Distribution and Habitat Suitability of Ross Seals in a Warming Ocean

Mia Wege1,2,3*, Horst Bornemann1, Arnoldus Schytte Blix4, Erling Sverre Nordøy4, Louise Biddle5 and Marthán Nieuwoudt Bester2

1 Alfred-Wegener-Institut, Helmholtz-Zentrum für Polar- und Meeresforschung, Bremerhaven, Germany, 2 Mammal Research Institute, Department of Zoology & Entomology, University of Pretoria, Pretoria, South Africa, 3 Gateway Antarctica, School of Earth and Environment, University of Canterbury, Christchurch, New Zealand, 4 Department of Arctic and Marine Biology, UiT The Arctic University of Norway, Tromsø, Norway, 5 Department of Marine Sciences, University of Gothenburg, Gothenburg, Sweden

Understanding the determinants of poorly studied species’ spatial ecology is fundamental to understanding climate change impacts on those species and how to effectively prioritise their conservation. Ross seals (Ommatophoca rossii) are the least studied of the Antarctic pinnipeds with a limited knowledge of their spatial ecology. We present the largest tracking study for this species to date, create the first habitat models, and discuss the potential impacts of climate change on their preferred habitat and the implications for conservation. We combined newly collected satellite tracking data (2016–2019: n = 11) with previously published data (2001: n = 8) from the Weddell, King Haakon VII and Lazarev seas, Antarctica, and used 16 remotely sensed environmental variables to model Ross seal habitat suitability by means of boosted regression trees for summer and winter, respectively. Five of the top environmental predictors were relevant in both summer and winter (sea-surface temperature, distance to the ice edge, ice concentration standard deviation, mixed-layer depth, and sea-surface height anomalies). Ross seals preferred to forage in waters ranging between −1 and 2°C, where the mixed-layer depth was shallower in summer and deeper in winter, where current speeds were slower, and away from the ice edge in the open ocean. Receding ice edge and shoaling of the mixed layer induced by climate change may reduce swimming distances and diving depths, thereby reducing foraging costs. However, predicted increased current speeds and sea-surface temperatures may reduce habitat suitability in these regions. We suggest that the response of Ross seals to climate change will be regionally specific, their future success will ultimately depend on how their prey responds to regional climate effects and their own behavioural plasticity.

Keywords: Antarctica, biologging, boosted regression trees models, climate change, habitat utilisation, species distribution model, remote sensing, Weddell Sea

INTRODUCTION

Climate change and extreme weather events are increasing and as a consequence are influencing the distribution and movements of species globally (Karl and Trenberth, 2003; Meehl et al., 2007). Species are shifting higher in altitude or poleward, but most species will not be able to shift their distributions or adapt their movement behaviours fast enough to keep up with climate shifts...
Elusive of the Antarctic seals (e.g., Würsig et al., 2018). They observations made during surveys from helicopters and ships distribution and habitat use of ice breeding seals are based on data because most of the available data and understandings of species’ habitat and distributions are biased toward haul-out (Bestley et al., 2020). Additionally, knowledge of pagophilic seal paucity in winter data), and ease of access to species [pack-ice of data on juveniles and non-breeders), timing of expeditions of these mobile consumers is biased by life-history phase (lack other components within this environment although, knowledge seabirds, and penguins) are relatively well studied compared to other well-studied Antarctic marine predators because they travel away from the Antarctic pack-ice and forage pelagically most of the year (Blix and Nordøy, 2007; Arcalís-Planas et al., 2015).

Given their pelagic foraging behaviour (Blix and Nordøy, 2007; Arcalís-Planas et al., 2015), Ross seals are expected to respond differently to climatic shifts and extreme weather events compared to other Southern Ocean predators. Because Ross seals spend most of their lives in the open ocean (Blix and Nordøy, 2007; Arcalís-Planas et al., 2015), it is suggested that their foraging behaviour is least likely to be negatively affected by changes in sea-ice coverage, unless the preferred prey species respond by shifting their distributional range (Siniff et al., 2008). A reduction in winter sea-ice extent may result in open pelagic waters that are closer to pack-ice areas where Ross seals haul out to breed and moult, which reduces swimming distance to their pelagic foraging grounds after breeding and moulting.

The lack of data for this species restricts the extent of our potential predictions of Ross seal behaviour in response to climate change. The limiting factor is baseline data on the oceanographic conditions that characterise the foraging habitats and availability of prey of Ross seals. Here, we combine recently collected tracking data (2016, 2018, 2019) from Ross seals in the eastern Weddell Sea and the area to the east, off Queen Maud Land (including the King Haakon VII and Lazarev seas) with the only other movement data of Ross seals from the same area (Blix and Nordøy, 2007). We build seasonal habitat suitability models using remotely sensed oceanographic variables to (1) identify oceanographic conditions that characterise summer and winter foraging habitat of Ross seals in the Weddell Sea and adjoining areas to the east; and (2) predict suitable foraging habitat for them during summer and winter, respectively in that area. We further discuss the implications of Ross seals’ foraging habitat in response to predicted climate change effects.

MATERIALS AND METHODS

Study Area
Ross seals were captured, instrumented and sampled in the King Haakon VII Sea off Queen Maud Land (6954’S–7220’S and 200’W–1746’W) and in two cases in the marginal sea-ice zone, north of the Lazarev Sea around ∼59°S and 5–21°E in spring 2019 (Supplementary Table 1). We combined tracking data of Ross seals collected within this region during South African National Antarctic Expedition S55 (December 2015–February 2016), the German Antarctic Expedition PS111 (January 2018–March 2018), and the South African Southern Ocean Seasonal Experiment a.k.a. SCALE (October–November 2019), with published data from Norwegian Antarctic Research
Capturing and Tag Deployment
Ross seals were captured and physically restrained on pack-ice floes in austral summers of 2016 \((n = 11)\), 2018 \((n = 2)\), and spring of 2019 \((n = 2)\) as the animals were encountered along the cruise track of the MV SA Agulhas II (2016, 2019) and RV Polarstern (2018, Alfred-Wegener-Institut Helmholtz-Zentrum für Polar- und Meeresforschung, 2017). An A-frame net, consisting of two stainless steel poles, hinged at the front end, with a nylon net in between, were used to capture the seals. After restraining the seal, a small hole was cut into the net where the top of the seal’s head was located, to place the satellite transmitter. Quick-setting Araldite Epox resin (AW2101/HW2951) was used to glue the transmitter to the seals’ heads. The capture and tagging procedure followed Arcalís-Planas et al. (2015) and animals were restrained for a maximum of 1 h. We determined the sex of the animals, and took standard length and girth measurements (Bonner and Laws, 1993). The age of individuals is unknown. The standard measures are not exactly comparable to the NARE 2000/01 data where animals were anaesthetized and therefore fully relaxed and extended (Blix and Nordøy, 2007), but are comparable to those of Arcalís-Planas et al. (2015). During the 2015/2016 summer voyage (hereafter S55) six SPOT6 and five Splash-series tags (Wildlife Computers, Redmond, WA, United States) Argos-linked (CLS, Toulouse, France) satellite trackers were deployed. During summer 2018 (hereafter PS111) two SPLASH10-309A tags, and during SCALE in spring 2019, two SPOT6 tags, were deployed. Satellite transmissions from the satellite tags were not duty cycled and were allowed data to be transmitted every day for all hours of the day, and were restricted to 500 accumulated transmissions per day. Transmissions were paused when the animal was hauled out for longer than 12 h, but allowed to transmit again if the haul out lasts longer than 8 days. Transmissions relayed data that were collected the previous 2 days. Methods on the capturing, immobilisation and deployment of satellite trackers during the Norwegian Ross seal study from 2001 to 2002 are in Blix and Nordøy (2007). The Norwegian dataset consists of Argos-collected satellite tracking data from 10 individuals fitted with SDR-T16 trackers (Wildlife Computers, Redmond, WA, United States).

