Dynamic ensemble models to predict distributions and anthropogenic risk exposure for highly mobile species

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Abstract
Aim: Advances in ecological and environmental modelling offer new opportunities for estimating dynamic habitat suitability for highly mobile species and supporting management strategies at relevant spatiotemporal scales. We used an ensemble modelling approach to predict daily, year-round habitat suitability for a migratory species, the blue whale (Balaenoptera musculus), and demonstrate an application for evaluating the spatiotemporal dynamics of their exposure to ship strike risk.

Location: The California Current Ecosystem (CCE) and the Southern California Bight (SCB), USA.

Methods: We integrated a long-term (1994–2008) satellite tracking dataset on 104 blue whales with data-assimilative ocean model output to assess year-round habitat suitability. We evaluated the relative utility of ensembling multiple model types compared to using single models, and selected and validated candidate models using multiple cross-validation metrics and independent observer data. We quantified the spatial and temporal distribution of exposure to ship strike risk within shipping lanes in the SCB.

Results: Multi-model ensembles outperformed single-model approaches. The final ensemble model had high predictive skill (AUC = 0.95), resulting in daily, year-round predictions of blue whale habitat suitability in the CCE that accurately captured migratory behaviour. Risk exposure in shipping lanes was highly variable within and among years as a function of environmental conditions (e.g., marine heatwave).

Main conclusions: Daily information on three-dimensional oceanic habitats was used to model the daily distribution of a highly migratory species with high predictive power and indicated that management strategies could benefit by incorporating dynamic environmental information. This approach is readily transferable to other species. Dynamic, high-resolution species distribution models are valuable tools for assessing risk exposure and targeting management needs.
1 | INTRODUCTION

The ocean provides a diverse and extensive suite of ecosystem services, all of which ultimately depend on a functioning ecosystem. Management strategies that enable human activity while conserving biodiversity remains a key challenge for ocean governance. Marine spatial planning implicitly addresses such trade-offs and as such has become a valuable approach for regulating multiple ocean activities while achieving conservation targets (Foley et al., 2010; White, Halpern, & Kappel, 2012). However, marine spatial planning is often static, despite recognition that marine habitats tend to be highly dynamic and can shift in space on timescales of days to weeks (Checkley & Barth, 2009; Kavanaugh et al., 2016). Thus, static management strategies do not account for shifting habitats or human activities and, importantly, offer only partial protection for highly mobile species (Dunn, Maxwell, Boustany, & Halpin, 2016; Hazen et al., 2018). Dynamic management has been identified as a potential solution to this problem by allowing management decisions to be updated in space and time in response to changing environmental or socioeconomic conditions (Lewison et al., 2015; Maxwell et al., 2015). In order to inform the ecological components of dynamic management, there is first a need to accurately describe the spatiotemporal distribution of species and populations (Foley et al., 2010).

Species distribution models (SDMs) are key tools for describing species habitats and distributions across marine and terrestrial systems (Elith & Leathwick, 2009; Robinson, Nelson, Costello, Sutherland, & Lundquist, 2017). Species distribution modelling involves using statistical tools to relate species occurrence or abundance to spatiotemporal patterns of environmental variation (Elith & Leathwick, 2009). Though the designs, uses and applications of SDMs in ecology are diverse, two methodological advancements hold particular promise for dynamic ocean management.

First, while marine SDMs have typically used a single-model type (Robinson et al., 2017), determining an appropriate modelling approach can be challenging given inherent trade-offs in the statistical methods available (Elith et al., 2006; Qiao, Soberón, & Peterson, 2015). Multi-model ensembles can reduce uncertainty by overcoming biases inherent in any one model type and providing a “consensus” approach to predictions (Araújo & New, 2007; Gritti, Duputié, Massol, & Chuine, 2013). Predictions generated from multi-model ensembles can also be more accurate than those of single models (Marmion, Parviainen, Luoto, Heikkinen, & Thuiller, 2009; Oppel et al., 2012; Scales et al., 2015). As a result, ensemble model approaches are increasingly recommended for marine species distribution modelling (Jones & Cheung, 2014; Robinson et al., 2017), motivating further evaluation of their application to dynamic management.

