lobe detections being identified spuriously as aggregations. It was not possible to remove seabed side-lobe detections prior to running the 3-D detection algorithm because the side-lobe detections varied from ping-to-ping. The 5-m minimum school dimension reduced spurious identification of side lobes and prevented spurious identification of krill swarm boundaries, but did occasionally split krill swarms. The 10-m minimum length setting degraded the detected krill swarm spatial resolution and eliminated small swarms. This degradation of krill swarm spatial structure and elimination of smaller krill swarms persisted with increasing minimum 3-D length settings.

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Consequently the 5-m minimum length setting was selected as a compromise.

A sensitivity analysis of processing threshold $S_v$ was not used because the number of background noise detections prevented the 3-D school detection algorithm from functioning correctly at low processing thresholds. The relationships between number of detected swarms and $S_v$ showed that more swarms were detected at lower $S_v$ (Fig. 4). The maximum median swarm height and volume occurred when $S_v = -52.7 \text{ dB re } 1 \text{ m}^{-1}$ (ca. 133 animals m$^{-3}$). At lower $S_v$, the 3-D detection metrics contain data from side lobe and noise detections that generally result in 3-D detections with smaller volumes than those attributed to 3-D krill swarm detections. At $S_v > -52.7 \text{ dB re } 1 \text{ m}^{-1}$, the lower density edges of detected krill swarms were eroded, resulting in reductions in all median krill swarm metrics. Given that at $S_v = -57.7 \text{ dB re } 1 \text{ m}^{-1}$ maximum swarm height and volume occurred, and the number of spurious side lobe detections fell, this was selected as the $S_v$ for 3-D schools detection of krill swarms. Therefore, for the purposes of this investigation, a krill swarm is defined as a collection of acoustic samples with $S_v = -57.7 \text{ dB re } 1 \text{ m}^{-1}$ with spherical dimensions $> 5 \text{ m}$. Using these 3-D detection parameters, a conventional vertically-downward looking SBE with a 7° beam would not sample an entire spherical krill swarm with a 5 m diameter until a depth of 57 m. Only 52% of MBE detected swarms were found deeper than 57 m, which illustrates the constraints prevalent with SBE: swarms of the 5-m example diameter cannot be entirely sampled with an SBE until a depth of at least 57 m. Despite the external envelope of swarms being defined using 3-D schools detection parameters of 5 m for shape and $S_v = -57.7 \text{ dB re } 1 \text{ m}^{-1}$
re 1 m⁻¹, the $S_v$ was removed when analysing the observations within the 3-D swarm boundary. This means the energetic measures of swarms are essentially unthresholded values.

### 3.2. Krill swarm morphology

As an initial investigation of swarm morphological types, the ratio of swarm length to height ($l/h$) was calculated. A continuous unimodal distribution was observed (Fig. 5), showing that with respect to $l/h$, there were no distinct krill swarm types. Further, it is evident from the $l/h$ distribution that the krill swarms were not spherical (i.e. spheres have $l/h=1$).

Krill swarms within the survey area exhibited a broad range of morphological variation, particularly in volume and area. Swarms generally were longer in the north-south direction, than east-west (Table 3), which may potentially be a consequence of current direction. On occasion, swarms filled up to 33% of the water column, but height and swarm minimum and maximum depths may otherwise have been constrained by water depth. The swarm energetic measures showed lower variation than morphological metrics, which may be biologically driven or a function of the sensitivity of the MBE (Table 3, coefficient of variation $S_v$, mean and $S_v$, max), or both. Swarm roughness ($R=\text{surface area}/\text{volume}$) had the lowest coefficient of variation (CV) for the morphological parameters, perhaps suggesting some underlying biological constraint (Cox et al., 2009).