State-Space Models for Basic Filtering
Location data collected through the ARGOS satellite system contain intrinsic errors. To account for this, we fitted a two-state, behaviourally switching, state-space model to individual tracks (Jonsen et al., 2005; Jonsen, 2016). This procedure filtered erroneous location estimates and provided interpolated tracks with estimated locations at 3 h time intervals. This time step was based on a combination of the median number of Argos location points per day (~30 points per day) and following logic on available temporal resolution of the environmental covariates (i.e., it does not make sense to have 30 points per day compared with, for example, only two values per day of sea-surface temperatures). Bayesian State-space models were fitted using Markov chain Monte Carlo in "rjags" (Plummer, 2016), via the "bsam" library (Jonsen et al., 2005; Jonsen, 2016) implemented in R (R Core Team, 2020). Two Markov chains were run in parallel, each of 55,000 iterations, only using every 50th value, while the first 10,000 values (i.e., burn-in) were excluded. Diagnostic plots were used to assess converging and appropriate mixing of the two Markov chains (Jonsen et al., 2013). Apart from filtering and interpolating the tracking data, state-space models also identify two hidden states in the movement data and classify location estimates into either “travelling” and “area-restricted search” behaviour, which is based on the animal’s swimming speed and turning angle (Jonsen et al., 2013). Animals travelling between points are assumed to swim faster and in a straight line to cover a large area in low detail, whereas animals searching, hunting or foraging, are expected to swim slower and turn more often to cover a small area in detail, making area-restricted search a proxy for prey searching or hunting behaviour (e.g., Fauchald and Tveraa, 2003; Patterson et al., 2008).

Kernel Density Estimation
We calculated 50% kernel utilisation distributions (i.e., core habitat use; Worton, 1989) for each individual split between the austral seasons (Summer: October–March; Winter: April–September) using the R library “adehabitatHR” (Calenge, 2006). H-values were selected using the ad hoc method (Silverman, 1986). Kernel densities were plotted for illustrative purposes and data exploration.

Environmental Correlates
We used 16 remotely sensed environmental covariates to describe the habitat use of Ross seals (Table 1 and

| Environmental variable (abbreviation) | Unit |
|--------------------------------------|------|
| Bathymetry (bathym)                  | m    |
| Ocean floor slope (slope)            | °    |
| Sea surface temperature (sst)        | °C   |
| Sea surface temperature anomalies (sstA) | °C  |
| Sea surface temperature gradient (sst_grad) | °  |
| Sea surface height anomalies (sshA)  | m    |
| Sea surface height gradient (ssh_grad) | °  |
| Mixed layer depth climatology (mld_clima) | m  |
| Geostrophic current velocity (currmag) | cm/s |
| Eddy kinetic energy (eke)            | cm²/s² |
| Wind magnitude (windmag)             | m/s  |
| Distance to ice edge (disticeedge)   | m    |
| Sea ice concentration (ice)          | %    |
| Sea ice concentration standard deviation (ice_sd) | NA  |
| Vertical mixing (vmix)               | w    |
| Vertical mixing standard deviation (vmix_sd) | NA  |

1Complete information of all variables can be found in the Supplementary Files.
Supplementary Table 2). Environmental variables chosen for the habitat modelling are known to affect marine predator foraging behaviour and are commonly used to model their habitat use (Raymond et al., 2015; Reisinger et al., 2018; Wege et al., 2019; Hindell et al., 2020). Variables were extracted from the Australian Antarctic Data Centre using the R library “raster” (Summer, 2015) and processed further if needed using R library “raster” (Hijmans, 2016). We matched the date and time of each seal location to the nearest environmental data in space and time for the dynamic environmental covariates (Supplementary Table 2). To avoid the inherent collinearity among environmental variables, we filtered for the most informative set by calculating variance inflation factors for the entire Weddell Sea region and immediately adjoining waters to the east and west (i.e., 40°S to 80°S; 80°W to 80°E) using the R library “fmsb” (Nakazawa, 2018). Variables with a variance inflation factor larger than 10 were excluded, because this is a good indication of strong collinearity (Nakazawa, 2018).

Species Distribution Modelling

Species distribution models were created using a supervised machine learning technique, boostd regression trees. We separated the summer and winter portions (see Kernel density estimation above) of the tracks to create a species distribution model for each season. Boosted regression trees were chosen because they are known to perform well in predicting species’ distributions (Reisinger et al., 2018; Wege et al., 2019; Hindell et al., 2020). The state-space modelled “area restricted search” or “travelling” estimates were used as the binary response variable inherently turning these models into classification tree ensembles with a Bernoulli distribution. Given that there are not enough tracks in different years, we did not consider any annual variations. We trained models using the “caret” library in R, which employs functions from the R library “gbm” (Ridgeway, 2015; Kuhn et al., 2019). For summer tracking data we held out 30% of the data (test data, nsummer = 3,241; nwinter = 4,784) with an equal number of area-restricted search and travelling points in each test data set. The remaining training data (nsummer = 11,669; nwinter = 11,163) was unbalanced between the number of “area-restricted search” and “travelling” points (area-restricted searchsummer = 7,835 vs. travellingsummer = 3,834; area-restricted searchwinter = 11,524 vs. travellingwinter = 2,384). We made use of a bootstrapped modelling approach. The bootstraps involved running 500 boosted regression tree models independently, using random sub-sample of 2000 area-restricted search and travelling points, respectively, which was repeated for summer and winter data. This method balanced the data sets, which reduces overfitting of models and the amount of spatial-autocorrelation that is inherent in tracking data (Hijmans, 2012). Each bootstrap model was tuned respectively using the “tuneGrid” function in the “caret” library, through compiling a range of candidate models and choosing the best model and optimal set of hyperparameters based on the lowest Area Under the Curve (AUC) value from the Random Operator Curve (ROC). Candidate models for each bootstrap made use of a 10-fold cross-validation approach. We calculated the out-of-bag AUC value as the goodness-of-fit measure for each of the bootstrapped models using the 30% hold-out test data.

To generate a prediction map of potential foraging habitat of Ross seals we calculated a seasonal mean value per grid square at the same spatiotemporal resolution applied to each environmental variable used in each of the model bootstraps (Reisinger et al., 2018; Wege et al., 2020). We predicted potential foraging habitat of Ross seals for the entire Weddell and adjacent seas and further to 80° easterly and westerly longitudes and from the coastline to 30°S Latitude and beyond for each bootstrap for summer and winter data, respectively using the “predict” function in the R library “raster” (Hijmans, 2016). Using the out-of-bag AUC value of each bootstrap as the weight of each bootstrap, we calculated a weighted mean average probability of area-restricted search behaviour of the 500 bootstraps using the “weighted mean” function in R library “raster” (Hijmans, 2016). The relative influence (%) of each environmental predictor variable was calculated for each bootstrap for the summer and winter ensembles and averaged across bootstraps. Partial dependence plots of the predictor variables, for each of the bootstraps were created in R library “pdp” (Greenwell, 2017) and averaged to create a mean ± standard deviation partial dependence plot.