Second, data-assimilative ocean circulation models offer the opportunity to better match species data to the underlying environmental processes at relevant spatiotemporal scales (Becker et al., 2016; Brodie et al., 2018). There has been increasing attention paid in the marine realm to the spatiotemporal scales of environmental observations and their relevance to the scales of animal response (Mannocci et al., 2017; Scales et al., 2016). Frequently, there is a mismatch between the spatiotemporal resolution of species occurrence and the spatiotemporal resolution of environmental observations. Such mismatch can lead to incorrect inferences and increased uncertainty (Scales et al., 2016). Moreover, designing dynamic management strategies for highly mobile species often relies on up-to-date, or “real-time,” estimates of species distributions (Laist, Knowlton, & Pendleton, 2014). Data-assimilative ocean circulation models can help solve this mismatch by providing gapless environmental data, often with higher spatial or temporal resolutions than those of processed remotely sensed data (Becker et al., 2016; Brodie et al., 2018). Ocean circulation models can also provide information about the vertical structure of the ocean that remotely sensed variables cannot, which can improve predictions of marine species’ distributions (Brodie et al., 2018).

Blue whales (Balaenoptera musculus) are both a highly migratory species and a species of conservation concern, highlighting the need to understand their habitat use and exposure to potential anthropogenic threats throughout their migrations. Blue whales are listed as Endangered under both the U.S. Endangered Species Act (1973) and the IUCN Red List of Threatened Species due to population depletion from commercial whaling (Reilly et al. 2008). In the Northeast Pacific, blue whales perform latitudinal migrations between tropical wintering/breeding grounds and productive foraging grounds at higher latitudes in the California Current Ecosystem (CCE) (Bailey et al., 2009; Balance, Pitman & Fiedler, 2006; Irvine et al., 2014; Mate, Lagerquist, & Calambokidis, 1999). While in the CCE, blue whales follow the spring and summertime progression of the availability of krill (Abrahms et al., 2019), their primary prey, and demonstrate temporal synchrony with krill availability (Croll et al. 2005; Fossette et al. 2017). While dynamic distribution data on krill are not available at the requisite spatial and temporal scales of our study, previous studies have shown that blue whale habitat in the CCE can be characterized by a combination of dynamic and static environmental characteristics, such as sea surface temperature, thermocline and seafloor depths and primary productivity (Becker et al., 2016; Hazen et al., 2017).

Though the eastern North Pacific blue whale population is recovering (Monnahan, Branch, & Punt, 2014), mortality from ship strikes in the CCE remains a major management concern (Rockwood,.....
Calambokidis, & Jahncke, 2017). In particular, the Southern California Bight (SCB; from San Diego to Point Conception, ~33–34.5°N) is a hotspot for ship strikes due to the high spatial and temporal overlap between blue whale summer foraging hotspots and shipping vessels travelling to and from southern California’s largest ports (Hazen et al., 2017; Redfern et al., 2013; Rockwood et al., 2017). Currently, voluntary seasonal slowdowns are the only tool used to manage ship strike risk in this region (Rockwood et al., 2017). The existence of multiple shipping lanes for marine traffic offers potential opportunity for redirecting vessels based on strike risk (Redfern et al., 2013), but existing blue whale models for the region are either limited by partial temporal coverage (Becker et al., 2018) or by coarse resolution (Hazen et al., 2017). In addition to better matching the timescales of species occurrences, finer scale distribution estimates may be better able to inform dynamic management decisions in the SCB given the migratory behaviour of blue whales (Bailey et al., 2009) and the variable biophysical conditions in the CCE (Checkley & Barth, 2009).

We used a multi-year (1994–2008) satellite tracking dataset on 104 blue whales along with data-assimilative ocean model output to develop a daily, year-round distribution model for blue whales in the CCE. We evaluated the utility of using multi-model ensembles relative to single-model approaches and used the largest compilation of independent blue whale sightings datasets to date to select and validate our final species distribution model. Finally, we quantified the spatial and temporal distribution of exposure to ship strike risk within shipping lanes in the SCB.