The first three components of the PCA accounted for 22, 15 and 13%, respectively; of the observed variance in krill swarm structure (Table 3). The first principal component was influenced significantly by swarm volume, surface area, north-south length, east-west length and height. The second component was

![Fig. 4. Sensitivity of krill swarm descriptive metrics to variations in the minimum detection threshold ($S_v$). Points are median metric values with 25% and 75% quantiles. The selected $S_v=23$ dB re 1 m⁻¹ is shown as black circles. X-axis scale is given in uncalibrated MBE $S_v$ (upper-scale) and ECDF mapped, approach one, MBE $S_v$ (lower-scale). Note *Method 1 calibration by comparison used (see Section 2.2). The 75% quantile for median swarm volume is 3,311 m³ at $S_v=23$ dB re 1 m⁻¹ and 3,421 m³ at $S_v=24$ dB re 1 m⁻¹.](image)
influenced significantly by the length to height ratio, and the third by geographical position and the swarm energetic parameters $S_v$ and $S_v_{\text{max}}$. No significant differences (at $p=0.05$) were found between the three-class cluster and other $n$-class clusters ($n=2$ to $10$) using Beale's pseudo $F$-statistic test (Johnson, 1998) for significant differences in the residual sums of squares of the intra-cluster distances between two possible $n$-class swarm clusterings (e.g., a comparison between a 2-class clustering and the 3-class clustering).

The swarm partition analysis was carried out in the reduced factorial space determined by PCA. Swarm type one contained 158 swarms, and types two and three contained 431 and 417 swarms, respectively (Table 4). Swarm type three was the most distinct, exhibiting little overlap with the other types in principal

<table>
<thead>
<tr>
<th>Variable</th>
<th>Mean (CV)</th>
<th>Range</th>
<th>PC1</th>
<th>PC2</th>
<th>PC3</th>
</tr>
</thead>
<tbody>
<tr>
<td>Roughness (m$^{-1}$)</td>
<td>3.3 (0.23)</td>
<td>1.2 to 8.1</td>
<td>0.115</td>
<td>-0.264</td>
<td>-0.016</td>
</tr>
<tr>
<td>Volume (m$^3$)</td>
<td>3695.7 (4.59)</td>
<td>46.2 to 406,709.8</td>
<td><strong>0.385</strong></td>
<td>0.28</td>
<td></td>
</tr>
<tr>
<td>Area (m$^2$)</td>
<td>11,024.7 (4.70)</td>
<td>218.6 to 1,222,048</td>
<td><strong>0.378</strong></td>
<td>0.278</td>
<td></td>
</tr>
<tr>
<td>North-south length (m)</td>
<td>120.2 (2.34)</td>
<td>9.9 to 4793.6</td>
<td><strong>0.384</strong></td>
<td>-0.271</td>
<td>-0.163</td>
</tr>
<tr>
<td>East-west length (m)</td>
<td>86.22 (2.03)</td>
<td>9.2 to 2959.5</td>
<td><strong>0.397</strong></td>
<td>-0.259</td>
<td>-0.14</td>
</tr>
<tr>
<td>Height (m)</td>
<td>10.6 (0.70)</td>
<td>3.9 to 77.5</td>
<td><strong>0.384</strong></td>
<td>0.147</td>
<td>0.217</td>
</tr>
<tr>
<td>Length/height</td>
<td>11.7 (1.34)</td>
<td>5.0 to 206.5</td>
<td>0.22</td>
<td><strong>0.374</strong></td>
<td>-0.244</td>
</tr>
<tr>
<td>Latitude (deg)</td>
<td>-62.44</td>
<td>-62.46 to -62.41</td>
<td>0.127</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Longitude (deg)</td>
<td>-60.7</td>
<td>-60.77 to -60.66</td>
<td>-0.14</td>
<td>-0.18</td>
<td><strong>0.516</strong></td>
</tr>
<tr>
<td>Seabed depth (m)</td>
<td>94.6 (0.12)</td>
<td>45.3 to 134.2</td>
<td>0.185</td>
<td>0.214</td>
<td>0.275</td>
</tr>
<tr>
<td>Position in water column</td>
<td>0.6 (0.32)</td>
<td>0 to 1</td>
<td>0.185</td>
<td>0.214</td>
<td>0.275</td>
</tr>
<tr>
<td>$S_v_{\text{mean}}$ (dB re 1 m$^{-1}$)</td>
<td>-56.17 (0.14)</td>
<td>-99.77 to -44.75</td>
<td><strong>0.468</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>$S_v_{\text{min}}$ (dB re 1 m$^{-1}$)</td>
<td>-101.40 (14.75)</td>
<td>-108.03 to -51.62</td>
<td>0.206</td>
<td>0.148</td>
<td>-0.123</td>
</tr>
<tr>
<td>$S_v_{\text{max}}$ (dB re 1 m$^{-1}$)</td>
<td>-51.04 (0.12)</td>
<td>-51.04 to -44.45</td>
<td><strong>0.393</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Swarm nearest neighbour distance (m)</td>
<td>36.8 (0.93)</td>
<td>3 to 329.5</td>
<td>0.205</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Time of day (GMT)</td>
<td>16:35</td>
<td>12:58 to 22:10</td>
<td>0.122</td>
<td>-0.347</td>
<td></td>
</tr>
<tr>
<td>Cross-track distance (m)</td>
<td>27.1 (0.70)</td>
<td>0.9 to 94.0</td>
<td>-0.192</td>
<td>0.172</td>
<td></td>
</tr>
</tbody>
</table>

