Foraging distribution of breeding northern fulmars is predicted by commercial fisheries

J. H. Darby1,2,*, S. de Grissac3, G. E. Arneill1,2,4, E. Pirotta1,5, J. J. Waggitt6, L. Börger3, E. Shepard3, D. Cabot1, E. Owen7, M. Bolton8, E. W. J. Edwards9,10, P. M. Thompson9, J. L. Quinn1,#, M. Jessopp1,2,#

1School of Biological, Environmental and Earth Sciences, University College Cork, Cork T23 N73K, Ireland
2MaREI Centre for Energy, Climate and Marine, Environmental Research Institute, University College Cork, Cork P43 C573, Ireland
3Biosciences Department, Swansea University, Swansea SA2 8PP, UK
4Green Rebel Group, Crosshaven Boatyard, Crosshaven P43 EV21, Ireland
5Department of Mathematics and Statistics, Washington State University, Vancouver, WA 98686, USA
6School of Ocean Sciences, Bangor University, Isle of Anglesey LL59 5AB, UK
7RSPB Centre for Conservation Science, Elive House, Inverness IV2 3BW, UK
8RSPB Centre for Conservation Science, The Lodge, Sandy SG19 2DL, UK
9Lighthouse Field Station, School of Biological Sciences, University of Aberdeen, Cromarty IV11 8YL, UK
10Marine Scotland Science, 375 Victoria Rd, Aberdeen AB11 9DB, UK

ABSTRACT: Habitat-use and distribution models are essential tools of conservation biology. For wide-ranging species, such models may be challenged by the expanse, remoteness and variability of their habitat, these challenges often being compounded by the species’ mobility. In marine environments, direct observations and sampling are usually impractical over broad regions, and instead remotely sensed proxies of prey availability are often used to link species abundance or foraging behaviour to areas that are expected to provide food consistently. One source of food consumed by many marine top predators is fisheries waste, but habitat-use models rarely account for this interaction. We assessed the utility of commercial fishing effort as a covariate in foraging habitat models for northern fulmars Fulmarus glacialis, a species known to exploit fisheries waste, during their summer breeding season. First, we investigated the prevalence of fulmar−vessel interactions using concurrently tracked fulmars and fishing vessels. We infer that over half of our study individuals associate with fishing vessels while foraging, mostly with trawl-type vessels. We then used hidden Markov models to explain the spatio-temporal distribution of putative foraging behaviour as a function of a range of covariates. Persistent commercial fishing effort was a significant predictor of foraging behaviour, and was more important than commonly used environmental covariates retained in the model. This study demonstrates the effect of commercial fisheries on the foraging distribution and behaviour of a marine top predator, and supports the idea that, in some systems, incorporating human activities into distribution studies can improve model fit substantially.

KEY WORDS: Fisheries · Discards · Marine conservation · Foraging behaviour · Habitat use · Anthropogenic food source

1. INTRODUCTION

Understanding the drivers of species’ distributions is a key objective in conservation biology. Statistical modelling allows us to identify these drivers (Elith & Leathwick 2009) and to subsequently predict distribution patterns (e.g. Scales et al. 2016). Ideally, habitat use models incorporate environmental covariates that are known to reflect food availability—for example, vegetation type in terrestrial systems (Vynne...
et al. 2011) and primary productivity in marine environments (Nur et al. 2011). In terrestrial systems, these descriptors work well because the associated producers are commonly fixed in space and time (e.g. Fryxell et al. 2004, Smit 2011). In marine environments, equivalent descriptors are more elusive because prey patches are transient and less predictable (Fauchald 2009). For example, chlorophyll a (chl a) concentration and sea surface temperature (SST) are often used as a proxy for productivity and prey abundance (e.g. Tremblay et al. 2009, Domalik et al. 2018, Serratosa et al. 2020) but with low predictive power for higher predator behaviour or distribution (e.g. Kane et al. 2020). This is possibly due to spatio-temporal lags between environmental conditions that promote productivity and prey aggregations targeted by marine top predators (e.g. Whitehead et al. 2010, McGowan et al. 2013, Torres et al. 2015). It may also be that these variables predict prey biomass rather than availability (Boyd et al. 2015, Waggitt et al. 2018). As a result, static variables such as water depth, distance to colony and seabed slope are often found to better represent marine predator distribution (Amorim et al. 2009, Critchley et al. 2020). Such habitat descriptors may function in combination to enhance prey availability (e.g. Stevick et al. 2008, Scott et al. 2013). Although many studies have examined the predictive power of these oceanographic features in explaining marine distributions (Tremblay et al. 2009), the extent to which these effects are moderated by human activities has seldom been investigated.

