# Frequency and consequences of individual dietary specialisation in a wide-ranging marine predator, the northern gannet

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ABSTRACT: Individual specialisations in animals are important contributors to a wide range of ecological and evolutionary processes, and have been particularly documented in relation to multiple aspects of foraging behaviours. Central-place foragers, such as seabirds, frequently exhibit pronounced specialisations and individual differences in a variety of foraging traits. In particular, the availability of fisheries discards alongside natural prey resources provides additional potential for differentiation and specialisation for opportunistically scavenging seabird species. However, the consequences of such specialisations for at-sea distributions and intraspecific interactions are not well known. Here, we investigated the links between the degree of dietary specialisation on natural or discarded prey and the foraging movements and spatial occupancy of northern gannets Morus bassanus in relation to differing intraspecific competition at 6 colonies of differing sizes. We found that, at most colonies, individuals with different dietary strategies concentrated foraging at differing levels of intraspecific competition. In addition, individuals pursuing different strategies were frequently, but not consistently, spatially separated, distinctions that were most acutely seen in females. However, this variation in individual strategy had no significant impact on current body condition. These analyses demonstrate how foraging-associated metrics need not covary within an unconstrained system. They also reveal that specialisation can have important consequences for the competitive regimes individuals experience, highlighting the complexity of examining interacting consequences at large spatial scales.

KEY WORDS: Fisheries  $\cdot$  Foraging  $\cdot$  GPS  $\cdot$  Individual specialisation  $\cdot$  Stable isotope analysis  $\cdot$  Seabird

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#### **INTRODUCTION**

Individual variation among animals is increasingly identified across a broad range of traits, and is key to understanding a range of ecological, evolutionary and applied issues (Van Valen 1965, Araújo et al. 2011, Wennersten & Forsman 2012). While examining variation at broader levels of classification, for example sex or age classes, can reveal relevant distinctions, significant variation is often left unexplained by such analyses, particularly in population-level generalists (Bearhop et al. 2004, Araújo et al. 2011). Individuallevel investigation can then be informative in explaining additional variation as, in many cases, organism responses, and their extent of specialisation across a range of attributes, differ among individuals (Bolnick et al. 2003, Araújo et al. 2011). Indeed, such variation among individuals frequently exceeds that within any one individual and, by spanning time or contexts, can result in long-term consistencies or behavioural syndromes (Dall et al. 2012, Sih et al. 2012).

Such specialisations have been shown to be theoretically and experimentally produced by increasing levels of competition (Svanbäck & Bolnick 2005, 2007, Bolnick et al. 2010), with inter-individual differences significant in reducing levels of competition among conspecifics (Durell 2000, Svanbäck & Bolnick 2007, Araújo et al. 2011, Tinker et al. 2012, Machovsky-Capuska et al. 2016a). Here, competitive interactions can affect individual prey choice preferences, producing differentiation between individuals within a single locality, and increasing the overall population niche width (Svanbäck & Bolnick 2007, Araújo et al. 2011, Ingram et al. 2011). These results typically rely on identifying whole food items, but there is also the potential for individuals to further specialise from a nutritional perspective through the selection of nutritionally complementary prey (Tait et al. 2014, Machovsky-Capuska et al. 2016a,b). Regardless of the level of selectivity, persistent differences in prey consumption among individuals can then lead to the establishment of dietary specialisations (Sih et al. 2012). Divergence in strategies can also occur through the movement of individuals displaced from areas of high competitive pressure. Such movements may lead to the discovery of different prey fields or foraging environments, with specialisations establishing among individuals in terms of their response to environmental cues and area occupancy across space or time (Bodey et al. 2014, Patrick et al. 2014, Wakefield et al. 2015). Thus, there are potential adaptive advantages to specialisation in many situations (Bolnick et al. 2011, Dall et al. 2012, Machovsky-Capuska et al. 2016b), but the links between competitive and environmental influences, and how these shape the consequences of specialisation, remain poorly understood.