RESULTS

Of the 25 satellite tags deployed across the four expeditions, 19 satellite tags successfully transmitted data for >5 days and produced 32,377 location estimates (Supplementary Table 1). Ross seals immediately travelled north after deployment and completing their annual moult (February and March) during minimum ice extent, and remained in open water just south of the Polar Front (defined by Orsi et al. (1995); Figure 1). In the winter, Ross seals migrated south toward the Marginal Ice Zone (i.e., the areas directly inward and outward from the ice boundary. They spent most of the winter in the open ocean within 500 km of the ice-edge and moved northward with the expanding sea-ice edge (Figure 1). Between September and November, assumed breeding individuals (n = 6 females) entered the consolidated pack-ice further south again to pup and mate, after which they travelled north again in December. In January, Ross seals travelled back into the remaining consolidated pack-ice against, but within 50 km of the Antarctic ice shelf, to moult (Figure 1). In summer, Ross seals spent on average 68.8% (sd: ±23.2%) in the open ocean (range: 35.0–100%), and in winter Ross seals spent on average 65.7% (sd: ±27.9%) in the open ocean (range: 23.3–100%).

Highest habitat use, from the kernel density analyses, was largely the same in summer and winter, in the open ocean and away from the Antarctic continent, but south of the Polar Front (Figure 2). The large area of the kernel is a product of their south–north movements throughout the year. The winter core kernel area shifted eastward by ∼15°. The most noticeable seasonal difference in core habitat use was the summer moult haul-out close to the Antarctic continent (Figure 2).
**FIGURE 1** State-space modelled location estimates for each of the 19 Ross seals in the King Haakon VII Sea that were used to model important foraging habitat in the Weddell Sea and surrounding areas. Colours represent the four different expeditions: orange: NARE 2000/01; blue: S55 2015/16; pink: PS111 2018; green: Scale 2019. The black line indicates the position of the Polar Front (Orsi et al., 1995) and the grey lines are GEBCO 1 arc-min bathymetrical contours (IOC et al., 2003). Inset: The position of this map in relation to the Antarctic continent.

**FIGURE 2** The summer (left) and winter (right) percentage utilisation distributions for 19 Ross seals are shown in relation to the Polar Front (black line) (Orsi et al., 1995) and the Antarctic Continent. The grey lines are GEBCO 1 arc-min bathymetrical contours (IOC et al., 2003).
Boosted regression tree model performance (AUC) was high among bootstraps, with summer models' mean ± standard deviation AUC = 0.90 ± 0.002 (range: 0.89–0.90) and winter models' mean ± standard deviation AUC = 0.93 ± 0.003 (range: 0.92–0.94). The predicted area-restricted search habitat suitability maps for summer (Figures 3A,B) and winter (Figures 3C,D) illustrate the seasonal differences between Ross seal potential foraging habitat and how this expands in summer and contracts in winter. Mean importance of predictor variables changed between seasons (Figure 4). During summer the top six variables based on mean relative influence (%) were sea surface temperature (sst; °C), ice concentration standard deviation (ice_sd), mixed-layer depth summer climatology (mld_clima; m), bathymetry (bathym; m), distance to the ice edge (dist_iceedge; m), and sea-surface height anomalies (sshA; m) (Figure 4A). The top six variables in winter based on mean relative influence (%) were the distance to the ice edge (dist_iceedge; m), standard deviation of ice concentration (ice_sd), sea surface temperature (sst; °C), sea surface height anomalies (sshA; m), mixed-layer depth winter climatology (mld_clima; m), and current magnitude (currmag; cm.s⁻¹) listed in order of decreasing importance.

Partial dependence plots show that in summer, area-restricted search probability peaked between 0 and 1.8°C and increased almost linearly with an increasing ice concentration standard deviation (Figure 5). Area-restricted search probability was highest at a mixed-layer depth of 20–40 m, increasing again at mld_clima above 80 m (Figure 5), while area-restricted search probability decreases as ocean depth became shallower up to 2000 m, but then peaked again between 1000 and 1800 m deep—just off the continental slope. Area-restricted search probability decreased the further seals moved away from the ice edge, with highest probabilities at 1500–2000 km away from the ice edge, whereas area-restricted search probability was highest between 0.025 and 0.075 m sshA, but was highly variable for values below 0.05 m (Figure 5). In winter, the relationship between area-restricted search and the distance to the ice edge was bimodal, peaking at 400 m and >1500 km away (Figure 6). Similar to summer, area-restricted search probability increased with increasing ice_sd and was highest at sea-surface temperatures between −1 and 2°C (Figure 6). At sshA values smaller than 0.04 m and larger than 0.08 m seals were more likely to display area-restricted search behaviour but the likelihood was also more

![Figure 3](image-url)
variable within these ranges. The mixed-layer depths where area-restricted search was likely was deeper than in summer (100–130 m), and was highest where current magnitude was 0 and decreases with increasing current speeds (Figure 6).

**DISCUSSION**

Conservation practices skewed toward certain species, age-classes and sexes can result in bias and ignore underrepresented species (e.g., McIntyre, 2014). The Ross seal is arguably one of the most data-deficient species in the Southern Ocean. In this study, we combine all the available data for Ross seals tracked in the eastern Weddell Sea and adjoining waters (n = 19) to create the first ever species distribution model. Environmental variables associated with Ross seal foraging areas in the summer and winter are used to identify the drivers of their movements during the unique pelagic phase. Habitat modelling provides novel insight into the year-round and large-scale distribution of the species and the potential impacts of climate change.

**Distribution and Diet**

Instrumented Ross seals travelled away from the pack-ice and spend most of their time in the open ocean (Blix and Nordøy, 2007; Arcalís-Planas et al., 2015, this study) and traversed the Atlantic sector of the Southern Ocean within a very narrow longitudinal band four times a year (Figure 1 and Supplementary Animations). The likelihood of area-restricted search behaviour in summer and winter covaried with ice concentration variability (Figures 5, 6), which serves as a proxy for sea-ice edge zones and accessibility of ice-covered areas (Wege et al., 2020). The sea-ice edge zones are productive areas due to the input of nutrients in the water column as the sea ice melts, which promote phytoplankton blooms which in turn trigger grazers and other mid-level and higher-order predators to aggregate forage (Nicol, 2006; Arrigo et al., 2008; Riekkola et al., 2019). The thick multi-year pack-ice present in the Weddell Sea and an increased swimming distance to the open ocean likely makes the southern Weddell Sea an unsuitable habitat for Ross seals, which would explain the low habitat suitability within the Weddell Sea embayment year-round (Figure 3). Bester et al. (2020) reviewed all observational data of Ross seals and found that during their annual moult (late summer, early autumn), they are absent within the inner reaches of the Weddell Sea south of 73°S and west of 30°W. Our habitat model showed that within the Weddell Sea embayment, a small area is still likely foraging habitat (Figure 3). Bester et al. (2020) used only observational data (i.e., Ross seals hauled out to moult/rest), while in contrast our study predicts potential foraging habitat based on environmental variables. It is also important to consider that a habitat model only predicts on the variables given to it, and that there are potentially unknown or currently immeasurable variables that dictate Ross seal foraging habitat.
In summer, the foraging behaviour of Ross seals can be split into two stages: a period close to the narrow continental shelf off Queen Maud Land during the annual moult and open ocean foraging between 55° and 65°S, south of the Polar Front. Open ocean foraging happens either post-moult when seals travel north for the winter, post-breeding (November–December) when breeding individuals return from the ice, and for non-breeders that remained at their northern range throughout the summer (Figures 2, 3). This explains why Ross seal foraging probability increases as the waters become deeper (negatively correlated) and preferred to forage over abyssal water (>4000 m deep; Figure 5). However, there is a peak in the probability of area-restricted search behaviour between 1000 and 1500 m depth (Figure 5), which is considered the continental slope and where Ross seals forage during their annual moult (Southwell, 2005; Blix and Nordøy, 2007; Arcalís-Planas et al., 2015).