2 | METHODS

2.1 | Movement data

Argos-linked satellite tags were deployed between 1994 and 2008 on 104 blue whales off of the Costa Rica Dome (N = 2), the Gulf of California (N = 3), the Santa Barbara Channel (N = 66) and northern California coast (N = 33) (Bailey et al., 2009; Hazen et al., 2017; Irvine et al., 2014). Duration of tracks averaged 101 ± 85 days (mean ± SD). All tracks were filtered for errors and smoothed to provide daily position estimates using a Bayesian switching state-space model (Bailey et al., 2009; Jonsen, Flemming, & Myers, 2005), resulting in 10,603 daily locations (Figure S1). Information on tag types and deployment duration is available in Bailey et al. (2009).

Because presence-absence models outperform presence-only models for species distribution modelling (Elith et al., 2006), use of artificial absence data (i.e., “pseudo-absences”) is recommended when true absence data are unavailable (Barbet-Massin, Jiguet, Albert, & Thuiller, 2012). To analyse habitat suitability in a presence-absence framework, we generated pseudo-absences following Hazen et al. (2017) by simulating 200 correlated random walks per whale track using empirical step-length and turn-angle distributions (Kareiva & Shigesada, 1983). This approach enabled matching the error structures of pseudo-absences and empirical movement data (Hazen et al., 2017; Jonsen et al., 2005; Scales et al., 2016; Willis-Norton et al., 2015). A flag value was assigned to each simulated track indicating its similarity to the empirical track based on distance and net angular displacement from the empirical track (Hazen et al., 2017; Willis-Norton et al., 2015). To ensure simulated tracks represented areas accessible to the whales, only simulated tracks in the upper 75th percentile of flag values were used for comparison (see Hazen et al., 2017 for detail). Two simulated tracks per empirical track were selected at random for inclusion in further analyses, resulting in 21,328 pseudo-absence points compared to 10,603 presence points (Barbet-Massin et al., 2012; Figure S2).

2.2 | Environmental data

Presence and pseudo-absence data were matched to dynamic surface and subsurface environmental variables as well as to static seafloor relief variables. Variables were examined based on hypothesized drivers of habitat and those shown to be significantly associated with blue whale space use (Becker et al., 2016, 2018; Hazen et al., 2017). Daily environmental data at 0.1° resolution were sourced from historical and near-real-time data-assimilative versions of the Regional Ocean Modelling System (ROMS) configured for the CCE (Moore et al., 2013; Neveu et al., 2016) obtained from oceanmodeling.ucsc.edu). Dynamic surface variables examined were sea surface temperature (SST), SST standard deviation (SST_sd), sea surface height (SSH), SSH standard deviation (SSH_sd), eddy kinetic energy (EKE) and wind stress curl. Dynamic subsurface variables were isothermal layer depth (ILD), similar to mixed layer depth and defined by a 0.5°C deviation from the SST, and bulk Brun Väisälä frequency (BVF), a measure of stratification averaged over the upper 200 m of the water column. In addition, the following static seafloor relief variables were sourced from ETOP01 (obtained from https://www.ngdc.noaa.gov/mgg/global/global.html; 0.1-degree resolution): bathymetry (z), standard deviation of bathymetry (z_sd), slope and aspect. Standard deviations of SST, SSH and bathymetry for each location were calculated using a 1° radius centred on that location. Our study area was matched to the ROMS model domain (30 to 48°N and from the coast to 134°W).