Colonially breeding marine vertebrates (e.g. seabirds and pinnipeds) are excellent test subjects for hypotheses about the consequences of individual specialisations, particularly with respect to foraging behaviours such as travel and prey searching (Ceia & Ramos 2015). The constraints of colonial breeding produce intraspecific competition for prey among colony members (Lewis et al. 2001, Villegas-Amtmann et al. 2013), and the presence of neighbouring colonies can also constrain foraging opportunities (Wakefield et al. 2013, 2017). Such conditions can favour individual tactics that reduce competition with conspecifics, and this may be more keenly seen in larger colonies where higher densities of individuals can produce stronger competitive effects (Tinker et al. 2012, Ceia & Ramos 2015, Kernaléguen et al. 2015). The consequences of specialisation in such central-place foragers may thus be seen either through sympatric differentiation in measures including colony niche width (Araújo et al. 2011, Bolnick et al. 2011), or through changes in spatial distribution. These differences in occupancy can be generated through both deliberate choice and competitive exclusion. For example, juvenile red knots Calidris canutus are forced to forage for longer durations, and in more dangerous localities, through direct competitive interference by adults (van den Hout et al. 2014). Alternatively, different foraging specialisations, including maintaining a generalist strategy, can represent equally successful approaches for avoiding interference in what are, amongst marine predators, often scramble competition situations (Woo et al. 2008, Machovsky-Capuska et al. 2016a). Importantly, the extent to which an individual pursues any specialist or generalist strategy can have a broad range of consequences. This is clearly seen in the exploitation of new foraging opportunities such as fisheries discards. Despite being novel from an evolutionary perspective, a number of seabird species now routinely exploit such anthropogenic resources (Oro et al. 1996, Bartumeus et al. 2010, Wagner & Boersma 2011, Bicknell et al. 2013, Bodey et al. 2014, Patrick et al. 2015, Pirotta et al. 2018), and specialisation on discards can dramatically affect an individual's long-term fitness, either directly through changes in adult body condition or mortality, or indirectly through effects on timing of reproduction or chick survival (Grémillet et al. 2008, Bicknell et al. 2013).

Here, we examined the consequences of specialism in foraging strategies at multiple colonies of the northern gannet Morus bassanus (hereafter gannet). We combined information from GPS loggers with stable isotope analysis (SIA) of blood samples from individuals from 6 colonies spanning more than one order of magnitude in size (~2000 to ~60 000 pairs) in differing oceanographic environments. We hypothesised that (1) different dietary specialisations, in terms of specific prey species consumed, will explain variation in foraging movement metrics because different prey are likely to be associated with different environmental cues (Scales et al. 2014, Cleasby et al. 2015a, Wakefield et al. 2015), and (2) individuals pursuing different foraging strategies will be more divergent in space use at larger colonies as a result of the increased competitive pressures present (Lewis et al. 2001, Wakefield et al. 2013). We also explored the consequences of different foraging strategies for seasonal measures of individual fitness (body condition and breeding performance). Anthropogenic resources have been suggested to be nutritionally inferior to naturally foraged prey (Annett & Pierotti 1999, Grémillet et al. 2008, Votier et al. 2010, Tait et al. 2014, Machovsky-Capuska et al. 2016a). We therefore hypothesised that (3) individuals that incorporate high proportions of discards (anthropogenic resources) in their diets will have poorer body condition than those that specialise on naturally available prey.