Stomach content analyses from earlier investigations suggested that Ross seals sampled in January off Queen Maud Land predominantly prey on Antarctic silverfish and squid (Skinner and Klages, 1994). Antarctic silverfish are pelagic and only occur close to the Antarctic continent, on the continental shelf, or on the shallow slope areas in cold water temperatures ranging between (−1.75 to −2°C) (DeWitt, 1970; Hubold, 1984; DeWitt et al., 1990; Ekau, 1990; Eastman, 1993; White and Piatkowski, 1993; Kellerman, 1996; Trunov, 2001; Donnelly et al., 2004; Davis et al., 2017; La Mesa et al., 2019), and higher temperatures of up to 4°C around islands of the Scotia Arc (Mintenbeck and Torres, 2017). Ross seals are likely only forage on Antarctic silverfish during their annual moult, while over
Wege et al. Habitat Suitability of Ross Seals

FIGURE 6 | Winter partial dependence plots showing how the probability of area restricted search (ARS) is influenced by the top six environmental variables in the winter models. Mean probability of ARS (solid line) ± sd (grey shaded) across the 500 bootstraps are shown. dist_iceedge, distance from the ice edge (m); ice_sd, ice concentration standard deviation; mld_clima, mixed layer depth climatology (m); sst, sea surface temperature (°C); sshA, sea surface height anomaly; currmag, current magnitude (m.s⁻¹).

the continental shelf and slope. Based on our understanding of Ross seal movements outside of the moult, these diet samples collected in January are not a reflection of the year-round diet. This agrees with stable isotope results (Rau et al., 1992; Zhao et al., 2004; Brault et al., 2019) and diving behaviour analyses, which also suggest that Ross seals are open ocean foragers that prey on mesopelagic fish and squid (Blix and Nordøy, 2007). Fisheries in the Southern Ocean primarily focus on krill (Euphausia spp.), toothfish (Dissostichus spp.) and squid (Agnew et al., 2005; Kock et al., 2007; Nicol et al., 2012; Chown and Brooks, 2019), which means Ross seals are not in direct competition with krill and toothfish fisheries. Currently, it is unknown what percentage of the Ross seal diet is comprised of cephalopods, but it is not their dominant prey species (Rau et al., 1992; Skinner and Klages, 1994; Zhao et al., 2004). Potential for conflict with cephalopod fisheries are also unknown, given the scarcity of data on cephalopod fishing in the open ocean around Antarctica (Agnew et al., 2005; Chown and Brooks, 2019).

Climate Change Implications
Siniff et al. (2008) suggested that the pelagic foraging behaviour of Ross seals makes them least susceptible to climate change compared to other Antarctic seals. They may even be considered as potential climate change winners, because the shrinking distance from the continent to the sea-ice edge in theory would require Ross seals not to swim as far to reach potential foraging grounds.
Ross seals preferred to spend time in waters between mixed layer depth and wind-driven sea-surface height anomalies. Probability of area-restricted search behaviour were the same in warm sea-surface temperature anomalies (Supplementary Figures 1, 2) during both seasons. Sea-surface temperature is a dominant predictor in most Antarctic predators’ movement behaviour (Hindell et al., 2020), and coincides with productive areas in the ocean, marine predator abundance (Block et al., 2011), and subsequent prey distribution and availability (Reisinger et al., 2018; Bestley et al., 2020).

During the summer, Ross seal area-restricted search probability decreased with an increased mixed-layer depth, while in winter area-restricted search probability increased with an increase in mixed-layer depth. The mixed-layer depth influences depth of prey aggregations and any fluctuations in the mixed-layer depth will influence how deep seals have to dive (and the amount of energy to expend) to find prey. The difference between the correlations of area-restricted search probability and mixed-layer depth across the seasons is likely linked to location of dives (continental shelf vs. off-shelf) and a general expected increase in mixed-layer depths during winter months (e.g., Sallée et al., 2010). In areas of the ocean covered in sea-ice, melting ice causes freshening of the water column and the mixed-layer depth to shoal (Sallée et al., 2013; Meijers, 2014). However, how the mixed-layer depth is likely to respond to climate change is not reliably known because of the inability of climate models to accurately represent stratification in the winter months, and are often biased towards shallower mixed layers (Sallée et al., 2013), which means climate models cannot always accurately predict what will happen to mixed-layer depths throughout the Southern Ocean. The interactions between processes that can affect stratification—primarily through momentum and buoyancy fluxes—are complex, and so how the mixed-layer depth will be affected by climate change differs by region and proximity to continental shelf. If, as suggested by climate models, the mixed layer depth shoals, it could potentially be beneficial to diving Ross seals and reduce the amount of energy expended while foraging (Reisinger et al., 2018; Bestley et al., 2020).

In summer, Ross seals displayed more area-restricted search behaviour around 500 and 1500 km away from the ice edge. This likely results from the seasonality in the expanding ice-edge: during the early winter months (April and May) Ross seals are still foraging at sea at their maximal range away from Antarctica and as the ice-edge expands, they move southward toward the ice edge, then following the growing seasonal ice (Animations in the Supplementary Files). If the ice edge recedes, this would mean that Ross seals will have to swim smaller distances to reach the open ocean and to return to the ice edge, which could potentially be beneficial to them and reduce the amount of energy expended swimming to foraging areas (Siniff et al., 2008). In summer, there was more predicted suitable foraging habitat available compared to winter for Ross seals (Figure 3). This is because of the seasonal pack-ice growth in winter and Ross seals’ preference for open ocean foraging, further indicating that receding ice edge under future climate scenarios will reduce the swimming distance to available foraging habitat, and that potentially more foraging habitat will become available. However, in the Weddell Sea and Ross Sea regions of the Southern Ocean the seasonal ice edge was expanding and sea-surface temperatures decreasing (but summer ice extent recently recorded to decrease (Turner et al., 2020)); while the Amundsen Sea ice losses are increasing along with rising sea-surface temperatures (Zhang, 2007; Bintanja et al., 2013; Mehl et al., 2016). These contrasting ice-dynamics will result in different responses in foraging behaviour by Ross seals to climate change.