Fig. 5. Histogram of length/height ratio of 3-D detected krill swarms.

Please cite this article as: Cox, M.J., et al., Three-dimensional observations of swarms of Antarctic krill (*Euphausia superba*) made using a multi-beam echosounder. Deep-Sea Research II (2010), doi:10.1016/j.dsr2.2009.10.003
### Table 4
Means of the three partition types for selected krill swarm metrics and associated coefficients of variation.

<table>
<thead>
<tr>
<th>Swarm metric</th>
<th>Swarm type</th>
<th>1</th>
<th>2</th>
<th>3</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number of swarms</td>
<td></td>
<td>158</td>
<td>431</td>
<td>417</td>
</tr>
<tr>
<td>Roughness</td>
<td></td>
<td>3.23 (7.78)</td>
<td>2.96 (5.07)</td>
<td>3.65 (4.34)</td>
</tr>
<tr>
<td>Seabed depth (m)</td>
<td></td>
<td>-51.89 (11.57)</td>
<td>-51.65 (7.75)</td>
<td>-57.44 (8.96)</td>
</tr>
<tr>
<td>Position in water column</td>
<td></td>
<td>0.78 (6.32)</td>
<td>0.73 (4.46)</td>
<td>0.55 (2.83)</td>
</tr>
<tr>
<td>Nearest-neighbour distance (m)</td>
<td></td>
<td>37.95 (1.39)</td>
<td>47.72 (1.11)</td>
<td>26.01 (1.16)</td>
</tr>
<tr>
<td>North-south length (m)</td>
<td></td>
<td>276.39 (0.41)</td>
<td>82.21 (1.17)</td>
<td>100.20 (2.06)</td>
</tr>
<tr>
<td>East-west length (m)</td>
<td></td>
<td>181.85 (0.43)</td>
<td>64.97 (1.22)</td>
<td>71.96 (2.02)</td>
</tr>
<tr>
<td>Height (m)</td>
<td></td>
<td>13.46 (1.21)</td>
<td>11.98 (1.57)</td>
<td>8.05 (2.24)</td>
</tr>
<tr>
<td>$S_{v_{\min}}$ (dB re 1 m/$C_0$)</td>
<td></td>
<td>-100</td>
<td>-100</td>
<td>-100</td>
</tr>
<tr>
<td>$S_{v_{\max}}$ (dB re 1 m/$C_0$)</td>
<td></td>
<td>-48.30 (11.88)</td>
<td>-48.32 (7.00)</td>
<td>-49.23 (13.75)</td>
</tr>
<tr>
<td>Cross-track distance (m)</td>
<td></td>
<td>33.87 (1.21)</td>
<td>29.80 (1.62)</td>
<td>40.42 (1.60)</td>
</tr>
<tr>
<td>Surface area (m$^2$)</td>
<td></td>
<td>29.197.01 (0.24)</td>
<td>11.554.54 (0.44)</td>
<td>5937.07 (0.24)</td>
</tr>
<tr>
<td>Volume (m$^3$)</td>
<td></td>
<td>9237.07 (0.24)</td>
<td>4220.67 (0.43)</td>
<td>1053.37 (0.86)</td>
</tr>
<tr>
<td>Length/height</td>
<td></td>
<td>19.46 (0.54)</td>
<td>7.24 (1.67)</td>
<td>13.80 (1.83)</td>
</tr>
<tr>
<td>Swarm depth (m)</td>
<td></td>
<td>54.6 (0.32)</td>
<td>58.0 (0.32)</td>
<td>58.5 (0.29)</td>
</tr>
<tr>
<td>Seabed gradient</td>
<td></td>
<td>0.19 (0.46)</td>
<td>0.23 (0.89)</td>
<td>0.21 (0.36)</td>
</tr>
</tbody>
</table>