Human activities have the potential to repel or attract species, for example through persistent habitat disturbance (Sauvajot et al. 1998) or provisioning of an extra food source (Newsome et al. 2015). In a marine setting, food sources derived from human activity usually originate from the fishing industry, either through depredation (e.g. Cosgrove et al. 2013) or through scavenging of offal and discards (Bicknell et al. 2013, Giménez et al. 2021). This association with human activity and waste can lead to negative effects, including plastic ingestion by urban scavengers (Caldwell et al. 2020), higher likelihood of human−wildlife conflicts (Cronin et al. 2016, Newsome & Van Eeden 2017) and the intake of food of lower nutritional value than natural prey (Grémillet et al. 2008).

Seabirds are widely distributed marine predators, but are experiencing global declines (Paleczny et al. 2015). Habitat-use and distribution models are essential tools in seabird conservation, as they can be used to identify areas of concern where seabirds and human activities co-occur (Critchley et al. 2018, Waggitt et al. 2020), and to prioritise areas for protection (Lascelles et al. 2012, McGowan et al. 2013). Many previous studies have focussed on the potentially harmful overlap between fisheries and seabirds in terms of bycatch risk (e.g. Tuck et al. 2011, Torres et al. 2013, Clay et al. 2019). However, at least 29 seabird species use fishing vessels as a source of food, through scavenging of discards or offal (Bicknell et al. 2013), or depredation of bait from baited gears (Dunn & Steel 2001), with studies showing how seabirds actively associate with vessels to forage on these products of fisheries (e.g. Bodey et al. 2014, Pirotta et al. 2018). Therefore, the distribution of commercial fishing effort might be a suitable predictor of foraging distribution for these marine predators. This could be through individuals co-occurring in the same productive areas as fishing vessels, but is more likely through vessel-attending species targeting areas of known vessel intensity in order to exploit fisheries waste (Collet & Weimerskirch 2020) or depredate catches.

The northern fulmar *Fulmarus glacialis*, hereafter ‘fulmar’, is a far-ranging, pelagic-foraging seabird that both benefits and suffers from fisheries interactions. It is one of the most commonly bycaught seabird species in the North Atlantic (Fangel et al. 2015, Hedd et al. 2016) and Alaskan fisheries (Dietrich et al. 2009). Fulmars are also vulnerable to other human activities and by-products, such as oil extraction (Fox et al. 2016) and plastic pollution (Acampora et al. 2013). Fulmars have undergone dramatic range expansion over the last 2 centuries, spreading from Iceland and St. Kilda through the UK, Ireland and on to the north coast of Europe (Burg et al. 2003). The drivers underlying this expansion likely include both the exploitation of fisheries waste (Fisher 1952, Phillips et al. 1999) and changing patterns of human exploitation (Gordon 1936, Thompson 2006). Currently, however, fulmars are in decline throughout much of their recently established range, observed both in colonies (Cordes et al. 2015, JNCC 2020) and at sea (Sherley et al. 2020). There is therefore an urgent need to better understand the drivers of their distribution when foraging at sea, and to investigate the causes of their recent declines.

We hypothesised that the foraging distribution of adult breeding fulmars could be better predicted using commercial fishing effort alongside a range of static and dynamic environmental variables. Using tracking data from fulmars and fishing vessels in Ireland and the UK, we estimated the prevalence of fulmar–fisheries interactions during the summer...
breeding season by looking at direct associations between the vessel and fulmar tracks. We then examined how fulmar foraging distribution was associated with commercial fishing effort over a broad spatio-temporal scale, comparing its importance as a habitat descriptor to other environmental variables. Finally, we incorporated fishing data and other significant environmental descriptors into models of fulmar foraging distribution over the range covered by our empirical data. The aim of this study was to infer that fulmar foraging distribution is influenced by commercial fishing effort, and that distribution models can be improved by acknowledging links between marine predators and human activity.

