RESEARCH ARTICLE

Static species distribution models in the marine realm: The case of baleen whales in the Southern Ocean

Ahmed El-Gabbas | Ilse Van Opzeeland | Elke Burkhardt | Olaf Boebel

Ocean Acoustics Group, Alfred-Wegener-Institut (AWI), Helmholtz-Zentrum für Polar- und Meeresforschung, Bremerhaven, Germany

Correspondence
Ahmed El-Gabbas, Ocean Acoustics Group, Alfred-Wegener-Institut (AWI), Helmholtz-Zentrum für Polar- und Meeresforschung, 27570 Bremerhaven, Germany.
Email: ahmed.el-gabbas@awi.de

Funding information
Bundesministerium für Ernährung und Landwirtschaft, Grant/Award Number: 2817HS004

Editor: Raimundo Real

Abstract

Aim: Information on the spatio-temporal distribution of marine species is essential for developing proactive management strategies. However, sufficient information is seldom available at large spatial scales, particularly in polar areas. The Southern Ocean (SO) represents a critical habitat for various species, particularly migratory baleen whales. Still, the SO’s remoteness and sea ice coverage disallow obtaining sufficient information on baleen whale distribution and niche preference. Here, we used presence-only species distribution models to predict the circumantarctic habitat suitability of baleen whales and identify important predictors affecting their distribution.

Location: The Southern Ocean (SO).

Methods: We used Maxent to model habitat suitability for Antarctic minke, Antarctic blue, fin and humpback whales. Our models employ extensive circumantarctic data and carefully prepared predictors describing the SO’s environment and two spatial sampling bias correction options. Species-specific spatial-block cross-validation was used to optimize model complexity and for spatially independent model evaluation.

Results: Model performance was high on cross-validation, with generally little predicted uncertainty. The most important predictors were derived from sea ice, particularly seasonal mean and variability of sea ice concentration and distance to the sea ice edge.

Main conclusions: Our models support the usefulness of presence-only models as a cost-effective tool in the marine realm, particularly for studying the migratory whales’ distribution. However, we found discrepancies between our results and (within) results of similar studies, mainly due to using different species data quality and quantity, different study area extent and methodological reasons. We further highlight the limitations of implementing static distribution models in the highly dynamic marine realm. Dynamic models, which relate species information to environmental conditions contemporaneous to species occurrences, can predict near-real-time habitat suitability, necessary for dynamic management. Nevertheless, obtaining sufficient species and environmental predictors at high spatio-temporal resolution, necessary for dynamic models, can be challenging from polar regions.
1 | INTRODUCTION

Information on marine species’ spatio-temporal distribution and their relationship to the environment is pivotal for well-informed, proactive management strategies and conservation actions (Becker et al., 2016; Guisan et al., 2013). However, obtaining sufficient data on marine mammal distribution across large spatial scales is challenging due to financial and logistic constraints, particularly in remote oceans (Kaschner et al., 2012; Robinson et al., 2011). Marine mammal occurrence data are frequently biased towards coastal areas and shallow waters (Robinson et al., 2011) or, for polar regions, to easy-to-access regions during summer months.

Species distribution models (SDMs) are empirical methods that relate information on species occurrence to environmental variables to predict potential species distribution and identify potential ecological factors governing their distribution (Phillips et al., 2006). SDMs are promising to further our limited knowledge of marine mammals’ distribution and support marine conservation prioritization, for example, identify biologically important areas (Guisan et al., 2013; Redfern et al., 2006; Smith et al., 2020). Although SDMs in marine environments are relatively less common compared to their application in the terrestrial realm, recent years showed a significant increase in SDM usage for marine habitats (Marshall et al., 2014; Melo-Merino et al., 2020; Redfern et al., 2006; Robinson et al., 2011). The main challenge to model the distribution of marine species is the availability of sufficient reliable species data (Dambach & Rödder, 2011; Robinson et al., 2011).

Two species information types are commonly used in SDMs: presence–absence and presence-only data. Presence–absence models (e.g. generalized additive models—GAMs) require carefully designed surveys and thus are more common in small-scale SDM studies (e.g. Esteban et al., 2013; de Stephanis et al., 2008). Absence data are hard to estimate correctly (Lobo et al., 2010), especially for highly mobile species and from remote areas (Smith et al., 2020). Marine mammals spend a vast amount of time submerged and can be visually detected only when on or near the water surface. Their detection is sensitive to species behaviour and oceanographic and meteorological conditions (Barlow et al., 2001). This imperfect detection can lead to false absences, which affect SDMs evaluation and bias species distribution inferences (Guilera-Arroita, 2017; Lobo et al., 2010). This is even more serious as the detectability of marine mammals varies in time and space (Guilera-Arroita, 2017). Furthermore, even dedicated surveys typically provide only a snapshot of species distribution and represent only a limited time and space range (Kaschner et al., 2006). Hence, not surprisingly, most SDMs use presence-only data. Presence-only models contrast species occurrences to a large sample of background locations to characterize the environment throughout the study area. Recent literature demonstrates the statistical validity of only a few presence-only SDM algorithms, including point process models and Maxent (Renner et al., 2015). The implementation of robust presence-only SDMs is particularly advantageous in the marine realm due to the difficulty of efficiently obtaining systematic presence-absence data (Smith et al., 2020).

The Southern Ocean (SO) is a biodiversity hotspot area, showing distinctive biogeographic features and high environmental variability (Convey et al., 2014; De Broyer et al., 2014; Fabri-Ruiz et al., 2019; Guillaumot et al., 2020). The SO’s sea ice environment represents a critical habitat for many threatened migratory and resident species, particularly for baleen whales (Filun et al., 2020; Thomisch et al., 2016; Van Opzeeland et al., 2013). Nevertheless, research efforts in the SO were limited because of its remoteness, vastness and sea ice coverage, posing considerable financial and logistical constraints (Bombsch et al., 2014; Scheidat et al., 2011). Our knowledge of the biodiversity in most SO areas seems to reflect sampling effort rather than the actual biodiversity status (Convey et al., 2014) and thus improving sampling effort deserves a high priority for Antarctic science (Guillaumot et al., 2018).

Spatio-temporal information on species distributions from the SO, necessary for conservation planning and management, is particularly patchy. Research efforts are generally biased towards relatively small areas of the SO (e.g. the West Antarctic Peninsula), repetitive ship tracks (e.g. to and from Antarctic stations) and mainly limited to summer months. Simultaneously, deep-sea and remote regions (e.g. the Bellinghausen and Amundsen Seas) remain largely underinvestigated (De Broyer et al., 2014). Most research vessels that operate in the SO are biased towards the operationally safe ice-free water and do not engage in the risk and costs of going deep into the sea ice (Herr et al., 2019; Williams et al., 2014), rendering modelling species distribution in the SO challenging (Guillaumot et al., 2018, 2020). Nevertheless, carefully implemented and evaluated presence-only SDMs can be a cost-effective tool to study species potential distribution and habitat and planning for future surveys in the SO.

In the SO, several baleen whale species have been extensively hunted to near extinction levels during the 20th-century commercial whaling, particularly Antarctic blue and fin whales (Kennis et al., 2016; Tulloch et al., 2018). Populations recovery is generally incomplete and shows variant recovery rates between species and SO regions, with some species exhibiting high recovery rates (e.g. humpback whales, Megaptera novaeangliae; Friedlaender et al., 2011; Tulloch et al., 2018) while others remain highly threatened (e.g. Antarctic blue whales, B. musculus intermedia; Branch et al., 2004; Tulloch et al., 2018). Information on the ecology and distribution of baleen whales in the SO is pivotal for

**KEYWORDS**
Antarctic blue whale, Antarctic minke whale, baleen whales, fin whale, humpback whale, Maxent, presence-only models, Southern Ocean, species distribution models, static species distribution models
the International Whaling Commission’s conservation efforts and measures addressing potential climate change impacts in polar ecosystems (Williams et al., 2014). However, such information is limited (Leaper & Miller, 2011); and thus, relatively few studies have modelled the distribution of baleen whales in the SO. Some species receive more attention, for example, humpback whales, while others, for example, fin and Antarctic blue whales, receive less attention (Širović & Hildebrand, 2011).