Fig. 6. Three-type partition analysis of krill swarms performed in principal component analysis reduced factorial space. The perceptive plot (top left) shows the first three principal components. Other panels show pairs of the first three principal components plotted on plane surfaces, illustrating changing overlap/separation between swarm types.
component planes 1 and 2, or planes 2 and 3 (Fig. 6). The other two swarm types showed increasing overlap, which is perhaps evidence that krill swarms were actually drawn from a continuum of swarm types, rather than a population of discrete swarm types.

Examining the geographic position of partitioned swarms revealed that type-one swarms were found entirely in the northwest of the survey area, whereas types two and three were found dispersed throughout the remainder of the site. A higher density of type-three swarms was found toward the east of the survey region, a region of particularly shallow water (depth=25 m, Fig. 7). Despite this cluster of type-three swarms in the shallow water area, seabed slope did not vary significantly between the swarm types (Table 4). No air-breathing predators were observed in the region around the type-one krill swarms (Fig. 7).

While analysis of the l/h failed to separate krill swarms into distinct morphological types, there were distinct differences in this ratio across the three partitioned types: type-one had almost double the l/h ratio of type-three (Table 4). Swarms in type-one, while being smaller, exhibited a different morphological structure that was not simply rescaled for larger swarm sizes. For example, the north-south and east-west lengths of type one swarms were more than double that of types-two and three, but heights were not doubled between swarm types. This suggests that swarm types are not simply driven by rescaling of a ‘standard’ swarm.

4. Discussion

This investigation has shown that MBE (200 kHz Simrad-Mesotech SM20) is capable of detecting Antarctic krill swarms, and that small boats are suitable platforms for conducting acoustic surveys in areas inaccessible to ocean-going research vessels. Further application of MBE technology deployed from a variety of platforms – e.g., boats, moorings, and autonomous underwater vehicles (AUVs) – has the potential to contribute much to krill research (Nicol and Brierley, 2010). Using the MBE, we found that krill swarms formed complex 3-D shapes, which were spatially segregated, and that, potentially, the nearshore waters of the South Shetlands contain a large biomass of krill.

Our analyses were based on an objectively-defined krill swarm boundary, identified using a 3-D detection algorithm, with parameters selected by sensitivity analysis (Table 2). Previous researchers have used automated ping-by-ping image recognition software to delineate pelagic aggregations observed by omnidirectional MBEs in 3-D (e.g., Misund et al., 1998; Brehmer et al., 2006). It is our understanding, however, that this investigation is the first MBE study that has assessed the influence of the parameters used in an automatic 3-D detection algorithm on the resultant aggregation metrics.

Previously, researchers have used single aggregation metrics, or ratios of aggregation metrics to describe aggregation type. For example, Gerlotto et al. (2004) acquired 3-D data from aggregations of anchovy (Engraulis ringens) and common sardine (Strangomera bentincki) using an MBE, and suggested that layers and schools could be partitioned by an aggregation length-to-height ratio of l/h=7. No such break point occurred in the observed l/h ratios of the krill swarms detected in this investigation (Fig. 6). Rather, the MBE results suggest a continuum of swarm l/h: the use of single swarm metrics or metric ratios are not useful discriminators for the krill swarms observed during this investigation. Consequently, multivariate analysis techniques were employed.