#### MATERIALS AND METHODS

#### Field data collection

Gannets were captured, and then recaptured for device removal (i.e. a total of 2 captures ind. $^{-1}$ ), at 6 island colonies over 38 d from late June to early August 2011, ensuring overlapping of tracking and sampling at all colonies (see Table 1). Chick-rearing adults (chicks  $\geq 2$  wk post-hatching [range 2-7 wk], identified from Nelson 2001) were caught at the nest during parental changeover using a brass noose or crook on the end of a carbon fibre pole. Passive GPS loggers (30 g; i-gotu GT200e; MobileAction Technology) or GPS radio frequency loggers (45 g; e-obs) were deployed, depending on colony accessibility. All devices were attached to the base of the central tail feathers using Tesa<sup>©</sup> tape, as used in previous studies at many of these colonies with no negative effects (Votier et al. 2010, Cleasby et al. 2015a, Bodey et al. 2018), and acquired locations every 2 min. Birds with passive loggers were recaptured using the same methodology approximately 12 d later (mean time over which trips were recorded: 11.5 d, range 4-15 d; see Table 1) for device removal. A small blood sample (0.2–1.0 ml) was taken from the tarsal vein from most individuals at both capture and recapture, for sexing and SIA. Blood samples were kept in a cooler (1–7 h) until undergoing centrifugation to separate red blood cells (RBC) from plasma. Separated samples were then kept at -20°C until being dried and homogenised for analysis. Diet samples were also collected from all colonies through opportunistic collection of spontaneous regurgitates from both handled birds and other breeding individuals disturbed during the capture process. These were necessarily limited in number by our focus on capturing departing adults, i.e. those that had already fed and

brooded their chick, often for many hours, and typically had empty stomachs, and by our ethical decision to not unduly disturb other birds at each colony. Prey items were identified to the lowest possible taxon and then stored at -20°C until undergoing lipid extraction prior to isotopic analysis (see Text S1 in Supplement 1 at www.int-res.com/articles/suppl/ m604p251\_supp.pdf).

#### **Determination of dietary specialisations**

Dietary specialisations were identified using Bayesian stable isotope mixing models fitted in the SIAR package (Parnell et al. 2010) to assign proportions of different prey species in the diets of individuals. This involved analysing the isotopic ratios of  $\delta^{15}N$ and  $\delta^{13}$ C for RBC from initial capture of individuals to determine the proportions of different food sources consumed, reflecting diet over approximately the previous month (Hobson & Clark 1992). Data from 149 individuals, comprising birds where GPS devices were both successfully and unsuccessfully retrieved, were included, with lipid-extracted prey samples from the specific colony of the individual in question used as sources because colony foraging areas are largely discrete (Wakefield et al. 2013). Using these estimates of dietary components, individuals were then classified as specialists if they met 2 a priori criteria: (1) the modal prey item estimate for an individual must be >1 SD above the average of all birds sampled at that colony; and (2) the prey item in question must comprise >30% of the individual's total diet. These criteria together accounted for both variation in resource availability across colonies, and dietary importance in a species with a broad foraging capability (Nelson 2001, Hamer et al. 2007), although they do not consider variation in the nutritional composition of prey that may add additional subtlety (Machovsky-Capuska et al. 2016b). Specialists were further categorised as either forage fish specialists (e.g. consumers of mackerel Scomber scombrus) or those that specialised on demersal discards (whitefish; see Supplement 1). Individuals with diets that did not meet these criteria were classed as generalists.

#### GPS data processing and movement metrics

Only complete foraging trips were included in analyses of foraging behaviour. In addition, all locations within 1 km of the colony were deleted as individuals only use these areas for bathing and rafting (Carter et al. 2016). Three metrics assessing different components of foraging behaviour were calculated from each individual trip: (1) total trip length (km), reflecting effort expended; (2) angle of departure (the average over the first 5 bearings >1 km from the colony, degrees), reflecting the extent to which an individual uses past knowledge; and (3) maximum distance from the colony (km), combining energy expended with both personal and public information use. In addition, for each GPS location  $L_0$ , speed (between  $L_{-1}$  and  $L_0$ ) and tortuosity (the degree to which the tracked animal's path diverges from a straight line between  $L_{-4}$  and  $L_{0}$ ) were determined. Putative foraging locations were then identified based on these parameters as described in Wakefield et al. (2013). Colony-specific utilisation distributions (UDs) were then estimated to enable investigation of the levels of intraspecific competition likely to be experienced by gannets foraging in different locations (see 'Habitat selection' below). The colony mean kernel density (KD) for all putative foraging locations was calculated based on a 2 km Lambert azimuthal equal-area grid using the R package 'adehabitatHR' (Calenge 2007). Individuals were tracked for different lengths of time, so the KD was estimated for each individual with the smoothing parameter *h* estimated by least-squares cross-validation. The mean smoothing parameter,  $\overline{h}$  was then used to estimate the KD for each individual, and this was averaged across individuals within colonies. UDs were then calculated for the 95, 75, 50 and 25% levels at each colony.