2. MATERIALS AND METHODS

2.1. Data collection

A total of 102 breeding adult fulmars were successfully tracked from Little Saltee, Ireland, the Isles of Scilly, England, and several Scottish mainland and island colonies between 2009 and 2019 (Table 1, Fig. 1). Fulmars were caught by hand, hand net or noose-pole from the nest. PathTrack Nanofix wireless enabled (10 g), Pathtrack Nanofix archival (15 g) or MobileAction iGotU gt-120 tags (~17 g) were attached to feathers on the centre of the bird’s back, directly above the centre of gravity, using Tesa® 4651 waterproof tape. All tag types record the same high accuracy (~3 m) GPS fixes. To mitigate potential negative impacts of tag attachment, total weight of the tag and attaching material were less than 3% of the total mass of the bird (2.19 ± 0.39% SD, max. 3.2%). Despite these considerations, we acknowledge that negative effects remain difficult to detect, control and quantify and could still lead to behavioural anomalies (Vandenabeele et al. 2014, Cleasby et al. 2021).

Tags were programmed to record locations at regular intervals. Depending on tag type and the intended duration of the deployment, this interval ranged from 1.5 to 10 min in the majority of birds, as well as a small subset of birds on Hirta, St. Kilda, with tags set to record every 15 or 20 min. Nests of tagged individuals were monitored, and data from tags were either remotely downloaded (PathTrack Nanofix wireless enabled) or retrieved after several days when the tagged bird had undertaken at least 1 foraging trip. All work was conducted under licences from the British Trust for Ornithology and the Irish National Parks and Wildlife Service. All data analyses were undertaken using R version 3.6.3 (R Core Team 2020). Foraging trips were originally defined as periods of >1 h that the bird spent >3 km from the colony. Distance from the colony and total distance travelled were calculated using the ‘raster’ package (https://cran.r-project.org/web/
packages/raster/; Tables S1 & S2 in the Supplement at www.int-res.com/articles/suppl/m679p181_supp.pdf). Data were further filtered to include foraging trips that contained >100 GPS fixes where birds travelled >10 km from the colony to exclude small trips that may have occurred because of disturbances at the colony. Points within 5 km of the colony were also removed as they likely encompassed colony-related behaviours (Bodey et al. 2014).

2.2. Fulmar–vessel foraging interactions

First, we aimed to establish the extent to which adult fulmars tracked in this study foraged in association with fishing vessels during the breeding season. Fishing vessel tracking data were sourced from the Vessel Monitoring System (VMS). Irish VMS data were sourced from the Irish Marine Institute, while UK VMS data were sourced from Marine Scotland Science. VMS provides GPS locations of fishing vessels >12 m long, approximately every 2 h. UK VMS data consisted of almost exclusively UK-flagged vessels, not detecting non-UK vessels in the UK Exclusive Economic Zone, likely leading to underestimates in fulmar–vessel encounters in these waters. Concurrent vessel and fulmar tracks were available from 2009 to 2018, and were linearly interpolated to the same resolution, either 3 or 10 min intervals depending on the resolution of the fulmar tracks from each colony. The coordinates of the nearest fishing vessel were extracted for each interpolated fulmar location in R, for a subset of complete tracks from 2018 and before (n = 92) for which sufficient VMS data were available. A frequentist implementation of the Pirotta et al. (2018) Bayesian approach, using similar constraints, was then used to fit a 7-state hidden Markov model (HMM) in ‘momentuHMM’ (https://cran.r-project.org/web/packages/momentuHMM/). This implementation was based on the fulmar example in the ‘momentuHMM’ vignette (McClintock & Michelot 2018) with the addition of a rest state. Briefly, our method assumes the occurrence of 7 states of movement: rest, transit (on outward journey, at fishing vessel or on return journey) and area-restricted search (ARS) (on outward journey, at fishing vessel or on return journey) (Table S3). These can be identified based on the step length and turning angle between interpolated points, while also incorporating distance and bearing to the nearest vessel and bearing to the colony. ARS is thought to correspond to foraging activity (Kareiva & Odell 1987, Weimerskirch et al. 2007, Bennison et al. 2018). Initial values of the model parameters were selected through \( k \)-means clustering of step lengths and turning angles into 3 clusters, intended to represent rest, ARS and transit states. The initial values of the parameters of the state-dependent distribution of distance to the nearest vessel \( d \) were selected based on the histogram of observed distances and prior knowledge that fulmars may direct their movement toward vessels from distances of up to 35 km (Pirotta et al. 2018; Table S4). The outputs of the 7-state HMM were used to estimate the proportion of time spent in vessel-associated ARS on a track-by-track basis, which was compared to the time spent in ARS away from detected vessels. Vessel gear type was not available for all VMS data, but where available, was used to investigate the frequency of occurrence of association with different types of fishing vessels. Gear types were grouped into 6 categories: trawlers, seines, longlines, gillnets, dredgers and traps/pots.