Despite the impacts on their energy expenditures (swimming distance and diving depth), their preferred foraging grounds are also influenced through changing sea-surface temperatures, increased wind and current magnitudes, increased wave height, and poleward shifting wind (Constable et al., 2014). How these changes will affect the distribution of prey is critical to understand how Ross seals will respond. Myctophids in the Southern Ocean are predicted to undergo species-specific shifts in their distribution, but overall most species shift toward the poles (Freer et al., 2019), which would require sub-Antarctic predators to travel south from their sub-Antarctic islands toward the ice to forage (Labrousse et al., 2017; Reisinger et al., 2018). Ross seals travel northward (this study), and a southward migration of prey would reduce travel distances and could be advantageous. Cephalopods, which also form a component of the Ross seal diet (Skinner and Klages, 1994), are unlikely to be affected by sea-ice changes, but they are vulnerable to mesoscale fluctuations and ocean acidification that are affecting entire ecosystems (Rodhouse, 2013; Xavier et al., 2018). How fish and squid species will respond to climate change and increased fishing pressures within mesopelagic ecosystems is largely still unknown and most studies are focussed on Antarctic krill (Euphausia superba) (Murphy et al., 2007, 2013).

Crab eater seals (Lobodon carcinophaga) rely on Antarctic krill for 90% of its diet (Hückstädt et al., 2012) while Antarctic fur seals (Arctocephalus gazella) breeding in the south Atlantic also...
predominantly prey on krill (Reid and Arnould, 1996). Dietary specialisation makes both species susceptible to any variances and shifts in prey distribution and abundance and are not able to buffer climate fluctuations as well as other Antarctic seals and seabirds (Forcada et al., 2008; Hückstädt et al., 2020). Generalist predators are more likely to switch between prey species are likely to fair better than specialist foragers (e.g., Angermeier, 1995; Terraube et al., 2011), like Weddell seals (Leptonychotes weddellii) for example, whose diet varies among many species of fishes, cephalopods, and crustaceans (Burns et al., 1998; Lake et al., 2003). But, due to Weddell seals’ association with fast-ice for breeding, they are likely to lose breeding habitat and increase in competition with emperor penguins (Aptenodytes forsteri) (LaRue et al., 2019; Jenouvrier et al., 2020; Trathan et al., 2020). Here, we only considered Ross seals foraging movements and potential energy expenditure in line with climate shifts. Like southern elephant seals, Ross seals are generalists and are free-roaming pelagic foragers, who can travel great distances to find prey, which is why elephant seals are also suggested to likely benefit from climate change (Costa et al., 2010). Unlike elephant seals who breed on sub-Antarctic islands, Ross seals breed on the pack-ice like crabeater and leopard (Hydrurga leptonyx) seals. Despite the energetic gains from swimming shorter distances to foraging locations, they are most likely to suffer breeding habitat loss, which might increasing breeding substrate competition with other pack-ice breeders, similar to Weddell seals and emperor penguins (LaRue et al., 2019). Southern elephant seals, who also prey on squid and fish (Daneri and Carlini, 2002; Van den Hoff et al., 2003) and can dive to depths of over 2000 m, beyond the diving reach of the Ross seal (Blix and Nordøy, 2007; McIntyre et al., 2010), are likely to compete with Ross seals for prey resources.

CONCLUSION

Overall, these results suggest that climate change will likely alter Ross seals’ foraging behaviour, but it is not clear whether it will be for the better. They will expend less energy while diving due to shoaling mixed-layers, and swimming shorter distances in the winter due to the contracting sea-ice, but in summer post-breeding individuals might swim further distances to reach open water. Their responses will be regionally specific to climatic shifts, rather than uniform. However, toward the end of the century under the “business as usual” models, sea-ice is predicted to decline around the entire Antarctic continent (Gutt et al., 2015) and temperatures of all water masses are predicted to increase (Meijers, 2014).

From a conservation perspective, currently an entire ecosystem assessment of the overall ecosystem effects of climate change in the Southern Ocean is still lacking and has been identified as one of the priority research questions for future research (Kennicutt et al., 2014; Gutt et al., 2015), especially given the planned marine protected area in the Weddell Sea (Tesche et al., 2020). It is therefore unknown how low trophic levels through to top predators will collectively respond to climate shifts and cumulatively affect each other. With expanding fisheries (Brooks, 2013) future conflict and competition between Ross seals and fisheries cannot be ruled out. Even so, it appears that the Ross seal will be the seal species whose foraging habitat is least affected by climate change. We only considered Ross seal foraging habitat and no other aspects of their life-history and their dependence on seasonal pack-ice for breeding substrate makes them vulnerable to climate change induced breeding habitat-loss (see above). Ultimately, several aspects of their life-histories remain unknown, which makes effective conservation planning for this data deficient species exceedingly difficult.

DATA AVAILABILITY STATEMENT

The original contributions presented in the study are publicly available. This data can be found here: https://doi.pangaea.de/10.1594/PANGAEA.929949.

ETHICS STATEMENT

The animal study was reviewed and approved by the University of Pretoria Animal Ethics Committee (Number EC082–15). The study was cleared under South African Department of Environmental Affairs Permit 04/2015–2019, pursuant to the provisions of Article 3 of the Protocol on Environmental Protection to the Antarctic Treaty, and Annex II and Annex V (Article 10(2)).

AUTHOR CONTRIBUTIONS

MB, HB, and MW contributed funding and collected the new data. AB and EN contributed previously collected data. MW analysed the data and wrote the manuscript. LB provided invaluable expertise on the oceanographic interpretations. All authors edited and contributed intellectual input to the final manuscript.

FUNDING

The National Research Foundation (NRF) South African National Antarctic Programme (SANAP) Grant Number 93088. The Alfred Wegener Institute for Polar and Marine Research through the Helmholtz Association Research Programme Polar Regions and Coasts in the changing Earth System II (PACES II), Topic 1, WP 6. The work on board RV Polarstern refers to Grant-No. AWI_PS111_00. The Norwegian Research Council and the Norwegian Polar Research Institute as part of the NARE 2000/2001 expedition for financial and logistical support. The Department of Environment, Forestry and Fisheries for logistical support within South African National Antarctic Programme (SANAP) and the Department of Science and Technology, through the National Research Foundation (NRF), for funding this project.
ACKNOWLEDGMENTS

Dr. Martin Postma, Dr. Nico Lübcker, and Wiam Haddad are thanked for their assistance in the field. The Officers and Crew of the MV SA Aquilas II and RV Polarstern extended every possible courtesy to us in support of our research objectives. Chief Scientists of the respective voyages, Dr. Thato Mtshali, Dr. Mike Schröder, and Dr. Tommy Ryan-Keogh are thanked for their support and the Department of Environment Affairs’ Co-ordinating Office (DCO) and Deputy DCO, for facilitation.

We acknowledge support by the Open Access Publication Funds of Alfred-Wegener-Institut Helmholtz-Zentrum für Polar- und Meeresforschung.

SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fmars.2021.659430/full#supplementary-material

REFERENCES

Alfred-Wegener-Institut Helmholtz-Zentrum für Polar- und Meeresforschung. (2017). Polar research and supply vessel POLARSTERN operated by the alfred-wegener-institute. J. Large Scale Res. Facil. 3: A119. doi: 10.17815/jlsrf-3-163

Agnew, D. J., Hill, S. L., Beddington, J. R., Purchase, L. V., and Wakeford, R. C. (2005). Sustainability and management of Southwest Atlantic squid fisheries. Bull. Mar. Sci. 76, 579–593.