2.3 | Species distribution modelling

Given potential differences in explanatory power and predictive skill (Derville, Torres, Iovan, & Garrigue, 2018; Fiedler et al., 2018), species distribution models were built using both Generalized Additive Mixed Models (GAMMs; “mgcv” R package) (Wood, 2017) and Boosted Regression Trees (BRTs; “dismo” R package) (Elith, Leathwick, & Hastie, 2008). Because seasonality in migratory behaviour has been shown to influence blue whale environmental preferences (Hazen et al., 2017), for both GAMMs and BRTs we explored year-round models as well as separate models for each migratory season (summer/fall—July–November; winter/spring—December–June). GAMMs were fitted using the binomial family and a logit link function, with individuals nested as a random effect. A tensor product smooth between latitude and longitude was explored as predictors in GAMMs to account for spatial autocorrelation (Becker et al., 2018). A latitudinal interaction term with SST was also considered in the GAMMs to account for the latitudinal temperature
gradient over the study area (Becker et al., 2018). Multiple candidate GAMM models were generated based on published and hypothesized predictor variable combinations (Becker et al., 2016; Hazen et al., 2017) and were initially evaluated using the Area Under the receiver operating Curve (AUC; Table S1) before selecting the top three GAMMs for further model evaluation. AUC statistics discriminate between true-positive and false-positive rates, and range from 0 to 1, where a score of >0.5 indicates better than random discrimination (Hanley & McNeil, 1982). Generalized variance inflation factors were used to ensure any collinear predictor variables were not included in the same GAMM. As such, SST and BVF were not included together in candidate GAMMs. A Bernoulli family distribution was used for BRTs, in which all environmental variables were included since BRTs can handle irrelevant predictors and any collinearity effects (Elith et al., 2008). GAMM-BRT ensemble model combinations were explored by ensembling the seasonal BRTs with each of the top three performing seasonal GAMMs. Each model type was given an equal weighting in the ensembles.

Because of the importance of using multiple metrics for SDM evaluation (Fourcade, Besnard, & Secondi, 2017), predictive performance for the BRTs, top three GAMMs, and ensemble model combinations was further evaluated using AUC and True Skill Statistic (TSS) metrics on three training and testing dataset combinations: (a) k-fold cross-validation with a 75%/25% training/testing data split over each of five folds, (b) "Leave One Out" cross-validation in which a year of data was iteratively left out from training and retained for testing and (c) the full tagging dataset tested on an independent blue whale sightings dataset (N = 3,413 observations; Table S3, Figures S3 and S4). To calculate AUC and TSS metrics for testing against sightings data, pseudo-absences were randomly generated at a 1:3 presence:absence ratio. As independent testing is recommended over cross-validation for evaluating SDM performance (Derville et al., 2018; Gregr, Palacios, Thompson, & Chan, 2018), the final model was chosen based on the sightings data metric averaged across seasons. Finally, we calculated the point biserial correlation between the final model's predictions and independent sightings versus pseudo-absences (Elith et al., 2006). Daily spatial predictions of blue whale habitat suitability were mapped onto the 0.1° gridded study domain using the "raster" R package.

### TABLE 1

Top seasonal GAMM, BRT and ensemble candidate models and diagnostic metrics (AUC/TSS) compared to independent sightings data for each season and averaged across seasons. Metrics for additional training/testing methods are provided in Table S1. Variable acronyms refer to sea surface temperature (SST), bathymetry (z), sea surface height (SSH), SSH standard deviation (SSH_sd), bathymetry standard deviation (z_sd), isothermal layer depth (ILD), bulk Brunt Väisälä frequency (BVF), and eddy kinetic energy (EKE). All GAMMs include a random effect for individual. The final model is highlighted in bold.