#### **Consequences of foraging strategies**

## Links between dietary specialisations and movement metrics

We used 3 generalised linear mixed models (GLMMs), one for each movement metric, to examine whether the identified foraging strategies significantly influenced foraging metrics. These models included sex and colony as fixed effects known to influence foraging behaviours (Stauss et al. 2012, Cleasby et al. 2015a) as well as a random individual effect. We examined whether there was an additional effect of dietary specialisation (n = 88 ind. spanning all colonies with full data required). Models were compared using an information theoretical approach, with the model with the lowest Akaike's information criterion for small samples (AIC<sub>c</sub>) score regarded as the top model. However, in instances where the top model included an extra term that did not improve

the model  $AIC_c$  score by more than 2 units, the most parsimonious model was also highlighted, as such additional terms can be regarded as uninformative (Arnold 2010). Goodness-of-fit was assessed using the likelihood-ratio based pseudo- $R^2$  (Nakagawa & Schielzeth 2013).

#### Habitat selection

We used habitat selection functions (HSFs) to model the foraging range usage by birds within each of the 3 dietary categories as a function of the level of competition experienced. HSFs compare spatial locations that are used vs. unused-but-available, adopting a logistic regression based approach with a casecontrol design (Aarts et al. 2008). This generates a binomial response that takes the value 1 for the *i*<sup>th</sup> data point if it belongs to the data set of putative foraging locations, or the value 0 if it belongs to the control data set. The control data set consisted of 5 pseudo-absences selected randomly within the 95% UD of each colony matched to each observed foraging location.

To estimate the level of competition experienced by gannets when foraging, we calculated the density of individuals at each point as as  $\hat{u}_{i,x}N_{i'}$  where  $\hat{u}_{i,x}$  is the estimated absolute density of use of cell x (cell size = 4 km<sup>2</sup>) by birds from colony  $i_i$  and  $N_i$  is the number of breeding pairs at the  $i^{\text{th}}$  colony (Wakefield et al. 2013). This approach incorporates information on colony size and allows for adjustment for how bird density declines within a colony's foraging range with increasing distance from the colony, and UDs thus calculated match data from at-sea surveys (Cleasby et al. 2015b). In addition, while we were not able to include data on prey availability, as fish distributions are not measured synoptically over the scale with which we tracked gannets, individuals from several of these study colonies are known to repeatedly cue in on stable oceanographic features (Scales et al. 2014, Cleasby et al. 2015a). As the foraging ranges of some colonies partially overlapped, we summed the spatial density estimates across grid squares at these locations (see Fig. S1 in Supplement 1). While small untracked colonies from which overlaps could not be calculated were located within the study area, these colonies represent <5% of the total birds foraging across the entire area, so additional competitive interactions will be minimal.

HSFs were estimated using a binomial generalised additive mixed model (GAMM) in the R package 'mgcv' (Wood 2006). The response variable was whether a location was used (1) or not (0), with the level of competition at each location included as a smoother. In our full model, we estimated separate competition smoothers for each foraging specialisation category by colony combination (e.g. 'bass rock – forage fish' or 'grassholm – generalist'). Bird identity nested within colony identity were included as random intercepts, and a thin-plate regression spline for the spatial coordinates of each data point was included to account for spatial auto-correlation (see Supplement 1). From this initial model, minimum adequate models were selected by backwards selection using *K*-fold cross-validation (K = 5; Supplement 1), using the summed log-likelihood values for the holdout data as a goodness-of-fit measure.