2.3. Identifying three primary states of fulmar behaviour

Next, we aimed to identify putative foraging behaviour using the features of fulmar tracks. All fulmar tracks were linearly interpolated to 10 min relocations, as this and subsequent analysis requires regular and uniform track point intervals. Trips were split into sections where gaps of >1 h were present in the raw GPS data to avoid interpolating over large time intervals. A separate 3-state HMM was fitted to these interpolated tracks to infer rest, ARS (putative foraging) and transit states (Figs. S1 & S2) irrespective of fishing vessel associations. In other words, while the 7-state HMM was used to differentiate specific vessel-associated ARS from other forms of ARS, this model was used to differentiate ARS from non-ARS more generally. Step lengths and turning angles between points were used to fit this HMM, with initial values of the parameters chosen using the same \( k \)-means procedure as for the 7-state model (Table S5). The ‘viterbi’ function in ‘MomentuHMM’ was used to obtain the most likely state sequence for each track. These inferred states were used to represent putative foraging and non-foraging behaviour, and to investigate foraging habitat preferences. One caveat of this approach is that it does not account for differences in search-type behaviour (ARS) during vessel-attendance or natural foraging, although the scales of movement are assumed to be similar (Pirotta et al. 2018).
2.4. Fulmar foraging habitat preference modelling

Fulmar foraging habitat preference was modelled using the outputs of the 3-state (rest, travel and ARS) HMM applied to the full tracking data. The response variable was presence or absence of ARS behaviour on each track point. To increase the accuracy in the absence data, only fixes classified as transit behaviour were retained to represent non-foraging, as rest can be difficult to discern from ARS without additional data, for example from time−depth recorders (Dean et al. 2013, Browning et al. 2018, Bennison et al. 2019). This response variable was modelled as a function of a set of fixed and dynamic covariates using binomial generalised additive mixed models (GAMMs) with a logit link function. Individual ID was included as a random effect. The ‘mgcv’ package was used for model fitting (https://cran.r-project.org/web/packages/mgcv/). The ‘bam’ function was used, because, while the autocorrelation estimation is more flexible in the ‘gamm’ function, the associated computation times were not feasible for this dataset. Skewed covariates were transformed towards normal using an optimised Box-Cox transformation factor (Box & Cox 1964; see Table S6), derived using the package ‘EnvStats’ (https://cran.r-project.org/web/packages/EnvStats/). Covariates highly correlated with one or more other covariates were identified and removed stepwise using the ‘concurvity’ function in ‘mgcv’. A high acceptable threshold of 0.8 was chosen for this process, as important partial effects may be expressed by related variables. This is especially true in biological settings with large sample sizes. The process by which ‘mgcv’ fits GAMMs also mitigates against the negative impacts of multicollinearity through backfitting of covariates (Wood 2008). The model was initially fit with a correlation parameter (\(\rho\)) of 0 (i.e. assuming no autocorrelation among residuals) grouped according to individual trip. The autocorrelation function (ACF) plot of the residuals of this model was then used to identify a suitable \(\rho\), the value of the correlation between consecutive residuals. The shape of the ACF plot was also used to verify that a first-order autoregressive (AR(1)) structure provided a good representation of the autocorrelation present. Thin-plate regression splines with shrinkage were used for all predictor variables, which return the simplest effective spline, with complexity further restricted by setting the gamma parameter to 1.2, which increases the null-space penalty when fitting the model (Wood 2003). This method avoids overfitting without having to arbitrarily constrain splines prior to model fitting. A whole-model approach using Akaike’s information criterion (AIC) was used for model selection using the ‘dredge’ function in the ‘MuMIn’ package (https://cran.r-project.org/web/packages/MuMIn/). Moran’s Index (MI) was calculated on the spatial distribution of residuals to investigate spatial autocorrelation in the model using the ‘ape’ package (https://cran.r-project.org/web/packages/ape/). A low MI value of 0.063 suggests that spatial autocorrelation was minimal, so no corrective spatial smooth was included in the model.