<table>
<thead>
<tr>
<th>Model name</th>
<th>Model description</th>
<th>Winter/spring</th>
<th>Summer/fall</th>
<th>Seas. Avg.</th>
</tr>
</thead>
<tbody>
<tr>
<td>GAMM 1</td>
<td>SST + SSH_sd + z + z_sd + ILD + EKE</td>
<td>0.974/0.846</td>
<td>0.908/0.755</td>
<td>0.941/0.801</td>
</tr>
<tr>
<td>GAMM 2</td>
<td>SST + SSH_sd + z + z_sd + ILD + EKE + lat*lon</td>
<td>0.914/0.724</td>
<td>0.892/0.727</td>
<td>0.903/0.706</td>
</tr>
<tr>
<td>GAMM 3</td>
<td>SST*lat + SSH_sd + z + z_sd + ILD + EKE</td>
<td>0.945/0.777</td>
<td>0.896/0.722</td>
<td>0.921/0.750</td>
</tr>
<tr>
<td>BRT</td>
<td>SST + SST_sd + SSH + SSH_sd + z + z_sd + ILD + EKE + curl + BVF + slope + aspect</td>
<td>0.983/0.880</td>
<td>0.903/0.704</td>
<td>0.943/0.792</td>
</tr>
<tr>
<td>Ensemble 1</td>
<td>GAMM 1 + BRT</td>
<td>0.985/0.875</td>
<td>0.913/0.732</td>
<td>0.949/0.804</td>
</tr>
<tr>
<td>Ensemble 2</td>
<td>GAMM 2 + BRT</td>
<td>0.972/0.852</td>
<td>0.910/0.722</td>
<td>0.941/0.787</td>
</tr>
<tr>
<td>Ensemble 3</td>
<td>GAMM 3 + BRT</td>
<td>0.978/0.854</td>
<td>0.906/0.725</td>
<td>0.941/0.790</td>
</tr>
</tbody>
</table>

### 2.4 Risk exposure in shipping lanes

Based on the output of the top-performing model or ensemble, we compared predicted habitat suitability within shipping lanes inside and outside of the Santa Barbara Channel in the SCB over the course of a year to evaluate spatiotemporal patterns in risk exposure to ship strikes. Vessels travelling into the SCB use either established routes inside the Santa Barbara Channel ("Northern approach") or a Western approach outside of the channel, following the implementation of an "Ocean-Going Vessel Fuel Rule" in 2009 that resulted in increased traffic using the Western approach (Redfern et al., 2013). Spatial layers for the alternate routes were provided by the Channel Islands National Marine Sanctuary. The average of predicted habitat suitability in all grid cells intersecting each lane was calculated for each day of the year in 2009 and 2015 to compare a year with average environmental conditions (2009) to a year in which a climatic anomaly occurred associated with a prolonged marine heatwave (2015) (Bond, Cronin, Freeland, & Mantua, 2015; Di Lorenzo & Mantua, 2016; Jacox et al., 2018, 2016).

All analyses were performed using R statistical computing (R Core Team, 2017).

### 3 RESULTS

#### 3.1 Species distribution model

Dynamic ensemble modelling for blue whales revealed that blue whale habitat use in the CCE is strongly influenced by temperature, seafloor topography and subsurface water properties (Table 1, Figures 1 and 2). Seasonal models outperformed year-round models on average for both GAMMs and BRTs (Table S2), and top models showed high predictive performance, with AUC scores using tracking and independent sightings datasets ranging from 0.84 to 0.99, and TSS scores ranging from 0.55 to 0.89 (Tables 1 and S1). The highest performing GAMM included SST, SSH_sd, z, z_sd, ILD and EKE (Table 1, Figure 1). The average explained deviance, a measure of descriptive performance, for the seasonal GAMMs was 43.4%. BRTs did not undergo variable selection and included all environmental
covariates, though SST, SSH_sd, z, z_sd and ILD contributed the highest explained deviance (Table 1, Figure 2, Figures S5 and S6). The average explained deviance for the seasonal BRTs was 60.1%.

GAMMs and BRTs showed general agreement in habitat preferences. For both model types, some differences in response curves between seasons were apparent: in winter/spring, habitat suitability was associated with SST >15°C, high SST standard deviations indicating thermal front activity, shallower ILDs (<50 m), low BVFs indicating weak stratification, seafloor depths <3,000 m, and areas of high seafloor ruggedness as measured by standard deviation of bathymetry (Figures 1 and 2, Figure S5). In summer/fall, habitat suitability was associated with SST between 16 and 20°C, weak stratification, shallower seafloor depths (<2,000 m), high seafloor ruggedness and high wind stress curl (Figures 1 and 2, Figure S6). In addition, sea surface height standard deviation (SSH_sd), a measure of mesoscale variability, was a significant contributor to the models in summer/fall, but not in winter/spring (Figures 1 and 2). Multi-model ensembles outperformed single models (Table 1). Based on AUC and TSS metrics using the independent sightings dataset, the final model used for spatial predictions of habitat suitability was an ensemble between the seasonal BRTs and highest performing seasonal GAMMs. Point biserial correlation confirmed a significant association with higher habitat suitability values for sightings versus pseudo-absences (Pearson's correlation coefficient = 0.547, p-value <0.001; Elith et al., 2006).