Aungermeier, P. L. (1995). Ecological attributes of extinction prone species: loss of freshwater species of fishes of Virginia. Conserv. Biol. 9, 143–158. doi: 10.1046/j.1523-1739.1995.0091014.x

Arcailis-Planas, A., Sveegaard, S., Karlsson, O., Harding, K. C., Wåhlin, A., Harkonen, T., et al. (2015). Limited use of sea ice by the Ross seal (Ommatophoca rossii), in Amundsen Sea, Antarctica, using telemetry and remote sensing data. Polar Biol. 38, 445–461. doi: 10.1007/s00300-014-1602-y

Arrigo, K. R., van Dijken, G. L., and Bushinsky, S. (2008). Primary production in Pacific Ocean sectors of the Southern Ocean in CMIP5 models: historical bias, forcing response, and state dependence. J. Geophys. Res. Ocean. 113:C08004.

Béring, C., and Weimerskirch, H. (2001). Emperor penguins and climate change. Nature 413, 183–186. doi: 10.1038/35075554

Béring, C., Weimerskirch, H., Guinet, C., and Jouventin, P. (2000). Effect of sea-ice extent on adult survival of an Antarctic top predator: the snow petrel Pagodroma nivea. Oecologia 125, 483–488. doi: 10.1007/3-540-30040-4_48

Bester, M. N., Wege, M., Oosthuizen, W. C., and Bornemann, H. (2020). Ross seal distribution in the weddel sea: fact and fallacy. Polar Biol. 43, 35–41. doi: 10.1007/s00300-019-02610-4

Bester, S., Ropert-Couzet, Y., Bengston Nash, S., Brooks, C. M., Cotté, C., Dewar, M., et al. (2020). Marine ecosystem assessment for the southern Ocean: birds and marine mammals in a changing climate. Front. Ecol. Environ. 8:56936. doi: 10.1389/fev.2020.56996

Bintanja, R., Van Oldenborgh, G. J., Drijfhout, S. S., Wouters, B., and Katsman, R., et al. (2013). Assessment of surface winds over the Atlantic, Indian, and Pacific Ocean sectors of the Southern Ocean in CMIP5 models: historical bias, forcing response, and state dependence. J. Geophys. Res. Atmos. 118, 547–562. doi: 10.1002/jgrd.50153

Brooks, C. M. (2013). Competing values on the antarctic high seas: CCAMLR and the challenge of marine-protected areas. Polar Biol. 36, 277–292. doi: 10.1007/11820896X.2013.854597

Burns, J. M., Trumble, S. J., Castellini, M. A., and Testa, J. W. (1998). The diet of Weddell seals in McMurdo sound, antarctica as determined from scat collections and stable isotope analysis. Polar Biol. 19, 272–282. doi: 10.1007/s00300005245

Calenge, C. (2006). The package adehabitat for the R software: a tool for the analysis of space and habitat use by animals. Ecol. Model. 197, 516–519. doi: 10.1016/j.ecolmodel.2006.03.017

Chen, I.-C., Hill, J. K., Olheimuller, R., Roy, D. B., and Thomas, C. D. (2011). Rapid range shifts of species associated with high levels of Climate Change. Science 333, 1024–1026. doi: 10.1126/science.1206432

Chown, S. L., and Brooks, C. M. (2019). The state and future of antarctic environments in a global context. Ann. Rev. Environ. Resour. 44, 1–30. doi: 10.1146/annurev-environ-101718-033236

Constable, A. J., Melbourne-Thomas, J., Corney, S. P., Arrigo, K. R., Barbraud, C., Barnes, D. K. A., et al. (2014). Climate change and Southern Ocean ecosystems I: how changes in physical habitats directly affect marine biota. Glob. Chang. Biol. 20, 3004–3025. doi: 10.1111/gcb.12623

Convey, P., and Peck, L. S. (2019). Antarctic environmental change and biological responses. Sci. Adv. 11: eaaz0888. doi: 10.1126/sciadv.aaz0888

Costa, D. P., Hückstädt, L. A., Crocker, D. E., McDonald, B. I., Goebel, M. E., and Fedak, M. A. (2010). Approaches to studying climatic change and its role on the habitat selection of antarctic pinnipeds. Integr. Comp. Biol. 50, 1018–1030. doi: 10.1093/icb/icq054

Cristofari, R., Liu, X., Bonadonna, F., Cherel, Y., Pistorius, P. A., Le Maho, Y., et al. (2018). Climate-driven range shifts of the king penguin in a fragmented ecosystem. Nat. Clim. Chang. 8, 245–251. doi: 10.1038/s41558-018-0084-2

Daneri, G. A., and Carlini, A. R. (2002). Fish prey of southern elephant seals, Mirounga leonina, at King George Island. Polar Biol. 25, 739–743. doi: 10.1007/s00300-002-0408-5

Davis, L. B., Hofmann, E. E., Klinck, J. M., Piñones, A., and Dinniman, M. S. (2017). Distributions of krill and Antarctic silverfish and correlations with environmental variables in the western Ross Sea, Antarctica. Mar. Ecol. Prog. Ser. 584, 45–65. doi: 10.3354/meps12347

de la Mare, W. K. (1997). A abrupt mid-twentieth century decline in Antarctic sea-ice extent from whaling records. Nature 389, 57–60. doi: 10.1038/37956

DeWitt, H. (1970). “The character of the midwater fish fauna of the ross sea, antarctica,” in Antarctic Ecology, ed. M. W. Holdgate (London: Academic), 305–314.

DeWitt, H., Heemstra, P., and Gon, O. (1990). “Nototheniidae,” in Fishes of the Southern Ocean, eds O. Gon and P. C. Heemstra (Grahamstown: JLB Smith Institute of Ichthyology), 279–331.

Donnelly, J., Torres, J. J., Sutton, T. T., and Simonelli, C. (2004). Fishes of the eastern Ross Sea, Antarctica. Polar Biol. 27, 637–650. doi: 10.1007/s00300-004-0632-2

Eastman, J. T. (1993). Antarctic Fish Biology: Evolution in a Unique Environment. San Diego, CA: Academic Press.