Physical habitat variables that may influence the availability of fulmar prey were chosen following Cox et al. (2018). These variables were depth, a seabed terrain ruggedness index (TRI; Wilson et al. 2007), stratification (Hunter-Simpson parameter; Simpson & Hunter 1974) and distance to the coast (km). These static environmental habitat predictors were handled as raster layers using the ‘raster’ package. Bathymetric data (TRI, depth) were sourced from a harmonised digital terrain model from the EMODnet database (https://portal.emodnet-bathymetry.eu). TRI identifies changes in seabed depth creating areas of complex currents, upwellings and internal waves, which are all known to enhance prey availability (Embling et al. 2012, Scott et al. 2013). Depth was included, as water column mixing is exaggerated in shallower waters, where tides interact more with the seabed (Cox et al. 2018). Stratification index was calculated as the Hunter-Simpson stratification parameter (Simpson & Hunter 1974) formulated from depth and maximum tidal current speed. Current information was provided at 1.5 km resolution, sourced from the Copernicus Analysis tool using the Atlantic Marginal Model (AMM15) available from the Marine Environmental Monitoring Service (Tonani et al. 2019). Maximum tidal current speeds were calculated from a 14 d spring–neap cycle. Stratification identifies mixed (<1.9), frontal (1.9) and stratified waters (>1.9), with fronts assumed to enhance prey availability (Scales et al. 2014, Waggitt et al. 2018). Prey availability may also be increased by intense mixing of the water column in areas of low stratification (Benjamins et al. 2015, Waggitt et al. 2016). TRI and stratification data were resampled to a 1 km resolution using a bilinear interpolation.

The Global Fishing Watch (GFW) Automatic Identification System (AIS) database was used to calculate commercial fishing effort across the region of interest at a 0.01 \(\times\) 0.01° spatial resolution. These AIS data differ from the VMS data used to run the 7-state HMM in that they provide the spatial distribution of commercial fishing effort with greater coverage,
while VMS returns the tracks of individual, identifiable fishing vessels. Fishing effort was calculated from hours fished only, so excluded vessels that were present but not thought to be engaged in fishing activity by the GFW process, which accurately differentiates fishing from non-fishing in >90% of cases (Kroodsma et al. 2018). We acknowledge that vessels may release offal/discards when in transit rather than when actively fishing, but this process sought to identify the broad spatial distribution of fisheries activity. The 7-state HMM inferred fulmars foraging in association with all major gear types (see Section 3). Another model was run with fisheries data split according to gear type, with model selection performed using the same process as for the generalised fishing effort model. The gear-type model was compared to the base model using AIC to assess the utility of gear-specific fisheries data in explaining fulmar foraging distribution. Fishing effort was observed to be spatially consistent across summer months (Tables S7 & S8, Figs. S3 & S4), and fishing hours from 2012 to 2018 were averaged across all of May to August (the breeding season for fulmars) in each grid cell to provide a representation of broad-scale fishing effort. The data were smoothed to 5 × 5 km rolling averages and assigned to track points using the ‘raster’ package. Point-specific monthly SST (°C) and chl $a$ (mg m$^{-3}$) values were sourced from NASA’s Ocean Biology Processing Group service (https://oceancolor.gsfc.nasa.gov/l3/), and appended to track points using Movebank’s bilinear interpolation and inverse distance-weighted processes, respectively (Dodge et al. 2013).

The goodness of fit (GOF) of the final model was assessed by measuring the area under the receiver operating characteristic curve (AUC). A confusion matrix and associated accuracy were also included using the packages ‘PresenceAbsence’ (https://cran.r-project.org/web/packages/PresenceAbsence/) and ‘caret’ (https://cran.r-project.org/web/packages/caret/), because AUC has limitations as a measure of GOF (Lobo et al. 2008). The effect of each covariate was plotted using the ‘mgcViz’ package (https://cran.r-project.org/web/packages/mgcViz/). The contribution of each covariate to model GOF was also estimated by removing the variable and calculating the change in AUC. The suitability of foraging habitat was then predicted over the range of the tracking data in each area by applying the model to a raster stack of static covariates. Chl $a$ and SST summer averages from 2002 to 2020 were used in this prediction and were again sourced from NASA’s OBPG (https://oceancolor.gsfc.nasa.gov/l3/).