The focus of this paper is to model the circumantarctic distribution of four baleen whale species that feature sufficient sighting data: Antarctic minke whale (AMW, *Balaenoptera bonaerensis*); Antarctic blue whale (ABW); fin whale (FW, *B. physalus*); and humpback whale (HW). We performed a rigorous screening of baleen whale circumpolar distribution data in the SO. We used Maxent (Phillips et al., 2006) as it is appropriate for the available presence-only data, with two ways of handling spatial sampling bias (no correction versus rarefaction). We used spatial-block cross-validation for independent model evaluation and optimizing model complexity to improve predictions. For each species, we predicted its circumantarctic habitat suitability and identified the most important predictors affecting their distribution and species suitability response to environmental changes. We compared our results with previous studies on these species in the SO and discuss reasons for observed differences. Finally, we evaluate the potential limitations of implementing static SDMs in the highly dynamic SDMs.

## Methods

### 2.1 Species data

Cetacean sightings south of 45°S were compiled from different sources. Only sightings after 1980 were considered to maintain a reasonable temporal match between environmental predictors and sightings. Data from three biodiversity repositories were quality controlled: the Global Biodiversity Information Facility (GBIF; https://www.gbif.org/), the Ocean Biodiversity Information System (OBIS, 2018) and OBIS-SEAMAP (Halpin et al., 2009). Other data sources include SO GLOBEC (2001–2002; http://www.ccpo.odu.edu/Research/globec_menu.html), SOWER cruises (https://iwc.int/sower; 2009–2010), RV Polarstern expeditions (https://www.awi.de/expedition/schiffe/polarstern.html) and data published in PANGAEA (https://www.pangaea.de/; details in Appendix S1–S5).

Data on baleen whales with sufficient sightings (AMW, ABW, FW and HW) were subjected to further quality control. We excluded erroneous occurrences or those with high uncertainty, for example, GBIF occurrences flagged with “known geospatial issues” and “possible” certainty level for Polarstern data. As biodiversity data repositories compile data from various sources, the same sighting can be duplicated within or between repositories. We excluded occurrences explicitly duplicated within and between data sources to avoid spurious high relative occurrence rates: only one instance of sightings with identical coordinates and date was retained. We excluded telemetry and catch data to avoid highly correlated occurrences, spatially or temporally. The final dataset consists of ~32 thousand sightings. The temporal distribution of species-specific sightings is shown in Figures S2, S7, S12 and S17. Note that figures in the Supporting Information are grouped by species (Figures S1–S20).

### 2.2 Environmental predictors

Potential predictors were obtained at the highest available spatial and temporal resolution (Table S1). We prepared ecologically relevant predictors summarizing environmental conditions in the SO and act as a proxy for prey availability (Redfern et al., 2006). We calculated monthly and seasonal mean and standard deviation of each dynamic predictor to explore temporal trends and intra-seasonal variability, respectively. Seasons were determined as three-month intervals from January, except for metrics representing sea ice (see below).

Bathymetry data were downloaded from GEBCO (Weatherall et al., 2015). From bathymetry, we derived slope, aspect and closest distances to coast, 500 m and 1,000 m isobaths. The Antarctic coast was defined as the ice shelf edge, that is, excluding any cavities under the ice shelves. The 1,000 m isobath was used to represent the location of the continental shelf break.

Chlorophyll-a concentrations (Chl-a) were downloaded as 8-day composites from OCCI (2002–2017; Sathyendranath et al., 2018). We only considered Chl-a mean and standard deviation in summer, as the spatial coverage in other seasons was rather poor, prohibiting the calculation of meaningful circumpolar averages.

Daily absolute dynamic topography (sea surface height, SSH) was obtained from Copernicus (https://copernicus.eu/; 1993–2017), following Bomboesch et al. (2014), from which daily (current) speed was estimated. We found only little inter-annual variability of SSH and speed, persuading us to use the annual mean and standard deviation of the whole period.

Temperature and salinity data at five standard depths (surface, 100 m, 200 m, 500 m and 1,000 m) were obtained from the World Ocean Atlas (1981–2010; Locarnini et al., 2019; Zweng et al., 2019).

Daily sea ice concentration (SIC) was obtained from Spreen et al. (2008). We used SIC data for complete years (2003–2010 and 2013–2017), with seasons customized according to the major phases of annual sea ice extent (https://seaice.uni-bremen.de/sea-ice-conce ntration/time-series/): season 1 (January–March, summer, lowest extent); season 2 (April, sea ice formation start); season 3 (May–November, high extent); and season 4 (December, high sea ice melting). We determined the closest distance to seasonally averaged sea ice edge (SIE), where SIE was identified as the largest polygon with mean SIC >15% (Parkinson, 2002). We assigned a value of zero to cells intersecting with SIE, positive values north of SIE (open water; SIC <15%) and negative values south of SIE (SIC >15%) (following Ainley et al., 2004).

All predictors were projected into equal-area projection at 10 × 10 km resolution. All analyses were restricted to south of
the climatological location of the Polar Front as defined by Orsi et al. (1995), which was chosen as a natural boundary of the SO with rather homogeneous hydrographic conditions south of it. Spatial gaps were interpolated using ordinary Kriging (Wackernagel, 1995) when necessary. After the rejection of less-informative predictors, as based on their temporal trends and personal experience, the initial list of predictors included 32 predictors (Table S1b).

We implemented predictor transformation when necessary (e.g. square root) to avoid the effect of few extreme values on model stability (Dormann & Kaschner, 2010). We excluded highly correlated predictors by maintaining a moderate maximum variance inflation factor of 4.5 (Zuur et al., 2010). This approach resulted in a total of 15 predictors used in the models (Figures S21–S22 and Table 1). Figure S23 shows environmental conditions at species-specific sightings against their full range in the study area.

2.3 | Species distribution models

We used MAXENT v3.4.1 (Phillips et al., 2017) to train two model sets: (1) using all occurrences to estimate habitat suitability under the point process modelling framework (following: Renner et al., 2015; ModelAll) and (2) using only one occurrence per cell (Model Unique). The latter is a special case of rarefaction, a commonly used method to correct for sampling bias and diminish the effect of spatial autocorrelation (Aiello-Lammens et al., 2015). It is expected that bias correction can lead to broader areas of suitable habitats (El-Gabbas & Dormann, 2018a; Phillips et al., 2009). Here, we used both models not to quantify the effect of sampling bias corrections, but to investigate whether and how they would affect our conclusions, under the assumption that differences in results reflect on model stability.

We used a 5-fold spatial-block cross-validation to evaluate model performance by maintaining spatial independence between training and testing dataset and to reduce the effect of spatial autocorrelation (Roberts et al., 2017). We determined block size and how to distribute blocks into cross-validation folds using blockCV R-package (Valavi et al., 2019): size was determined as median spatial autocorrelation range of environmental conditions at sighting locations; blocks were distributed into folds balancing the number of occurrences (Figure S24).