Ekau, W. (1990). Demersal fish fauna of the Weddell Sea. Antarctica. Antarct. Sci. 2, 129–137. doi: 10.1017/S0954102090000165
Freer, J. J., Genner, M. J., Tarling, G. A., Collins, M. A., and Partridge, J. C. (2019).
Forcada, J., Trathan, P. N., and Murphy, E. J. (2008). Life history buffering
Fauchald, P., and Tveraa, T. (2003). Using first-passage time in the analysis of area-
Wege et al. Habitat Suitability of Ross Seals
Gutt, J., Bertler, N., Bracegirdle, T. J., Buschmann, A., Comiso, J., Hosie, G., et al.
Hindell, M. A., McMahon, C. R., Bester, M. N., Boehme, L., Costa, D. P.,
raster: Geographic Data Analysis and Modeling. R
Hijmans, R. J. (2012). Cross-validation of species distribution models: removing spatial sorting bias and calibration with a null model. Ecology 93, 679–688.
doi: 10.1890/11-0826.1
Hijmans, R. J. (2016). raster: Geographic Data Analysis and Modeling. R
Hindell, M. A., Sumner, M., Bestley, S., Bravington, M. V., Patterson, T. A., Pedersen, M. W., et al. (2013). State-space models for bio-loggers: a methodological road map. Deep Sea Res. Part II Top. Stud. Oceanogr. 88–89, 34–46. doi: 10.1016/j.
doi: 2012.07.008
Jonsen, I. D., Fleming, J. M., and Myers, R. A. (2005). Robust state-space modeling of animal movement data. Ecology 86, 2874–2880. doi: 10.1890/04-1852
Karl, T. R., and Tremberth, K. E. (2003). Modern global climate change. Science 302, 1719–1723.
Kellerman, A. K. (1996). “Midwater fish ecology,” in Foundations for Ecological Research West of the Antarctic Peninsula, Antarctic Research Series, Vol. 76, eds R. M. Ross, E. E. Hofman, and L. N. Quetin (Washington D.C.: American Geophysical Union), 231–256. doi: 10.1029/97pp00231
Kennon, C. R., Griffen, K. E., Stirling, I., and Cullen, J. J. (2016). Six priorities for Antarctic adaptation. Phil. Trans. R. Soc. Lond. B. Biol. Sci. 371, 1–14. doi: 10.1098/rstb.2015.0386
Koch, K.-H., Reid, K., Croxall, J. P., and Nicol, S. (2007). Fisheries in the Southern Ocean: an ecosystem approach. Philos. Trans. R. Soc. Lond. B. Biol. Sci. 362, 2333–2349. doi: 10.1098/rstb.2006.1954
Kuhn, M., Wing, J., Weston, S., Williams, A., Keefer, C., Engelhardt, A., et al. (2019). caret: Classification and Regression Training. R Package Version 6.0-83. Available online at: https://CRAN.R-project.org/package=caret (accessed February 1, 2020).
La Mesa, M., Piepennburg, D., Pineda-Metz, S. E. A., Riginella, E., and Eastman, J. T. (2019). Spatial distribution and habitat preferences of demersal fish assemblages in the southeastern Weddell Sea (Southern Ocean). Polar Biol. 42, 1025–1040. doi: 10.1007/s00300-019-02495-3
Labrouse, S., Argeret, F., Solow, A. R., Barbraud, C., Bost, C. A., Sallée, J. B., et al. (2019). First odyssey beneath the sea ice of juvenile emperor penguins in East Antarctica. Mar. Ecol. Prog. Ser. 609, 1–16. doi: 10.1134/meps12831
Labrouse, S., Sallée, J. B., Fraser, A. D., Massom, R. A., Reid, P., Hobbs, W., et al. (2017). Variability in sea ice cover and climate elicit sex specific responses in an Antarctic predator. Sci. Rep. 7:43236. doi: 10.1038/srep43236
Labrouse, S., Vacqué-Garcia, J., Heerah, K., Guinet, C., Sallée, J. B., Authier, M., et al. (2015). Winter use of sea ice and ocean water mass habitat by southern elephant seals: the length and breadth of the mystery. Prog. Oceanogr. 137, 52–68. doi: 10.1016/j.pocean.2015.05.023
Lake, S., Burton, H., and van den Hoff, J. (2003). Regional, temporal and fine-scale spatial variation in Weddell seal diet at four coastal locations in east Antarctica. Mar. Ecol. Prog. Ser. 254, 293–305. doi: 10.3354/meps254293
LaRue, M. A., Salas, L., Nur, N., Ainley, D. G., Stammerjohn, S. E., Barrington, L., et al. (2019). Physical and ecological factors explain the distribution of Ross Sea Weddell seals during the breeding season. Mar. Ecol. Prog. Ser. 612, 193–208. doi: 10.3354/meps12877
McIntyre, T. (2014). Trends in tagging of marine mammals: a review of marine mammal biologging studies. Afr. J. Mar. Sci. 36, 409–422. doi: 10.2989/1814232X.2014.976655
McIntyre, T., de Bruyn, P. J. N., Ansorge, I. J., Bester, M. N., Bornemann, H., Plotz, J., et al. (2010). A lifetime at depth: vertical distribution of southern elephant seals in the water column. Polar Biol. 33, 1037–1048. doi: 10.1007/s00300-010-0782-3
McMahan, C. R., and Burton, H. R. (2005). Climate change and seal survival: evidence for environmentally mediated changes in elephant seal. Mloungi leonina, pup survival. Proc. R. Soc. Lond. B. 272, 923–928. doi: 10.1098/rspb.2004.3038
Meehi, G. A., Arblaster, J. M., Bitz, C. M., Chung, C. T. Y., and Teng, H. (2016). Antarctic sea-ice expansion between 2000 and 2014 driven by tropical Pacific decadal climate variability. Nat. Geosci. 9, 590–595. doi: 10.1038/ngeo2751
Meehi, G. A., Stocker, T. F., Collins, W. D., Friedlingstein, P., Gaye, T., Gregory, J. M., et al. (2007). “Global climate projections,” in IPCC, 2007: Climate Change 2007: The Physical Science Basis. Contribution of Working Group I to the Fourth Assessment Report of the Intergovernmental Panel on Climate Change, eds S. Solomon, D. Qin, M. Manning, Z. Chen, M. Marquis, K. B. Averyt, et al. (Cambridge: Cambridge University Press), 747–846.
Meijers, A. J. S. (2014). The Southern ocean in the coupled model intercomparison project phase 5. Philos. Trans. R. Soc. A Math. Phys. Eng. Sci. 372:20130296. doi: 10.1098/rsta.2013.0296
Meijers, A. J. S., Meredith, M. P., Murphy, E. J., Chambers, D. P., Belcher, M., and Young, E. F. (2019). The role of ocean dynamics in king penguin range estimation. Nat. Clim. Chang. 9, 120–121. doi: 10.1038/s41558-018-0388-2
Meredith, M., Sommerkorn, M., Cassotta, S., Derksen, C., Ekyakim, A., Hollowed, A., et al. (2019). “International panel of climate change scientific report: ocean and cryosphere,” in IPCC Spec. Rep. Ocean Cryosph. a Chang. Clim. - Summ.
Patterson, T. A., Thomas, L., Wilcox, C., Ovaskainen, O., and Matthiopoulos, J. (2018). State-space models of individual animal movement. *Trends Ecol. Evol.* 33, 1239ñ1243. doi: 10.1016/j.tree.2017.10.009

Péron, C., Weimerskirch, H., and Bost, C.-A. (2012). Projected poleward shift of king penguins' (*Aptenodytes patagonicus*) foraging range along the Crozet Islands, southern Indian Ocean. *Proc. R. Soc. B Biol. Sci.* 279, 2515ñ2523. doi: 10.1098/rspb.2011.2705

Pinsky, M. L., Worm, B., Fogarty, M. J., Sarmiento, J. L., and Levin, S. A. (2013). Marine taxa track local climate velocities. *Science* 341, 1239ñ1243. doi: 10.1126/science.1239352

Plummer, M. (2016). *rjags*: Bayesian Graphical Models using MCMC. *R* package version 4-6. Available online at: https://cran.r-project.org/package=rjags (accessed February 1, 2020).

R Core Team (2020). *R: A Language and Environment for Statistical Computing*. Vienna: R Foundation for Statistical Computing.