3. RESULTS

We obtained tracking information from 102 breeding fulmars between 2009 and 2019. A total of 184 partial or full foraging trips were recorded from tracked individuals. No distinct differences in distance travelled or proportion of time in ARS were observed between different geographical areas (Tables S1 & S2). The distribution of tracks covered extensive areas of the southeast Celtic Sea, northwest North Sea and northeast Atlantic (Fig. 2)

3.1. Fishing vessel association and behaviour

Of the 102 tagged birds, 76 had complete foraging trips required for the 7-state HMM to infer direct vessel association. Of these 76 individuals, 41 were identified as engaging in vessel-associated ARS. Fig. 3 shows an example of a fulmar track inferred to have engaged in both natural and vessel-associated ARS over the course of 54 h in the Celtic Deep, a heavily fished area in the Celtic Sea. Within all trips analysed, a greater proportion of time was allocated to non-vessel-associated ARS (29.9%) than to vessel-associated ARS (9.5%) (Fig. 4). Within the trips in which vessel-associated ARS was detected, almost half of the time spent in ARS was spent in association with vessels (46.2 ± 30.1% SD). Most of the vessel-associated ARS occurred in the presence of trawlers (83.2%), consistent with the prevalence of trawler fisheries in the study area and the quantity of discards and offal they produce (Marine Institute & Bord...
Iascaigh Mhara 2011). Fulmars showed vessel interactions with all gear types, although longliners (<0.1%), trappers and potters (0.9%) and gillnetters (1.5%) only accounted for a small proportion of interactions (Fig. 5).

The outputs of the simplified HMM suggested that ARS and non-ARS states were roughly evenly distributed within trips, with 46% ARS (or putative foraging), 32% transit and 22% rest across the full tracking dataset.

### 3.2. Fulmar foraging habitat modelling

Commercial fishing effort was selected as a predictor of putative foraging distribution and behaviour, and environmental variables retained were stratification, seabed roughness, SST and chl a concentration (Table 2). Commercial fishing effort made a much greater contribution to model GOF than other variables retained (Table 2). The effect of each retained covariate is shown in Fig. 6. ARS was more likely in poorly stratified/well-mixed waters with a stratification index <3 and unlikely in highly stratified waters. The probability of engaging
in ARS increased with increasing commercial fishing effort, chl $a$ concentration, SST and seabed roughness. Model AUC was 74.6% and the prediction accuracy was 69%. These figures both suggest that the model has a moderate predictive power. A confusion matrix was constructed to compare predicted vs. actual values from the model outputs (Table 3).

When commercial fishing effort was split according to gear type, model fit improved according to AIC (11 198 with generalised fishing effort, 11 174 with fishing effort split by gear types). Trawler and gillnet fishing effort were retained by the model, with other fishing gears dropped by the model selection process (Table S8). Non-trawler fishing effort covariates were heavily positively skewed due to the reduced prevalence of these fisheries within the range of the tracked fulmars, leading to high power transformation factors (Box & Cox 1964) and limiting the utility of these covariates. Generalised fishing effort alone provided a better explanatory variable than retained gear-specific efforts (Table 2; Table S8), and because the tracked fulmars were shown to interact with all major fishing gear types, the model including generalised fishing effort was used for subsequent foraging habitat prediction.

Table 2. Retained habitat-preference generalised additive mixed model (GAMM) covariates associated with area-restricted search (ARS) behaviour of fulmars tracked from Irish, Scottish and Scilly Island colonies. Estimated degrees of freedom (EDF) represents the complexity of the model term, while $\chi^2$ represents the effect the term has on the model output. A p-value of <0.05 is considered significant. Change in the area under the receiver operating characteristic curve (AUC) of the model on removal of each variable is also included to represent variable effect on model goodness of fit. This table is ordered by $\chi^2$. See Fig. 6 for the marginal effects plots for each covariate.

| Variable            | EDF | $\chi^2$ | p     | Change in AUC (%) |
|---------------------|-----|----------|-------|-------------------|
| Fishing effort      | 1.5 | 199.4    | <0.001| 1.7               |
| Sea surface temp.   | 0.9 | 25.7     | <0.001| 0.1               |
| Stratification      | 0.9 | 23.3     | 0.001 | <0.1              |
| Chlorophyll $a$     | 0.8 | 9.1      | 0.015 | 0.1               |
| Seabed roughness    | 0.9 | 8.8      | 0.003 | 0.3               |