To improve model performance, we tuned Maxent’s parameters using cross-validation (Merow et al., 2013). We used ENMeval R-package (Muscarella et al., 2014) to estimate the best combination of feature classes (transformation of predictors) and regularization multiplier (model complexity). For each model type and species, we used 40 combinations: five feature classes (L/LQ/H/LQH/LQHP; where “L” linear, “Q” quadratic, “H” hinge and “P” product transformation) and eight regularization multiplier values (0.5 to 4, with 0.5 increment). The combination with highest testing AUC (area under the ROC curve) using cross-validation was used in the final models (Table S2). We present the mean habitat suitability along with the coefficient of variation (ratio between standard deviation and mean prediction) as a measure of predictive uncertainty. In addition to cross-validation, we ran full models that used all occurrences. In each model, we estimated predictor importance using permutation importance and jackknifing. We show the results of the full models in the main text and cross-validated models in the Supporting Information.

| Table 1 | List of environmental predictors used in the models |

| Predictor | Statistics | Season | Transformation | Abbreviation | VIF |
|-----------|------------|--------|---------------|--------------|-----|
| Bathymetry |            |        |               | Bathymetry   | 3.0 |
| Slope     |            |        | Natural log   | Slope        | 1.5 |
| Distance to coast |            |        |              | DistCoast    | 2.2 |
| Distance to 1,000 m isobath |            |        | Square root   | DistIsobath1000 | 2.8 |
| Chlorophyll-a | Mean | Summer | Natural log   | Chl-a        | 1.9 |
| Sea ice concentration | Mean | Summer | Natural log   | SIC_Mean_S1  | 3.7 |
| Distance to the ice edge |            |        | Summer        | DistIceEdge_S1 | 3.8 |
| Sea ice concentration | SD | April  |              | SIC_SD_S2    | 2.6 |
| Sea ice concentration | SD | May–November |              | SIC_SD_S3    | 2.8 |
| Sea ice concentration | SD | December |              | SIC_SD_S4    | 3.3 |
| Temperature (200 m) | Mean | Annual |              | Temp_200     | 3.4 |
| Surface salinity | Mean | Annual |              | Sal_Surf     | 1.9 |
| Sea Surface Height | Mean | Annual |              | SSH_Mean     | 4.4 |
| Sea Surface Height | SD  | Annual |              | SSH_SD       | 2.6 |
| Speed     |            |        |              | Speed        | 3.8 |

Note: Statistics: type of statistics used to calculate each predictor (SD = standard deviation); season: which season or month range was used; transformation: transformations implemented to maximize uniformity of the data; abbreviation: the abbreviation used in the figures; VIF: the value of variance inflation factor. Summer was defined as from January to March. See Table S1 for more information on the predictors used.
FIGURE 1 Predicted habitat suitability of four baleen whale species in the Southern Ocean using Model\textsubscript{All} (all occurrences, left) and Model\textsubscript{Unique} (removed duplicate sightings, right). These maps represent predictions from the respective “full model,” calibrated without cross-validation. Mean prediction from cross-validated models and their coefficient of variation are shown in the Supporting Information. Map colours range from blue (low suitability) to red (high suitability). All maps are on Maxent’s cloglog scale.

3 | RESULTS

In general, both model types (Model\textsubscript{All} and Model\textsubscript{Unique}) give similar results, with Model\textsubscript{Unique} resulted in a broader range of suitable habitats and slightly lower testing AUC, as expected after bias correction (Figure 1 and Table S2). Generally, the most important predictors were sea ice related (Figure 2). The uncertainty of cross-validated predictions was generally low and did not show a pronounced spatial pattern, reflecting the stability of these sub-models.
3.1 | Antarctic minke whale

Models predicted a circumantarctic habitat of AMW, with a general preference closer to the Antarctic coast except for a small patch southwest of the Balleny Islands and the Amundsen Sea coast towards the Ross Sea. Most of the southern part of the Weddell Sea was predicted less suitable (Figure 1 and Figure S1). The most important predictors were distance to summer SIE, mean summer SIC and SIC variability (Figure 2 and Figure S3). AMW was shown to prefer locations close to SIE and moderate SIC (<50%; Figures S4 and S5).

3.2 | Antarctic blue whale

Suitable areas for ABW were near the Antarctic coast (yet 50–300 km offshore), ranging from 30°W eastwards to 170°W (Figure 1 and Figure S6), that is, along the East Antarctic coast and notably rather sparsely off West Antarctica. Other suitable areas include small patches in the Bellingshausen and Amundsen Seas and between Elephant and the South Sandwich Islands. The most important predictors were SIC variability, mean summer SIC and distance to 1,000 m isobath (Figure 2 and Figure S8). Other relatively important predictors were bathymetry, temperature at 200 m and distance to summer SIE. Suitable habitats were predicted in areas with high SIC variability in December (c.a. 35%–45%) and low mean summer SIC (<40%) or low-to-moderate distance to 1,000 m isobath (<250 km; Figures S9 and S10). ABW habitat is more suitable close to SIE (with lower suitability south of it), at high temperature at 200 m (3–5°C) and locations with moderate depths (3,500–4,500 m; Figure S9).

3.3 | Fin whale

The most suitable areas for FW extend eastwards from Elephant Island to South Georgia Island, near Bouvet Islands, small patches close to the Antarctic coast from 30°E eastwards to 180°E and offshore of the Ross Sea (Figure 1 and Figure S11). Important predictors were distance to summer SIE, mean summer SIC, SIC variability, distance to coast and SSH variability (Figure 2 and Figure S13). Highest
suitability was shown north of the SIE (<200 km and at ~1,500 km from it), only <100 km from the coast) or locations with low SIC (<50%), low temperature at 200 m (<-1.5°C) or low SSH variability (Figures S14 and S15).

3.4 | Humpback whale

The effect of sampling bias correction on predicted distribution was most evident for HW, due to intensive sampling west of the Antarctic Peninsula and in East Antarctica. Generally, suitable areas are the Western Antarctic Peninsula eastwards to the South Orkney Islands, around the South Sandwich and Bouvet Islands, and a strip close to the coast from 15°W eastwards to 170°W (Figure 1 and Figure S16). The most important predictors were distance to summer SIE, SIC variability from April to November and summer SIC. Other important predictors include distance to coast, distance to 1,000 m isobath and SSH variability (Figure 2 and Figure S18). HW suitability was higher at locations close to SIE at summer SIC <60%. On the open water side of SIE, high suitability was found only at locations with high SIC variability (Figures S19 and S20). Moderate suitability is predicted <300 km from 1,000 m isobath and locations close (<100 km) or far (>1,000 km) from the coast (Figure S19).

4 | DISCUSSION

4.1 | Baleen whale habitats in the Southern Ocean

Overall, the most important predictors affecting baleen whales’ habitat suitability in the SO are those derived from SIC. Sea ice cover varies within and between years, and this variability plays an integral role in whale distribution (Thiele et al., 2004). Our use of seasonal variability of SIC can be considered as a proxy for site accessibility for whales; the higher the SIC standard deviation, the more accessible for whales (Wege et al., 2020). SIC variability affects prey (krill) survival, population dynamics and abundance (Fraser & Hofmann, 2003; Thiele et al., 2004), with highest observed abundances close to the SIE (Brierley et al., 2002; Murase et al., 2002; Thiele et al., 2004). Obtaining reliable data on the distribution and abundance of prey, particularly krill, are currently not possible at the circumantarctic scale (Robinson et al., 2011), rendering most studies dependent on remotely sensed predictors as a proxy for prey availability (Herr et al., 2019).

Transition zones, for example SIE and continental shelf break, are known high-productivity areas (Beekmans et al., 2010). The use of predictors describing distance to them can serve as a proxy for prey availability. The majority of visual observation data available to us were recorded using vessels unsuited for penetrating the ice, except...
a few sightings obtained from icebreaker vessels (e.g. Polarstern) and icebreaker-supported helicopter surveys (e.g. Herr et al., 2019). This explains why only little sightings came from the south of the SIE. Nevertheless, distance to SIE was one of the most important predictors for the models of the four study species. In the following, we briefly compare our species-specific results with results of other studies (summarized in Tables 2 and S3) to evaluate the models’ reliabilities in general.

### 4.1.1 | Antarctic minke whale

Although AMWs are thought to be the most abundant cetacean species in the SO (Williams et al., 2014), they are among the least studied marine mammal populations (Risch et al., 2019). AMWs have a circumantarctic distribution and are considered the major consumer of Antarctic krill in the SO (Beekmans et al., 2010; Kasamatsu, Matsuoka, et al., 2000; Williams et al., 2014). Highest AMW density was estimated in the Western Antarctic Peninsula and the Weddell and Ross Seas (Dominello & Širović, 2016; Risch et al., 2019). The AMW is a year-round resident in the SO and occurs throughout a wide range of SIC (Filun et al., 2020; Friedlaender et al., 2011; Herr et al., 2019; Thiele et al., 2004), preferring the SIE area (Dominello & Širović, 2016; Herr et al., 2019; Kasamatsu, Ensor, et al., 2000; Scheidat et al., 2011; Williams et al., 2014). It has been observed both within the pack ice region and in open water (although in lower numbers) (Beekmans et al., 2010; Ensor, 1989; Friedlaender et al., 2006; Herr et al., 2019; Thiele & Gill, 2004; Williams et al., 2014). They can exploit pack ice and forage krill through sea ice, which is mostly unavailable to other baleen whales, due to their compact small-sized body, hard and pointed nostrum and high maneuverability (Ainley et al., 2012; Friedlaender et al., 2014).