The spatial dynamics of the whales' latitudinal migratory behaviour were evident in the predictions and matched patterns previously described in the literature (Bailey et al., 2009; Burtenshaw et al., 2004; Irvine et al., 2014; Mate et al., 1999). Blue whale habitat suitability in the CCE generally remained low January–April and began increasing in the SCB by May and June. Habitat suitability continued to increase northward through late summer and early fall, reflecting the whales' northward progression as more northerly habitats became productive in accordance with the seasonal upwelling cycle (Bograd et al., 2009). By November, habitat suitability contracted southward into the SCB and remained low through winter, in concordance with whales returning to tropical breeding grounds. Consistent with previous studies (Bailey et al., 2009; Hazen et al., 2017; Irvine et al., 2014), hotspots of habitat suitability within the CCE were observed in the SCB, Monterey Bay (~37°N), Gulf of the Farallones (~38°N), and in the vicinity of Cape Mendocino (~40°N) and Cape Blanco (~43°N) (Figure 3). Spatial predictions for 2016 during the spring and summertime upwelling season/period of intensive blue whale foraging in the California Current (May–September) are presented in Figure 3. Daily spatial predictions for the years 2009, 2015 and 2016 can be viewed interactively and downloaded via RShiny (S7).
3.2 | Seasonal timing of risk exposure in shipping lanes

Predicted whale habitat suitability in the shipping lanes varied substantially within and between years as a function of environmental conditions (Figures 4 and 5, Figure S8). Habitat suitability within the shipping lanes in 2009 matched the expected pattern of whale migratory behaviour as well as the observer sightings data, with very low habitat suitability December–April and a sustained period of high suitability June–October (Figure 5). In contrast, the same analysis during the Northeast Pacific marine heatwave of 2015 showed high periods of suitability distributed throughout the annual cycle, which were similarly concordant with the sightings data (Figure 5).

4 | DISCUSSION

Dynamic management approaches, in which management strategies are adjusted in concert with relevant biological, environmental and socioeconomic processes, are increasingly proposed to balance the trade-offs between human activities and species conservation (Lewison et al., 2015; Maxwell et al., 2015). Because dynamic management strategies often rely on an understanding of how the spatial distribution of a species or population changes with time (Howell, Kobayashi, Parker, Balazs, & Polovina, 2008; Maxwell et al., 2015), dynamic SDMs are emerging as an important natural resource management tool (Hazen et al., 2018). Dynamic SDMs can not only elucidate a species’ habitat preferences and distribution in relation to shifting environmental conditions, but they can also help identify the spatial and temporal dynamics of species’ risk exposure (Zydelis et al., 2011). Our study highlights the utility of dynamic ensemble modelling using high-resolution environmental data to identify time-varying species distributions and guide dynamic management of a highly migratory species.

4.1 | Model performance

Based on a suite of testing metrics, including validation against an extensive compilation of independent sightings data, our final seasonal model yielded accurate year-round predictions of daily blue whale habitat suitability in the CCE and realistically reproduced the whales’ expected latitudinal migratory behaviour. The top GAMM and BRT models showed strong descriptive and predictive performance, and an ensemble of the two models increased overall performance (Table 1). A previous blue whale model relating the same satellite tracking dataset to monthly remotely sensed variables obtained a full dataset AUC score of 0.86 (Hazen et al., 2017); the present model represents a substantial improvement in predictive performance based on that metric (Full AUC = 0.95), as well as compared to independent observations (Sightings AUC = 0.95). In addition, the use of year-round satellite tracking data spanning over ten years with daily ocean-modelled environmental variables enabled predictions of blue whale habitat suitability with unprecedented temporal coverage and resolution in the CCE, representing a significant step towards finer scale dynamic management applications.