Table 3. Confusion matrix to assess the goodness of fit of the fulmar foraging habitat preference model by comparing model predictions to hidden Markov model (HMM)-estimated behaviours. Agreement between the habitat preference model prediction and HMM is seen as validation of that prediction’s accuracy, while disagreement is assumed to represent habitat preference model inaccuracy. The prediction accuracy of this model is 69%. ARS: area-restricted search

HMM-estimated transit | HMM-estimated ARS |
-----------------------|-------------------|
Predicted transit      | 5770              | 3751              |
Predicted ARS          | 2715              | 8641              |

Fig. 6. Marginal effects of each model covariate on the probability of area-restricted search (ARS) behaviour prediction in fulmars. Dotted lines and shaded grey represent 95% confidence intervals. The rugplots at the base of the graphs represent the distribution of values of each covariate in the model dataset. SST: sea surface temperature; TRI: terrain ruggedness index
The spatial predictions from the model for each study area are shown in Fig. 7. Large oceanographic features, such as the continental shelf edge, the Rockall Bank and the Norwegian Trench are highlighted as important foraging habitat. Intensely fished areas, such as the Celtic Deep, much of the continental shelf edge and the central North Sea, are also areas of abundant foraging opportunities for breeding fulmars. Closer to shore, areas with a high degree of mixing are identified as important, such as the southeast corner of Ireland (around Little Saltee) and the Pentland Firth between the Scottish mainland and Orkney.

4. DISCUSSION

Our study highlights high foraging effort in areas of persistently high fishing effort as well as a prevalence of vessel interactions in breeding fulmars. Of the tracked fulmars, 54% associated with vessels on foraging trips, and for those birds known to forage at vessels, roughly half of their time spent in ARS was in the presence of vessels. This provides important context for the elevated bycatch levels for this species (Dunn & Steel 2001, Fangel et al. 2015), as well as the range expansion of the last centuries (Fisher 1952). Not only does this finding improve our understanding of the distribution of fulmars and their use of anthropogenic food subsidies, it also provides insight into how the foraging distribution of this species may be changing with shifting fisheries distribution, policies and practices. On a broader scale, our study also suggests that anthropogenic factors should be considered when discussing the distribution and behaviour of species, such as the fulmar, that alter their habits in response to human activities.

The high degree of putative foraging in association with fishing vessels points to the potential importance of fisheries byproducts (discarded fish, offal, bait) in the diet of this species (Ojowski et al. 2001). This is likely exaggerated in breeding individuals, whose foraging range is central-place constrained by needing to return to the nest for incubation, chick-rearing and provisioning duties (Edwards et al. 2016). Dupuis et al. (2021) showed that fulmars interact with vessels throughout the non-breeding season, but that interactions were more prevalent in parts of their range closer to breeding colonies. This suggests that vessel attendance may be more common when central-place foraging compared to less restricted periods of their annual cycle. Fisheries waste and depredation opportunities may increase food availability within their restricted breeding range (Bicknell et al. 2013) and widen diet to include species otherwise physically out of reach (Hudson & Furness 1988, Thompson et al. 1995, Phillips et al. 1999) of this surface-feeding species. However, vessel interactions may also result in bycatch, and rates of vessel-associated foraging noted in this study are consistent with the high incidence of bycatch of fulmars relative to other seabird species (Dunn & Steel 2001, Fangel et al. 2015). Over half of the tracked fulmars associated with vessels, and of all ARS detected in the analysis, 30% was associated with fishing vessels. This is also likely to be an underestimate given that VMS data do not include vessels under 12 m length or those fishing illegally, as well as UK VMS data not capturing non-UK vessels (see Section 2.2). There may also be
a mismatch in the quantity of VMS data available across different time periods, as proportionally more vessels in the European fleet are fitted with VMS transponders over time (O’Shea & Thompson 2006), with similar patterns apparent in AIS data (Fig. S4).

The degree to which fulmars associated with vessels across our tracking dataset is also reflected in persistent fishing effort being a significant predictor of fulmar foraging habitat. According to model outputs, anthropogenic fishing effort was a better predictor of fulmar habitat preference than oceanographic covariates typically used in species distribution models, including chl a concentration, SST, seabed roughness and stratification (e.g. Camphuysen & Garthe 1997, Skov & Durinck 2001, Kane et al. 2020).