Accordingly, we found high importance of SIC-derived predictors, particularly distance to summer SIE and summer SIC. Sea ice is an essential habitat for AMW and affects their distribution and foraging behaviour in the SO (Friedlaender et al., 2014; Herr et al., 2019; Kasamatsu, Ensor, et al., 2000; Risch et al., 2019). Thus, with potential future climate change being expected to affect Antarctic krill population dynamics, the AMW’s suitable habitats will shrink throughout the SO (Ainley et al., 2012; Herr et al., 2019; Risch et al., 2019; Williams et al., 2014). Our models show high predicted habitat suitability at SIC up to 50% and low at higher SIC values. Bombosch et al. (2014) showed that AMW habitat suitability was consistently predicted in sea ice-covered areas in the SO. Ainley et al. (2012) showed a consistent positive effect of sea ice cover on AMW suitability, and similarly, Filun et al. (2020) found a strong positive correlation between SIC and AMW acoustic presence in the Weddell Sea, with the highest acoustic activity occurring at SIC >75%. Distance to SIE was also important in other SDM studies with highest suitability close to it (Beekmans et al., 2010; Bombosch et al., 2014; Friedlaender et al., 2011; Herr et al., 2019; Kasamatsu, Ensor, et al., 2000; Murase et al., 2013; Williams et al., 2014). In contrast, Filun et al. (2020) reported very little acoustic activity near the SIE area in the Weddell Sea during December and January. Some studies discussed the important role of the Antarctic ice shelf break: higher suitability closer to it with a strong decline with increasing distance (Ainley et al., 2012; Beekmans et al., 2010; Herr et al., 2019; Murase et al., 2013). In our model, distance to 1,000 m isobath had very low importance, although with a similar (albeit weak) relationship pattern. We found low importance of bathymetry (negative relationship), distance to coast, slope, Chl-a (negative relationship) and positive relationship for salinity and water temperature at 200 m, but see Kasamatsu, Ensor, et al. (2000), Friedlaender et al. (2011), Ainley et al. (2012) and Murase (2014) for contradicting results.

### 4.1.2 | Antarctic blue whale

ABW was once an abundant species in the SO, but is currently extremely rare after its intensive exploitation during the whaling industry era from 1904 until 1978 (Branch et al., 2007; Double et al., 2015; Kasamatsu, 1988; Miller et al., 2015). After the ceasing of the whaling industry, the circumpolar ABW abundance was reported to be depleted to only less than 1% of its original abundance before whaling (Branch et al., 2004, 2007), making the ABW one of the most endangered baleen whale species in the SO (Leaper & Miller, 2011). Little is known on the distribution and migration patterns of ABW in the SO and its relationship with krill (Branch et al., 2007; Double et al., 2015; Thomisch et al., 2016). ABWs were visually sampled relatively infrequently in the SO in comparison with other baleen whales (Murase, 2014; Širović & Hildebrand, 2011), but their calls can be accurately detected (Thomisch et al., 2016). This allowed some studies to model ABW distribution using passive acoustic data (Shabangu et al., 2017; Širović & Hildebrand, 2011).

We found the most important predictors are SIC-derived and distance to 1,000 m isobath. Other SDM studies provide limited information on the effect of sea ice on ABW’s suitability. High ABW habitat suitability was predicted at low SIC (<40%) and close to summer SIE (Figure S9). Similarly, Širović et al. (2004) and Thomisch et al. (2016) reported a negative correlation between sea ice coverage and the number of detected ABW calls in the Western Antarctic Peninsula and the Weddell Sea, respectively. Nevertheless, ABW was also acoustically present in areas with high winter SIC (90%) in the Weddell Sea (Thomisch et al., 2016) and under non-navigable ice conditions in the Ross Sea (Double et al., 2015). This suggests the overwintering of ABW in highly ice-covered areas, potentially in local recurring polynyas (Thomisch et al., 2016). A high encounter rate of ABW near the SIE was also reported by other studies (Branch et al., 2007; Kasamatsu, 1988; Kasamatsu, Matsuoka, et al., 2000; Rankin et al., 2005; Širović et al., 2004).

We found moderate importance for bathymetry and temperature at 200 m. Highest suitability was found at around 5,000 m and lower elsewhere. Širović and Hildebrand (2011) found that in the
### Table 2
Summary of the comparison between this study’s results and like studies on the Antarctic minke whale in the SO

| Other studies                                | This study | High importance (>5%) | Low importance (<5%) |
|-----------------------------------------------|------------|-----------------------|----------------------|
| Near and S. of summer SIE                     | ⊖ ⊖        | Unclear               | ⊖ ⊖                  |
| Ainley et al. (2012)                          | ✓          | High imp.             | High imp./high imp.  |
| Beekmans et al. (2010)                        | ⊖          | Low imp.              | ⊖                    |
| Bombosch et al. (2014)                        | ✓ ⊖        | Moderate imp. / high  | Moderate imp.        |
| Friedlaender et al. (2011)                    | Low        | Most imp. predictor   | Moderate imp.        |
| Filun et al. (2020)                           | Little     | Low to moderate imp.  | Moderate imp.        |
| Herr et al. (2019)                            | ✓          | High imp.             | ✓                    |
| Kasamatsu, Ensor, et al. (2000)               | ✓          | High imp.             | ✓                    |
| Kasamatsu, Matsuoka, et al. (2000)            |            |                       |                      |
| Murase (2014)                                 |            |                       |                      |
| Murase et al. (2013)                          | ✓          | Low imp.              | ✓                    |
| Williams et al. (2014)                        | ✓          | Low imp.              | ✓                    |

Note: Important predictors, as identified by this study, are shaded dark grey in the column header (>5% permutation importance for the full models, in descending order). Cell colours represent the agreement between our study results and other studies: green, high agreement; orange, some disagreement; red, high disagreement. Empty cells represent situations when the given predictor was not tested. Similar results for other species are shown in Table S3.

Abbreviations: ⊖ = positive relationship; ⊖ = negative relationship; ✓ = similar results; Chl-a, chlorophyll-a concentration; Dist., distance; imp., importance; SD, standard deviation; SIC, sea ice concentration; SIE, sea ice edge; SSH, sea surface height; Temp., water temperature.
Pacific along the Western Antarctic Peninsula acoustic presence is more suitable at greater depths. Similarly, Murase (2014) found a high abundance peak at depth \(-4,000\) m, but with an additional peak near 0 m, that is, close to the coast. In contrast, Shabangu et al. (2017) found the least suitability at around 5,000 m. We found a positive relationship with temperature at 200 m, which coincides with results for calling presences by Širović and Hildebrand (2011). In contrast, Kasamatsu, Matsuoka, et al. (2000) reported a high encounter rate at lower temperatures, and Shabangu et al. (2017) showed high suitability of calling whales at \(-0°C\) sea surface temperature (SST). We found moderately low importance of distance to coast, SSH (positive relationship) and Chl-\(a\) (positive relationship). Similarly, Širović and Hildebrand (2011) found a non-significant relationship between Chl-\(a\) and calling ABW off the Western Antarctic Peninsula. In contrast, Shabangu et al. (2017) showed that these predictors were among the most important predictors for call detections: peak suitability close to coast, then sharply declined until \(-1,000\) km; low suitability at SSH around \(-1.5\) m and high elsewhere and high suitability at low Chl-\(a\).