Though our satellite tracking dataset ended in 2008, comparison with independent sightings data through 2017 suggested that the
FIGURE 3  Comparison of spatial predictions at 0.1-degree resolution of blue whale habitat suitability (range 0–1) in the California Current from the top-performing GAMM, BRT and ensemble models for the first day of the month in May, July and September 2016. Bottom right panel shows points of interest, from north to south: Cape Blanco, Cape Mendocino, Gulf of the Farallones, Monterey Bay (stars) and Southern California Bight (box)
extrapolative performance of our model into novel years and conditions is credible. Additional observations could be used to further evaluate extrapolative performance and monitor any changes in whale behaviour or environmental preferences as conditions in the CCE continue to change (Di Lorenzo, Miller, Schneider, & McWilliams, 2005; Di Lorenzo & Ohman, 2013). New data sources such as acoustic monitoring arrays (Širovic et al., 2015) could be integrated into both model building and validation. Blue whale locations collected from satellite tag data showed highest blue whale densities near the coast (Figure S1), which may be due in part to nearshore tagging locations (Bailey et al., 2009). Telemetry-based models may therefore underestimate offshore habitat suitability, though model performance remained high when compared to sightings data that were more broadly distributed offshore (Figure S3). Testing the performance of models developed using a combination of data types such as telemetry, transect survey and acoustic monitoring data would therefore be a valuable exercise for exploring biases based on data types used (Fithian, Elith, Hastie, & Keith, 2014; Yamamoto et al., 2015).

Our model evaluation procedure also highlights the value of using multi-model ensembles. Though top-performing GAMMs and BRTs yielded similarly high diagnostic scores (Table 1), fine-scale differences were evident in the spatial predictions (Figure 3). Different modelling approaches have various strengths and weaknesses, and in particular display trade-offs between the ability to explain fitted data versus predict novel data (i.e., descriptive versus predictive performance, respectively) (Derville et al., 2018). For instance, machine learning techniques like BRTs typically have strong descriptive power but can suffer from overfitting (Derville et al., 2018). Indeed, the unsmoothed response curves from our BRT models (Figures S5 and S6) show abrupt changes typical of the recursive binary splits of regression trees (Elith et al., 2008). Such abruptness in response curves can indicate overfitting, though the performance of these models tested against novel sightings data suggests this is not the case. In contrast to machine learning models, regression models like GAMMs may have lower descriptive performance but have been shown to have good predictive performance (Derville et al., 2018; Gregr et al., 2018; Qiao et al., 2015). GAMMs have also been proposed as effective tools for predicting into novel conditions (Becker et al., 2018; Derville et al., 2018). Ensemble models therefore provide an approach for balancing these trade-offs and can highlight areas of consensus between models (Araújo & New, 2007; Gritti et al., 2013; Marmion et al., 2009; Scales et al., 2015). In order to be relevant to management applications, our interest here was primarily in predictive rather than descriptive performance, and indeed, we demonstrate that a multi-model ensemble yielded higher predictive
showed strong spatial consistency with known krill hotspots in the CCE (Becker et al., 2016). Indeed, May–September predictions of blue whale habitat suitability were able to detect general patterns of whale habitat preferences. Furthermore, increased prey availability is likely contributing to the association of higher predicted blue whale densities in these areas (Becker et al., 2016; Hazen et al., 2017). Blue whales’ apparent preference for areas with shallower ILD values is also consistent with higher predicted blue whale densities in these areas (Becker et al., 2016), an association likely driven by increased prey availability. May–September predicted blue whale habitat suitability showed strong spatial consistency with known krill hotspots in the SCB, Monterey Bay, and downstream of Cape Mendocino and Cape Blanco (Santora, Sydeman, Schroeder, Wells, & Field, 2011; Santora, Zeno, Dorman, & Sydeman, 2018). Thus, while our model did not include prey data, it was able to realistically identify blue whale foraging hotspots via its combination of physical proxies.