The recent decline of this species, as well as importance of intensely fished areas for foraging fulmars, highlights the need for targeted conservation that may involve adaptation of fishing gears or fishing practices to mitigate bycatch risk (Løkkeborg 2011, Domingo et al. 2017, Da Rocha et al. 2021). This study would also suggest that negative effects of other threats to fulmars such as oil pollution could be compounded if present in areas of intense fishing activity.

Despite fulmars being known consumers of fishery discards (Ojowski et al. 2001), the described effects of fishing vessels on fulmar foraging behaviour and distribution could arguably be due to both fulmars and fisheries targeting similar areas. While we cannot rule this out entirely, a recent study on similarly sized, generalist, surface-feeding procellariiform Cory’s shearwater Calonectris borealis (Granadeiro et al. 1998) found very little overlap with fishing activity within its range (Pereira et al. 2021). Cory’s shearwaters are not thought to habitually associate with fishing vessels, and low overlap would suggest their natural prey is obtained in areas that are not heavily fished. This supports the suggestion that fulmar foraging habitat being associated with fishing effort is due to fulmars concentrating ARS over areas of known intense fisheries, as Collet & Weimerskirch (2020) demonstrated in black-browed albatrosses Thalassarche melanophris, which were shown to direct foraging effort towards areas where they had previously encountered fishing vessels. Black-browed albatrosses are also known to take fishery discards (Mariano-Jelicich et al. 2014).

Fulmars still directed considerable foraging effort towards natural prey, with more foraging occurring in the absence of detected vessels. This is reflected in the contribution of environmental covariates to the foraging habitat preference model. Although natural prey species of fulmars are trophically distinct from primary producers (Furness & Todd 1984), chl a concentration was nevertheless identified as a significant predictor of fulmar foraging behaviour, consistent with studies in other procellariiforms (Kane et al. 2020) as well as other marine predators (Russell et al. 1999, Cox et al. 2016). Areas of low stratification, or well-mixed waters, may be of benefit to fulmars through the accumulation and aggregation of zooplankton by complex flow structures (Benjamins et al. 2015). Seabed roughness may also contribute to the complexity of flow structures and increase the availability of prey (Cox et al. 2018) as well as being unsuitable for many demersal fisheries due to risk of loss or damage to fishing gears through snagging on complex seabed terrain.

Fisheries are dynamic, and influenced by governance, market demand, policy and infrastructure (ports, harbours, etc.) as well as fish stock levels and distribution shifts due to climate change (Perry et al. 2005). The foraging distribution of fulmars and other discard-reliant species is therefore linked to the same range of socioeconomic and environmental factors. The European Union initiated a discard ban, or landing obligation, in 2015, calling on fishing vessels to cease the disposal of bycaught non-target species at sea (Borges 2021). Fulmars are known to take a range of fisheries byproducts, including offal and longline bait (Phillips et al. 1999), though our study has shown that fulmars have a clear affinity towards trawler vessels, the major proprietor of discards in the study region (Marine Institute & Bord láscaigh Mhara 2011). With discard rates presumably decreasing, it would be expected that fulmars may shift their fishing effort either towards natural prey, or towards other fishing gears with available byproducts. This may lead to increased associations with longline, gillnet and purse seine vessels, where fulmars can depredate bait or assimilated prey, although this in turn may cause increased rates of bycatch from fulmars getting caught in these gears (Dunn & Steel 2001).

Including fishing effort as a covariate in foraging habitat preference analysis may be useful for a range of species that take discards, target similar prey species to fisheries, or even actively avoid vessels. More generally, including anthropogenic factors in habitat models can improve our understanding of species distribution and behaviour across a range of taxa. For example, Russell et al. (2014) showed that windfarms can influence the foraging behaviour of seals by creating artificial reefs and excluding fisheries, both thought to lead to increased abundance of prey. Sim-
ilarly, Lieber et al. (2019) showed how a persistent artificial wake created by a wave power harvester created a foraging hotspot for surface-feeding seabirds through persistent mixing of the water column. Human industry is shaping the habits and habitats of species, and this study highlights the value of considering such anthropogenic factors when investigating species’ ecology.

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