### 4.1.3 Fin whale

Although FW was the most caught species in the SO during the 20th-century commercial whaling (>718 K whales taken), there is limited information on its distribution, abundance, demographics and environmental variables affecting its ecology (Herr et al., 2016; Santora et al., 2014). A relatively recent estimation of FW population in the SO has shown that it is currently at only 2% of the presumed pre-whaling estimated abundance (Leaper & Miller, 2011). We found that the most important predictors are SIC-derived predictors, distance to coast, SSH variability and temperature at 200 m. We found highest (although moderate) suitability close and far (\(-1,500\) km) from the coast. In contrast, Williams et al. (2006) found that abundance increases with the distance from coast, with the lowest intensity close to it off the northern Antarctic Peninsula. Santora et al. (2014) reported FW preference for more complex bathymetry off the northern Antarctic Peninsula. Murase (2014) found three abundance peaks at depths of 4,500 m, 2,200 m and 0 m, while Williams et al. (2006) reported a low intensity in depths <1,000 m. However, we found low importance of bathymetry, with two low suitability peaks at around 4,500 m and near 0 m. We found highest suitability at cold water (<\(-1.5°C\)). In contrast, Santora et al. (2014) found a positive correlation between SST and FW abundance off the Western Antarctic Peninsula, and similarly, Kasamatsu (1988) and Kasamatsu, Matsuoka, et al. (2000) reported a higher encounter rate at warmer temperatures (\(>1°C\)). We found low importance of Chl-\(a\) with no clear relationship, which conforms with Murase (2014) except at high Chl-\(a\) at which high FW abundance was predicted. We found two moderate suitability peaks in open water, either close or far from (\(-1,500\) km) SIE. In contrast, other studies noted that FWs are rarely reported near the SIE (Širović et al., 2004); Kasamatsu, Matsuoka, et al. (2000) found a high encounter rate far from SIE, and, similarly, Scheidat et al. (2011) reported that the FW majority was observed >140 km from SIE.

### 4.1.4 Humpback whale

Although HWs were highly exploited during the 20th-century whaling industry, with >150,000 caught whales between 1904 and 1966 (Nowacek et al., 2011), the population has been increasing since the cessation of the whaling industry (Friedlaender et al., 2011). HWs are the most common whale species in the Western Antarctic Peninsula area in summer (Scheidat et al., 2011) and seem to be absent from the Ross Sea (Branch, 2011; Leaper & Miller, 2011). This conforms with the areas predicted as suitable habitats by our models (Figure 1 and Figure S16). Important predictors were SIC-derived, distance to coast and 1,000 m isobath, as well as SSH variability. Highest suitability is predicted at locations with low SIC or locations either close to SIE or far from it on the sea ice-free side (higher). Schall et al. (2020) found a weak correlation between SIC and HW acoustic presence in the Atlantic sector of the SO, while Van Opzeeland et al. (2013) reported HW acoustic presences at high SIC values (>90%) at an Antarctic coastal recording site during winter. Friedlaender et al. (2011) and Thiele et al. (2004) found highest suitability close to SIE in the Western Antarctic Peninsula. Bombosch et al. (2014) found that HW suitable habitats are primarily in ice-free areas and follow the sea ice retreat. There is apparently a lagged effect of sea ice dynamics on the habitat suitability of HW, suggesting that HWs do not actively track the location of SIE recent retreat, but instead the high productivity that occurs up to two months after sea ice melting (Riekkola et al., 2019). Andrews-Goff et al. (2018) found that predictors associated with the marginal ice zone as the main predictors for HW foraging habitat in the Pacific sector of the SO, particularly mean SIC one month prior to HW arrival to the SO, SIC variability two months prior to arrival, and the distance to SIE (highest at \(-65\) km), while Riekkola et al. (2019) similarly found that distance to SIE two months prior to arrival as an important driver for HW behaviour in the SO.

High HW suitability was predicted at areas very close to the coast, which conforms with the frequent sightings of HWs in coastal areas of the Antarctic Peninsula (Dalla Rosa et al., 2008; Nowacek et al., 2004; Thiele et al., 2004) and near the Greenwich Meridian (Van Opzeeland et al., 2013). We found low importance of bathymetry and slope, with moderate suitability in shallow areas (<1,000 m) and low elsewhere. Similar results were also shown by Murase (2014), while Friedlaender et al. (2011) reported that HW occupies rugged topography around Marguerite Bay, Western Antarctic Peninsula, with bathymetry and slope among the most important predictors. Murase (2014) found highest abundance at low salinity, two peaks at moderate and high SSH, and high Chl-\(a\). Owen et al. (2019) found slope, Chl-\(a\) and SST among the most important predictors for HW foraging behaviour in East Antarctica. In contrast, we found no clear
relationship with salinity, Chl-α and temperature at 200 m, and negative with SSH, but neither of them was an important predictor. In concordance to our results, Riekkola et al. (2019) found a negative relationship between HW foraging behaviour in the Pacific sector of the SO and SSH and low importance of speed, while Kasamatsu, Matsuoka, et al. (2000) found no relationship between HW density and SST.

4.2 | Reasons for discrepancies between studies

Unambiguously asserting reasons for the discrepancies between the results discussed above is challenging, as we do not know the true preferred niche of these species. Generally, inconsistency can be attributed to data and methodological reasons. Most studies used occurrences from a limited time frame (e.g. from within summer months of 1–2 years) or covered only a small section of the SO, for example, the northern Antarctic Peninsula (Santora et al., 2014; Williams et al., 2006), the Western Antarctic Peninsula (Friedlaender et al., 2011; Kasamatsu, 1988; Murase et al., 2013; Širović & Hildebrand, 2011; Thiele et al., 2004), East Antarctica (Owen et al., 2019), the Pacific sector of the SO (Andrews-Goff et al., 2018; Riekkola et al., 2019), the Weddell Sea (Filun et al., 2020; Schall et al., 2020; Thomisch et al., 2019; Van Opzeeland et al., 2013; Williams et al., 2014), the Ross Sea (Murase et al., 2013) and the Bellingshausen and Amundsen Seas (Kasamatsu, Ensor, et al., 2000). The use of spatially or temporally limited sightings and environmental data makes it difficult for these models to capture the full range of species niche (e.g. causing truncated or biased response curves; Barbet-Massin et al., 2010; Thuiller et al., 2004). Although it is technically possible for these models to predict potential distributions at the circumantarctic scale, the necessary extrapolation to novel conditions or new combinations increases prediction uncertainty (Zurell et al., 2012).

Contrastingly, this study used circumantarctic visual observation data, covering a wide range of baleen whale suitable environmental conditions (and their combinations) in the SO. To date, only a few studies investigated the distribution and niche characteristics of baleen whales at the circumantarctic scale (e.g. Bomboš et al., 2014; Branch, 2011; Branch et al., 2007), possibly due to challenges obtaining sufficient data. SDM studies at large scales such as the SO assume stationary species–environmental relationships through space and time, that is, same niche characteristics at smaller areas of the SO or between seasons (Dormann et al., 2012; El-Gabbas & Dormann, 2018B; Osborne et al., 2007). The distribution of baleen whales varies between seasons and spatial divisions of the SO (Riekkola et al., 2019; Thiele et al., 2004). For example, Beekmans et al., (2010) found inconsistent relationships between environmental predictors and AMW density at circumantarctic and regional scales, suggesting that the relationships between AMW and environmental conditions can be best studied at a regional rather than circumantarctic scale.

The vast majority of our sightings were made from the end of December to the end of February (Figure S25). This evident temporal bias towards summer months seems inevitable when using only visual observation data. Passive Acoustic Monitoring (PAM), however, has provided ample evidence for the (near-) year-round presence of several species in this area (Filun et al., 2020; Schall et al., 2020; Thomisch et al., 2016; Van Opzeeland & Hillebrand, 2020; Van Opzeeland et al., 2013). Although we attempted to correct for spatial sampling bias using rarefaction, the absence of visual observations from the Weddell Sea has affected model predictability in this area (Figure 1). The integration of other data types in SDMs, for example, from tagged animals (e.g. Hindell et al., 2020) and PAM, will be able to fill this gap and forward our understanding of year-round niche preferences of these species.

Moreover, studies implemented different response types (e.g. presence-only vs. presence-absence), modelling techniques (e.g. GAMs vs. Maxent), spatial and temporal resolutions, predictor’s combinations, environmental bias patterns and data quality and sampling methods.