4.3 | Implications for dynamic management

Despite the dynamism of human activities and species affected by them, most boundaries used for ocean management, such as shipping lanes, are static. If dynamic spatial boundaries are unfeasible for management, the spatiotemporal patterns of overlap with anthropogenic threats should be assessed to understand where and when greatest risk occurs (Redfern et al., 2013; Rockwood et al., 2017). Specifically, information on spatiotemporal patterns of risk could be used to guide the timing of management actions, such as the implementation of slowdown rules, where spatial management boundaries are otherwise inflexible. Previous year-round models for the Eastern North Pacific blue whale population are valuable for long-term planning (Hazen et al., 2017), but moving to finer spatiotemporal resolution offers a better match with the timescales of oceanographic variability and human activities in the region. We found that predicted whale habitat under average environmental conditions overlapped with southern California shipping lanes primarily during June–October, with greater risk exposure shifting from the western to the northern lane later in the year (Figures 4 and 5). However, the anomalous warming of 2015 (Bond et al., 2015; Di Lorenzo & Mantua, 2016; Jacox et al., 2018, 2016) resulted in dramatic changes in the timing of overlap, with risk exposure increasing earlier in the year under anomalously warm conditions (Figure 5). These predictions were mirrored by the date ranges of observed blue whale sightings in those years (Figure 5, Figure S4).

Such results suggest that environmental and climatic conditions should be considered in management planning and that slowdown rules or alternate shipping lanes could be employed dynamically based on time-varying risk of ship strike. Dynamic slowdown areas based on high whale use have been shown to effectively reduce North Atlantic right whale mortalities from ship strikes, and 15-day temporary management areas were employed at short notice to protect unpredictable whale aggregations (Laist et al., 2014). However, these measures were only effective when slowdowns were mandatory. As slow-steaming or use of alternative routes can be costly for vessels, dynamic implementation of these management options has the potential to be more economical than static or even seasonal rules. Daily distributions of whale habitat suitability could be combined with vessel movement data derived from shipboard monitoring systems (e.g., AIS; Kroodsma et al., 2018) to refine strike risk and evaluate alternative management options. Future studies should also investigate the correspondence in whale sightings or observed strikes to particular thresholds in whale habitat suitability that can be used to trigger management interventions.

Finally, our study relied on fine-scale, gapless environmental data derived from a data-assimilative ocean model developed for our study region. This type of model output is becoming more commonly available regionally and similar products exist globally at coarser

![Figure 5](https://example.com/figure5.png)
resolutions; we encourage development, dissemination and uptake of output from these ocean models for applications like the one demonstrated here. In the absence of gapless data, analytical techniques such as Boosted Regression Trees have been successfully applied to deal with missing remotely sensed data, for example due to cloud cover, in a dynamic species distribution modelling context (Hazén et al., 2018; Welch et al., 2018). With the increase in ocean modelling or remote sensing technologies and computational power, there is greater opportunity to implement dynamic management approaches that are more responsive to changing environmental conditions, species’ movements and patterns of human activity (Hazén et al., 2018; Maxwell et al., 2015). Such efforts thereby exemplify opportunities for cross-disciplinary collaboration. Dynamic, high-resolution species distribution models provide a valuable tool for assessing the spatio-temporal patterns of risk exposure to achieve management objectives.

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DATA AVAILABILITY

All spatial predictions from the habitat suitability model for 2009, 2015 and 2016 and shipping lane layers used in this study are available for download as raster grids via RShiny: https://heatherwelch.shinyapps.io/benioff_app/. Sample data and code for model fitting and evaluation are available via GitHub: https://github.com/briana-abrahms/DynamicEnsembleSDM.

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Biosketch

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Supporting information

Additional supporting information may be found online in the Supporting Information section at the end of the article.