Rau, G. H., Ainley, D. G., Bengston, J. L., Torres, J. J., and Hopkins, T. L. (1992). 15N/14N and 13C/12C in Weddell Sea birds, seals, and fish: implications for diet and trophic structure. *Mar. Ecol. Prog. Ser.* 84, 1ñ8. doi: 10.3354/meps08401

Reisinger, R. R., Raymond, B., Hindell, M. A., Bester, M. N., Crawford, R. J. M., Orsi, H., Whitworth, T. III, and Nowlin, W. D. Jr. (1995). On the meridional extent of *Euphausia superba* along the Antarctic Peninsula and South Georgia. *J. Mar. Syst.* 109ñ110, 22ñ42. doi: 10.1016/j.jmsys.2012.03.011

Murphy, E. J., Trathan, P. N., Watkins, J. L., Reid, K., Meredith, M. P., Forcada, J., et al. (2017). Climatically driven fluctuations in Southern Ocean ecosystems. *Proc. R. Soc. B* 284, 20171180. doi: 10.1098/rspb.2017.1180

Nakazawa, M. (2018). *fmsb*: Functions for Medical Statistics Book With Some Demographic Data. R Package Version 0.6.3. Available online at: https://cran.r-project.org/package=fmsb.

Nicol, S. (2006). Krill, currents, and sea ice: *Euphausia superba* and its changing environment. *Bioscience* 56, 111ñ120. doi: 10.1641/0006-3568(2006)056[0111:kcsea]2.0.CO;2

Nicol, S., Foster, J., and Kagawuchi, S. (2012). The fishery for Antarctic krill - recent developments. *Fish Fish.* 13, 30ñ40. doi: 10.1111/j.1467-2979.2011.00406.x

Orsi, H., Whitworth, T. III, and Nowlin, W. D. Jr. (1995). On the meridional extent and fronts of the Antarctic circumpolar current. *Deep Sea Res. I Oceanogr. Res. Pap.* 42, 641ñ673. doi: 10.1016/0967-0637(95)00201-W

Patterson, T. A., Thomas, L., Wilcox, C., Ovaskainen, O., and Mathiopoulos, J. (2008). Space-space models of individual animal movement. *Trends Ecol. Evol.* 23, 87ñ94. doi: 10.1016/j.tree.2007.10.009

Prévot, I., Defaix, O., Barbraud, C., Atkinson, C., Le Maho, Y., and Ainley, D. G. (2010). Bottom-up regulation of a pole-ward migratory front of the Antarctic Circumpolar Current. *Glob. Change Biol.* 16, 2513ñ2530. doi: 10.1111/j.1365-2486.2010.01951.x

R Core Team (2020). *R: A Language and Environment for Statistical Computing*. Vienna: R Foundation for Statistical Computing.

Rau, G. H., Ainley, D. G., Bengston, J. L., Torres, J. J., and Hopkins, T. L. (1992). 15N/14N and 13C/12C in Weddell Sea birds, seals, and fish: implications for diet and trophic structure. *Mar. Ecol. Prog. Ser.* 84, 1ñ8. doi: 10.3354/meps08401

Raymond, B., Lea, M. A., Patterson, T. A., Andrews-Goff, V., Sharples, R., Chiaradia, A., Atkinson, C., Le Maho, Y., and Ainley, D. G. (2010). Bottom-up regulation of a pole-ward migratory front of the Antarctic Circumpolar Current. *Glob. Change Biol.* 16, 2513ñ2530. doi: 10.1111/j.1365-2486.2010.01951.x

R Core Team (2020). *R: A Language and Environment for Statistical Computing*. Vienna: R Foundation for Statistical Computing.

Rau, G. H., Ainley, D. G., Bengston, J. L., Torres, J. J., and Hopkins, T. L. (1992). 15N/14N and 13C/12C in Weddell Sea birds, seals, and fish: implications for diet and trophic structure. *Mar. Ecol. Prog. Ser.* 84, 1ñ8. doi: 10.3354/meps08401

Raymond, B., Lea, M. A., Patterson, T. A., Andrews-Goff, V., Sharples, R., Chiaradia, A., Atkinson, C., Le Maho, Y., and Ainley, D. G. (2010). Bottom-up regulation of a pole-ward migratory front of the Antarctic Circumpolar Current. *Glob. Change Biol.* 16, 2513ñ2530. doi: 10.1111/j.1365-2486.2010.01951.x

R Core Team (2020). *R: A Language and Environment for Statistical Computing*. Vienna: R Foundation for Statistical Computing.

R Core Team (2020). *R: A Language and Environment for Statistical Computing*. Vienna: R Foundation for Statistical Computing.
are not determined by habitat characteristics. *BMC Ecol.* 19:36. doi: 10.1186/s12898-019-0252-x

Wege, M., Salas, L., and LaRue, M. A. (2020). Citizen science and habitat modelling facilitates conservation planning for crab eater seals in the Weddell Sea. *Divers. Distrib.* 26, 1291–1304. doi: 10.1111/ddi.13120

White, M. G., and Piatkowski, U. (1993). Abundance, horizontal and vertical distribution of fish in eastern Weddell Sea micronekton. *Polar Biol.* 13, 41–53.

White, W. B., and Peterson, R. G. (1996). An Antarctic circumpolar wave in surface pressure, wind, temperature and sea-ice extent. *Nature* 380, 699–702. doi: 10.1038/380699a0

Worton, B. J. (1989). Kernel methods for estimating the utilization distribution in home-range studies. *Ecology* 70, 164–168. doi: 10.2307/1938423

Würsig, B., Thewissen, J. G. M., and Kovacs, K. M. (2018). *Encyclopedia of Marine Mammals*, 3rd Edit Edn. San Diego, CA: Elsevier Press.

Xavier, J. C., Cherel, Y., Allcock, L., Rosa, R., Sabirov, R. M., Blicher, M. E., et al. (2018). A review on the biodiversity, distribution and trophic role of cephalopods in the Arctic and Antarctic marine ecosystems under a changing ocean. *Mar. Biol.* 165, 1–26. doi: 10.1007/s00227-018-3352-9

Young, I. R., and Ribal, A. (2019). Multiplatform evaluation of global trends in wind speed and wave height. *Science* 364, 548–552. doi: 10.1126/science.aav9527

Young, I. R., Zieger, S., and Babanin, A. V. (2011). Global trends in wind speed and wave height. *Science* 332, 451–455. doi: 10.1126/science.1197219

Younger, J. L., van den Hoff, J., Wienecke, B., Hindell, M. A., and Miller, K. J. (2016). Contrasting responses to a climate regime change by sympatric, ice-dependent predators. *BMC Evol. Biol.* 16:61. doi: 10.1186/s12862-016-0630-3

Zhang, J. (2007). Increasing Antarctic sea ice under warming atmospheric and oceanic conditions. *J. Clim.* 20, 2515–2529. doi: 10.1175/JCLI4136.1

Zhao, L., Castellini, M. A., Mau, T. L., and Trumble, S. J. (2004). Trophic interactions of Antarctic seals as determined by stable isotope signatures. *Polar Biol.* 27, 368–373. doi: 10.1007/s00300-004-0598-0

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

**Copyright © 2021 Wege, Bornemann, Blix, Nordøy, Biddle and Bester.** This is an open-access article distributed under the terms of the Creative Commons Attribution License (CC BY). The use, distribution or reproduction in other forums is permitted, provided the original author(s) and the copyright owner(s) are credited and that the original publication in this journal is cited, in accordance with accepted academic practice. No use, distribution or reproduction is permitted which does not comply with these terms.