Furthermore, marginal response curves used to describe species response can be deceptive. To estimate a species’ marginal response curve for any predictor, each other predictor is fixed at one value, neglecting the true multi-dimensionality of the environmental space. For example, Maxent uses a predictor-specific mean value at training observations. The response curve shape can be sensitive to the values at which other predictors are fixed, especially when using limited or biased data or correlated predictors. To overcome this caveat, we show pairwise mean habitat suitability in the environmental space of the most important three predictors, while allowing all predictors to vary together (e.g. Figure S5). However, Maxent quantifies permutation importance based on training AUC drop after permutation (Phillips, 2017). Thus, spatio-temporal biases in species data can highly affect this estimate.

4.3 | Static SDMs in highly dynamic marine environments

The majority of SDMs, particularly when covering large spatial scales, including this study, are static. Static models use predictors summarizing environmental conditions over long periods (seasonal or annual averages over >10–50 years; e.g., Sbrocco & Barber, 2013), irrespective of the exact time of species sighting (Bateman et al., 2012). They assume species–environment relationships fixed in space and time and that locations with species detections represent suitable year-round habitats, which likely is a rather poor assumption, especially for migratory species (Bateman et al., 2012; Reside et al., 2010).

Static models are more appropriate in highly static environments (as is the case for many terrestrial settings) and for modelling less mobile resident species (e.g. plants and lizards). However, the marine environment is immensely dynamic and undergoes significant changes over short periods, which likely affects the distribution of highly mobile species (Fernandez et al., 2017). Static models can neither capture environmental dynamics nor predict near-real-time species distribution necessary for dynamic
ocean management. In a dynamic setting, static models can only provide a fictitious representation (in time) of species suitability for the period over which the model is calibrated. To obtain robust SDMs, it is necessary to maintain a spatio-temporal match between species occurrences and environment (dynamic SDMs; Fernandez et al., 2017; Reside et al., 2010). This is particularly important for highly mobile marine species whose distribution is defined by both short- and long-term variations in ocean conditions (Mannocci et al., 2017). In contrast to conventional static models, dynamic SDMs capture the year-round species–environment relationships and allow predicting habitat suitability at finer temporal resolution (day-week-month).

The environment in polar regions, particularly the SO, is highly dynamic due to the seasonal waxing and waning of sea ice (Dayton et al., 1994). It hence appears intuitive to use dynamic, rather than static, SDMs to study habitat preference of migratory whales in the SO. However, obtaining many circumantarctic oceanographic variables at fine spatial and temporal resolution is challenging, compromising dynamic models’ feasibility. Many variables are limited to the sea surface and are not available at high temporal resolution (e.g. daily or weekly) (Fernandez et al., 2017). For example, daily or weekly salinity and productivity data are not available from the SO, and daily oceanic temperatures are limited to the water surface. Other variables show inconsistent and incomplete spatial coverage year-round. For example, Chl-a data are highly patchy and limited to summer months, which constrains its use in year-round dynamic models.

The unavailability of sufficient, less temporally and spatially biased sightings hinders efficient use of dynamic models and can, in part, explain modellers’ preference for static over dynamic models (Milanesi et al., 2020). High spatio-temporal resolution of some environmental predictors became available only recently. For example, daily or weekly salinity and productivity data are not available from the SO, and daily oceanic temperatures are limited to the water surface. Other variables show inconsistent and incomplete spatial coverage year-round. For example, Chl-a data are highly patchy and limited to summer months, which constrains its use in year-round dynamic models.

Averaging (during the calculation of predictors) over highly varying environments can diminish the influence of environmental variability on the model, possibly leading to over- or under-prediction (Zimmermann et al., 2009). We attempted to diminish the impact of temporal mismatch between sightings and environmental conditions by including the environmental temporal variation (standard deviation) where appropriate. Seasonal variability in combinations with means can express extreme conditions and improve models’ predictive power (Zimmermann et al., 2009). Nevertheless, we emphasize that incorporating environmental variability in static SDMs is inevitably insufficient to capture the SO’s high dynamics. For example, although seasonal SIC variability had high importance in our models, it is unfit to determine species preference to sea ice, and its response curve is hard to interpret. Hence, including predictors representing environmental variabilities may improve predictions, but they are fall short of explaining. Further, summarizing some highly dynamic variables can be challenging, even on a seasonal or monthly scale. For example, we estimated distance to seasonal SIE from seasonal mean SIC. However, sea ice cover (and SIE with it) varies at a high temporal (daily) frequency. Therefore, a single line describing long-term average SIE is not a good representative of the true SIE in any season or month (Figure 3 and Figure S26–S27). Although we found high importance of SIC and distance to SIE in summer, relating species observations to their concomitant environmental conditions should be of higher priority in SDMs (Figure S28).

5 | CONCLUSION

In this study, we used presence-only SDMs (Maxent) to model the circumantarctic habitat of four baleen whale species and identified important predictors affecting their distribution in the Southern Ocean. Model performance was high (Table S2), with generally little predicted cross-validated uncertainty. Unsurprisingly, models identified sea ice-derived predictors and distance to continental shelf break as the main predictors. The indispensable role of sea ice in the lives of many Antarctic species, particularly krill-dependent predators, makes whale species sensitive to future changes in the distribution and the dynamics of the sea ice (Herr et al., 2019; Leaper & Miller, 2011; Thiele et al., 2004). Such environmental change signals have already been reported from polar regions, for example, the warming of the West Antarctic Peninsula area (Gutt et al., 2015; Vaughan et al., 2003), and the predicted shrinkage of sea ice in the Antarctic under all future climate change scenarios (Gutt et al., 2015; Leaper & Miller, 2011; Solomon et al., 2007). This emphasizes the need for more studies on the spatio-temporal distribution of baleen whales in the SO to understand the potential impact of climate change on these species. We compared our species-specific results with results of other studies in the SO and provided reasons for results discrepancy, which is generally attributed to the use of different species data quality and quantity, different study area extent and methodological reasons.

Maxent is known for its high predictive accuracy and considered one of the most frequently used technique in marine SDM studies (Melo-Merino et al., 2020). Our models back the usefulness of presence-only SDMs like Maxent as a cost-effective tool for studying the distribution of migratory whales (e.g. Smith et al., 2020). The current work further supports the pivotal role of crowdsourcing data from biodiversity repositories (e.g. GBIF and OBIS) and circumantarctic dedicated surveys (e.g. SO GLOBEC and SOWER) to strengthen our knowledge about the distribution and niche of migratory whales in less-surveyed oceans (Beekmans et al., 2010). Nevertheless, future surveys should be prioritized towards less studied areas and the pack ice region, especially beyond the summer months. Alternative data sources, such as PAM and from tagged animals, form a useful addition for studying marine mammals’ habitat preferences year-round, but still require work before these data can be integrated. PAM is particularly useful in the SO for detecting rarely visually sighted species like ABW and covering difficult-to-access areas (e.g. the ice-covered Weddell Sea). PAM data have already been used in SDMs for odontocete species producing clicks which propagate over short distances allowing to use environmental
data from the recording sites (e.g. Gallus et al., 2012; Soldevilla et al., 2011). To date, only few applications have included baleen whales of which calls propagate over long distances causing uncertainty in the interpretation of the relationship between whales and the environment due to this potential mismatch in scales (e.g. Širović & Hildebrand, 2011; Stafford et al., 2009). Nevertheless, the use of PAM data in SDMS, particularly for species in polar waters, holds great potential that calls for exploring this further.

ACKNOWLEDGEMENTS

This work was financially supported by the German Federal Ministry of Food and Agriculture (BLE) through the Federal Office for Agriculture and Food (BLE), grant number 2817HS004. We thank Andy Traumüller for helping in environmental variables preparation and all data providers for making their observation data publicly available, especially captains and nautical officers onboard Polarstern. An earlier version of this manuscript was improved by comments of Alaaeldin Soultan. This work was partially performed on the HPC computer facility of the Alfred Wegener Institute. We thank Natalja Rakowsky for technical support on the HPC computers. Open Access funding enabled and organized by Projekt DEAL.