The results of the multivariate analysis techniques employed during this research (Tables 3 and 4, Fig. 7) demonstrated that groups may occupy different geographical areas (Fig. 7), and that spatial segregation between groups can occur over small distances (ca. 3 km) and have sharp boundaries (Fig. 7). Through the use of two tie lines, surveyed perpendicularly to the main survey lines (Fig. 1 and Fig. 7), that cut through the south-east and north-west areas of the study site, we have been able to discount a day affect in our observations. If swarm segregation was simply caused by a day affect, then type-one swarms would have been detected in the eastern end of the survey site and types-two and three in the western area of the...
survey site. However, only one swarm was detected in the western area along a tie line, and this was a type-one swarm providing strong evidence that the swarm type segregation was truly spatial and not a day effect (Fig. 7). The temporal persistence of swarm type, elucidated by data collected along the surveyed tie lines, suggests that the causes of swarm-type segregation may be spatially persistent, such as water depth or prevailing water current.

We cannot be certain of the causes of the spatial persistence of krill swarms observed during this investigation. The next step would be to sample the physical environment and to make targeted trawls through individual krill swarms to examine possible relationships between swarm metrics and characteristics of krill within swarms such as length-distribution, stage, and sex ratio. This would enable assessment of the role of swarm size-sorting mechanisms, as suggested by Watkins et al. (1992) and previously observed on the west of the Antarctic Peninsula by Lascara et al. (1999).

The spatial segregation of krill-swarm types in nearshore areas may be of significance for land-based, air-breathing krill predators that have constrained foraging-trip durations during the breeding season. Should swarms of different types be comprised of krill that are of different sizes, then the predators may gain energetic benefits from focussing foraging effort on particular swarm types. Conversely, escape responses of individual krill in a swarm may also vary by swarm type: naïve individuals in a swarm may have ineffective escape response (O’Brien, 1989; Ritz, 1994). The potential differences in krill escape response, coupled with the swarm spatial segregation observed during this study (Fig. 7), suggest that krill catchability by predators may vary considerably at small scales (ca. 3 km), and may explain why no air-breathing predators were observed foraging in the vicinity of type-one swarms (Fig. 7).

The 3-D classification of krill swarm types may in the future assist in the partitioning of acoustic echoes to separate those arising from krill swarms and those from other species. Such an analysis was carried out by Woodd-Walker et al. (2003) using 2-D observations of krill swarms from conventional, vertically downward-oriented, single-beam echosounders. However, problems with the classification of acoustic targets remains. Fielding et al. (2007) suggested that Mackeral icefish (Champsocephalus gunnari) have been incorrectly acoustically classified as Antarctic krill using a two-frequency dB-difference technique (Brierley et al., 1998), thus positively biasing krill areal density estimates, \( \hat{p} \). Considering 3-D krill-swarm metrics, rather than just a conventional two-frequency dB-difference technique, may enable the acoustic echoes from mackerel icefish to be eliminated from \( \hat{p} \), thus reducing the positive bias that arises from incorrect identification of acoustic echoes.

This research has demonstrated the krill can be detected with MBEs, and that the data are useful for ecological studies, e.g., determining the spatial segregation of swarm types. Considerable work is still required before MBE data can be used for directly calculating \( \hat{p} \), such as in devising field-calibration procedures. Fortunately, some progress is being made with: MBE calibration procedures (e.g., Foote et al., 2005); estimating krill target strength at any insonified angle (e.g., Demer and Conti, 2005); and accounting for the directivity of krill scatter perceived by an MBE (e.g., Cutter et al., 2009). Even now, however, 3-D swarm type information derived from MBE observations may enable researchers to improve identification of krill swarms and exclude other species from biomass estimates, thereby reducing bias. MBEs provide a wider-eyed view of the world of krill, and we expect MBEs to add much to our understanding of this key Antarctic species.

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References


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