PEER REVIEW

The peer review history for this article is available at https://publo ns.com/publon/10.1111/ddi.13300.

DATA AVAILABILITY STATEMENT

The dataset analysed during the current study is already publicly available (details in Appendix S1–S5). The interactive distribution of cleaned occurrences is available at this web application: https://awi oza.shinyapps.io/BaleenWhales/.

ORCID

Ahmed El-Gabbas https://orcid.org/0000-0003-2225-088X
Ilse Van Opzeeland https://orcid.org/0000-0001-8369-7234
Elke Burkhardt https://orcid.org/0000-0002-5128-4176
Olaf Boebel https://orcid.org/0000-0002-2259-0035

REFERENCES

Aiello-Lammens, M. E., Boria, R. A., Radosavljevic, A., Vilela, B., & Anderson, R. P. (2015). spThin: An R package for spatial thinning of species occurrence records for use in ecological niche models. Ecography, 38(5), 541–545. https://doi.org/10.1111/eco.01132
Ainley, D. G., Jacobs, S. S., Ribic, C. A., & Gaffney, I. (2004). Seabird distribution and oceanic features of the Amundsen and southern Bellingshausen seas. Antarctic Science, 10(2), 111–123. https://doi.org/10.1017/s0954102098000169
Ainley, D. G., Jongsomjit, D., Ballard, G., Thiele, D., Fraser, W. R., & Tynan, C. T. (2012). Modeling the relationship of Antarctic minke whales to major ocean boundaries. Polar Biology, 35(2), 281–290. https://doi.org/10.1007/s00300-011-1075-1
Andrews-Goff, V., Bestley, S., Gales, N. J., Laverick, S. M., Paton, D., Polanowski, A. M., Schmitt, N. T., & Double, M. C. (2018). Humpback whale migrations to Antarctic summer foraging grounds through the southwest Pacific Ocean. Scientific Reports, 8(1), 12333. https://doi.org/10.1038/s41598-018-30748-4

Barbet-Massin, M.,Thuiller, W., & Jiguet, F. (2010). How much do we overestimate future local extinction rates when restricting the range of occurrence data in climate suitability models? Ecography, 33(5), 878–886. https://doi.org/10.1111/j.1600-0587.2010.06181.x
Barlow, J., Gerodete, T., & Forcada, J. (2001). Factors affecting perpendicular sighting distances on shipboard line-transect surveys for cetaceans. Journal of Cetacean Research and Management, 3, 201–212.
Bateman, B. L., VanDerWal, J., & Johnson, C. N. (2012). Nice weather for bettongs: Using weather events, not climate means, in species distribution models. Ecography, 35(4), 306–314. https://doi.org/10.1111/j.1600-0587.2011.06871.x
Becker, E., Forney, K., Fiedler, P., Barlow, J., Chivers, S., Edwards, C., Moore, A., & Redfern, J. (2016). Moving towards dynamic ocean management: How well do modeled ocean products predict species distributions? Remote Sensing, 8(2), 149. https://doi.org/10.3390/rs8020149
Beekmans, B. W. P. M., Forcada, J. F., Murphy, E. J., de Baar, H. J. W., Bathmann, U., & Fleming, A. H. (2010). Generalised additive models to investigate environmental drivers of Antarctic minke whale (Balaenoptera bonaerensis) spatial density in austral summer. Journal of Cetacean Research and Management, 11(2), 115–129.
Bomboesch, A., Zitterbart, D. P., Van Opzeeland, I., Frickenhaus, S., Burkhardt, E., Wisz, M. S., & Boebel, O. (2014). Predictive habitat modelling of humpback (Megaptera novaeangliae) and Antarctic minke (Balaenoptera bonaerensis) whales in the Southern Ocean as a planning tool for seismic surveys. Deep Sea Research Part I: Oceanographic Research Papers, 91, 101–114. https://doi.org/10.1016/j.dsr.2014.05.017
Branch, T. A. (2011). Humpback whale abundance south of 60°S from three complete circumpolar sets of surveys. Journal of Cetacean Research and Management, 3, 53–69.
Branch, T. A., Matsuoka, K., & Miyashita, T. (2004). Evidence for increases in Antarctic blue whales based on Bayesian modelling. Marine Mammal Science, 20(4), 726–754. https://doi.org/10.1111/j.1748-7692.2004.tb01190.x
Branch, T. A., Stafford, K. M., Palacios, D. M., Allison, C., Bannister, J. L., Burton, C. L. K., Cabrera, E., Carlson, C. A., Galletti vernazzani, B., Gill, P. C., Huckle-gaete, R., Jenner, K. C. S., Jenner, M.-N.- M., Matsuoka, K., Mikhailov, Y. A., Miyashita, T., Morrice, M. G., Nishiwaki, S., Sturrock, V. J., ... Warneke, R. M. (2007). Past and present distribution, densities and movements of blue whales Balaenoptera musculus in the Southern Hemisphere and northern Indian Ocean. Mammal Review, 37(2), 116–175. https://doi.org/10.1111/j.1365-2907.2007.00106.x
Brierley, A. S., Fernandes, P. G., Brandon, M. A., Armstrong, F., Millard, N. W., McPhail, S. D., & Griftiths, G. (2002). Antarctic krill under sea ice: Elevated abundance in a narrow band just south of ice edge. Science, 295(5561), 1890–1892. https://doi.org/10.1126/science.1068574
Convey, P., Chown, S. L., Clarke, A., Barnes, D. K. A., Bokhorst, S., Cummings, V., Ducklow, H. W., Frati, F., Green, T. G. A., Gordon, S., Griffiths, H. J., Howard-Williams, C., Huiskes, A. H. L., Laybourn-Parry, J., Lyons, W. B., McMinn, A., Morley, S. A., Peck, L. S., Quesada, A., ... Wall, D. H. (2014). The spatial structure of Antarctic biodiversity. Ecological Monographs, 84(2), 203–244. https://doi.org/10.1890/12-2216.1
Dalla Rosa, L., Secchi, E. R., Maia, Y. G., Zerbini, A. N., & Heide-Jørgensen, M. P. (2008). Movements of satellite-monitored humpback whales on their feeding ground along the Antarctic Peninsula. Polar Biology, 31(7), 771–781. https://doi.org/10.1007/s00300-008-0415-2
Dambach, J., & Rödder, D. (2011). Applications and future challenges in marine species distribution modeling. Aquatic Conservation: Marine and Freshwater Ecosystems, 21(1), 92–100. https://doi.org/10.1002/ac.1160
Dayton, P. K., Mordida, B. J., & Bacon, F. (1994). Polar marine communities. American Zoologist, 34(1), 90–99. https://doi.org/10.1093/ icb/34.1.90
De Broyer, C., Koubbi, P., Griffiths, H. J., Raymond, B., Udekkem d’Acouz, C., D., Van de Putte, A. P., Danis, B., David, B., Grant, S., Gutt, J., Held, C., Hosie, G., Huettmann, F., Post, A., & Ropert-Coudert, Y. (Eds.). (2014). Biogeographic Atlas of the Southern Ocean. SCAR. https://www.biodiversity.org/atais/
deStephanis, R., Cornulier, T., Verborgh, P., Salazar Sierra, J., Gimeno, N. P., & Guinet, C. (2008). Summer spatial distribution of cetaceans in the Strait of Gibraltar in relation to the oceanographic context. Marine Ecology Progress Series, 352, 275–288. https://doi.org/10.3354/meps07164
Dominello, T., & Širović, A. (2016). Seasonality of Antarctic minke whale (Balaenoptera bonaerensis) calls off the western Antarctic Peninsula. Marine Mammal Science, 32(3), 826–838. https://doi.org/10.1111/mms.12302
Dormann, C. F., & Kaschner, K. (2010). Where’s the sperm whale? A species distribution example analysis. http://www.mcderm-ecolgy.org/?page_id=355
Dormann, C. F., Schymanski, S. J., Cabral, J., Chuiine, I., Graham, C., Hartig, F., Kearney, M., Morin, X., Römermann, C., Schröder, B., & Singer, A. (2012). Correlation and process in species distribution models: Bridging a dichotomy. Journal of Biogeography, 39(12), 2119–2131. https://doi.org/10.1111/j.1365-2699.2011.02659.x
Double, M. C., Miller, B. S., Leaper, R., Olson, P., Cox, M. J., Miller, E., Calder, S., Collins, K., Donnelly, D., Ensor, P., Goetz, K., Schmitt, N., Andrews-Goff, V., Bell, E., & O’Driscoll, R. (2015). Cruise report on blue whale research from the NZ/Aus Antarctic Ecosystems Voyage 2015 of the Southern Ocean Research Partnership. Paper SC/66A/SH/7 presented to the IWC Scientific Committee. https://iwc.int/
El- Gabbas, A., & Dormann, C. F. (2018a). Improved species-occurrence predictions in data-poor regions: Using large-scale data and bias correction with down-weighted Poisson regression and Maxent. Ecography, 41(7), 1161–1172. https://doi.org/10.1111/ecog.03149
El- Gabbas, A., & Dormann, C. F. (2018b). Wrong, but useful: Regional species distribution models may not be improved by range-wide data under biased sampling. Ecology and Evolution, 8(4), 2196–2206. https://doi.org/10.1002/ece3.3834
Ensor, P. H. (1989). Minke whales in the pack ice zone, East Antarctica, during the period of maximum annual ice extent. https://iwc.int/
Esteban, R., Verborgh, P., Gauffier, P., Giménez, J., Afán, I., Cañadas, A., García, P., Murcia, J. L., Magalhães, S., Andreu, E., & de Stephanis, R. (2013). Identifying key habitat and seasonal patterns of a critically endangered population of killer whales. Journal of the Marine Biological Association of the United Kingdom, 94(6), 1317-1325. https://doi.org/10.1017/s002531541300091x
Fabri-Ruiz, S., Danis, B., David, B., Saucède, T., & Treml, E. (2019). Can we generate robust species distribution models at the scale of the Southern Ocean? Diversity and Distributions, 25(1), 21–37. https://doi.org/10.1111/ddi.12835
Fernandez, M., Yesson, C., Gannier, A., Miller, P. I., & Azevedo, J. M. N. (2017). The importance of temporal resolution for niche modelling in dynamic marine environments. Journal of Biogeography, 44(12), 2816–2827. https://doi.org/10.1111/jbi.13080
Flun, D., Thomisch, K., Boebel, O., Brey, T., Sirovic, A., Spiesecce, S., & Van Opzeeland, I. (2020). Frozen verses: Antarctic minke whale (Balaenoptera bonaerensis) calls predominantly during austral winter. Royal Society Open Science, 7(10). https://doi.org/10.1098/rsos.192112
Frischauer, B. J., Hofmann, E. E. (2003). A predator’s perspective on causal links between climate change, physical forcing and ecosystem response. Marine Ecology Progress Series, 265, 1–15. https://doi.org/10.3354/meps265001
Friedlaender, A. S., Goldbogen, J. A., Nowacek, D. P., Read, A. J., Johnston, D., & Gales, N. (2014). Feeding rates and under-ice foraging strategies of the smallest lunge filter feeder, the Antarctic minke whale (Balaenoptera bonaerensis). The Journal of Experimental Biology, 217(Pt 16), 2851–2854. https://doi.org/10.1242/jeb.106682
composition in the Weddell Sea, Antarctica. Marine Ecology Progress Series, 638, 191–206. https://doi.org/10.3354/meps13258
Van Opzeeland, I., Van Parijs, S., Kindermann, L., Burkhardt, E., & Boebel, O. (2013). Calling in the cold: Pervasive acoustic presence of humpback whales (Megaptera novaeangliae) in Antarctic coastal waters. PLoS One, 8(9), e73007. https://doi.org/10.1371/journal.pone.0073007
Vaughan, D. G., Marshall, G. J., Connolley, W. M., Parkinson, C., Mulvaney, R., Hodgson, D. A., King, J. C., & Pudsey, C. J. (2003). Recent rapid regional climate warming on the Antarctic Peninsula. Climatic Change, 60(3), 243–274. https://doi.org/10.1023/A:1026021217991
Wackernagel, H. (1995). Ordinary Kriging. In H. Wackernagel (Ed.), Multivariate geostatistics (pp. 74–81). Springer.
Weatherall, P., Marks, K. M., Jakobsson, M., Schmitt, T., Tani, S., Arndt, J. E., Rovere, M., Chayes, D., Ferrini, V., & Wrigley, R. (2015). A new digital bathymetric model of the world’s oceans. Earth and Space Science, 2(8), 331–345. https://doi.org/10.1002/2015ea000107
Wege, M., Salas, L., & LaRue, M. (2020). Citizen science and habitat modelling facilitates conservation planning for crabeater seals in the Weddell Sea. Diversity and Distributions, 26(10), 1291–1304. https://doi.org/10.1111/ddi.13120
Williams, R., Hedley, S. L., & Hammond, P. S. (2006). Modeling distribution and abundance of Antarctic baleen whales using ships of opportunity. Ecology and Society, 11(1). https://doi.org/10.5751/ES-01534-110101
Williams, R., Kelly, N., Boebel, O., Friedlaender, A. S., Herr, H., Kock, K.-H., Lehner, L. S., Maksym, T., Roberts, J., Scheidat, M., Siebert, U., & Bierley, A. S. (2014). Counting whales in a challenging, changing environment. Scientific Reports, 4, 4170. https://doi.org/10.1038/srep04170
Zimmermann, N. E., Yoccoz, N. G., Edwards, T. C., Meier, E. S., Thuiller, W., Guisan, A., Schmatz, D. R., & Pearman, P. B. (2009). Climatic extremes improve predictions of spatial patterns of tree species. Proceedings of the National Academy of Sciences of the United States of America, 106(Suppl 2), 19723–19728. https://doi.org/10.1073/pnas.0901643106
Zurell, D., Elith, J., & Schröder, B. (2012). Predicting to new environments: Tools for visualizing model behaviour and impacts on mapped distributions. Diversity and Distributions, 18(6), 628–634. https://doi.org/10.1111/j.1472-4642.2012.00887.x
Zuur, A. F., Ieno, E. N., & Elphick, C. S. (2010). A protocol for data exploration to avoid common statistical problems. Methods in Ecology and Evolution, 1(1), 3–14. https://doi.org/10.1111/j.2041-210X.2009.00001.x
Zweng, M. M., Reagan, J. R., Seidov, D., Boyer, T. P., Locarnini, R. A., Garcia, H. E., Mishonov, A. V., Baranova, O. K., Weathers, K. W., Paver, C. R., & Smolyar, I. V. (2019). A. Mishonov (Ed.), World Ocean Atlas 2018, Volume 2: Salinity (50 pp). A. Mishonov Technical Ed. NOAA Atlas NESDIS 82. https://www.ncei.noaa.gov/sites/default/files/2020-04/woa18_vol2.pdf

BIOSKETCH
The authors are members of the ocean acoustics group at the Alfred Wegener Institute, Helmholtz Centre for Polar and Marine Research. They are interested in exploring the distribution and behaviour of marine mammals and their underwater acoustic environment in the polar oceans using long-term (multi-year) visual observations and passive acoustic recordings. They aim at providing reliable information on year-round distributions of marine mammals in polar oceans to inform conservation, management and science. They are particularly interested in using static and dynamic species distribution models in polar areas using visual and PAM data.

Author contributions: All authors conceived the research; A.E.-G. performed data curation, planned and conducted the analyses; A.E.-G. wrote the first draft, with all authors contributing to the writing of the final manuscript.

SUPPORTING INFORMATION
Additional supporting information may be found online in the Supporting Information section.

How to cite this article: El-Gabbas A, Van Opzeeland I, Burkhardt E, Boebel O. Static species distribution models in the marine realm: The case of baleen whales in the Southern Ocean. Divers Distrib. 2021;00:1–17. https://doi.org/10.1111/ddi.13300