#### Body condition

Body condition was measured in the field as a seasonal fitness proxy, as offspring recruitment rates and lifetime individual breeding success are not known in this system. This was estimated using the scaled mass conditional index (Peig & Green 2009). Body mass was measured  $(\pm 50 \text{ g})$  on initial capture when the stomach was empty, and scaled to the mean maximum tarsus length (see Supplement 1). This index was calculated using data from 176 individuals across all colonies. It is hypothesised that a higher scaled mass is an indicator of individuals with higher fitness because breeding is a demanding process which is likely to reduce body condition. The effect of specialisation on scaled mass was assessed using a general linear model (GLM) with a Gaussian error structure, and the full model included all 2-way interactions between colony, sex and dietary type. Simplified models were compared using AIC<sub>c</sub> scores, with consideration of both the top ranked and the most parsimonious models. Normal Q-Q plots confirmed that all model residuals conformed to assumptions of normality, and all analyses were conducted in R v.3.4.3 (R Core Team 2017).

## RESULTS

A total of 112 individuals were successfully tracked across the 6 colonies (mean  $\pm$  SD ind. colony<sup>-1</sup>: 19  $\pm$  8), producing 810 complete foraging tracks (range ind.<sup>-1</sup>: 2–20; Table 1). Blood samples were taken from 149 individuals (mean  $\pm$  SD ind. colony<sup>-1</sup>: 25  $\pm$  11, including 98 successfully tracked individuals). The majority of individuals were categorised as general-

except for the Grassholm situation	e, which was surveyed i	in 2009 (http	p://jncc.defra.gov.uk/smp/ stable isotope aı	). Retrieved device nalysis	s are those	e from whi	ch data were s	successfully re-	covered. SIA:
Colony name	Geographic location	Colony size (AON)	Devices retrieved with multiple complete trips (deployment dates)	Median number of trips ind. <sup>-1</sup>	Individ sampled 1 Males	luals for SIA Females	Generalists	Forage fish specialists	Discard specialists
Great Saltee, Ireland Bull Rock, Ireland Ailsa Craig, Scotland, UK Little Skellig, Ireland Grassholm, Wales, UK Bass Rock, Scotland, UK	52° 06' N, 06° 37' W 51° 35' N, 10° 18' W 55° 15' N, 05° 06' W 51° 46' N, 10° 30' W 51° 43' N, 05° 28' W 56° 05' N, 02° 24' W	2400 3700 27100 29700 39300 55500	18 (2–19 Jul) 14 (28 Jun–15 Jul) 16 (7–22 Jul) 9 (11–23 Jul) 30 (25 Jun–29 Jul) 25 (26 Jun–2 Aug)	9 12.5 6	13 5 5 14	20 11 5 9	26 (79%) 15 (56%) 11 (69%) 6 (60%) 22 (55%) 19 (82%)	$\begin{array}{c} 4 \ (12 \%) \\ 6 \ (22 \%) \\ 2 \ (13 \%) \\ 3 \ (30 \%) \\ 8 \ (20 \ \%) \\ 4 \ (18 \%) \end{array}$	$\begin{array}{c} 3 \ (9\%) \\ 6 \ (22\%) \\ 3 \ (19\%) \\ 1 \ (10\%) \\ 10 \ (25\%) \\ 0 \ (0\%) \end{array}$

Table 1. Fieldwork locations, sample sizes and foraging strategy categorisations of northern gannets. Colony sizes are apparently occupied nests (AON) counted in 2004

ists, with the proportion of specialists of either kind varying substantially between colonies (Table 1).

# Links between dietary specialisations and movement metrics

The top models for all movement metrics contained the effects of sex and colony, confirming the known increase in foraging distances at larger colonies (Lewis et al. 2001), and reflecting the fact that females typically travel greater distances than males (Cleasby et al. 2015a) (Fig. 1, Table 2). Dietary specialisation had an important effect only on the maximum distance birds moved from their colony (Fig. 1, Table 2 & Table S1 in Supplement 1). Females tended to travel further than males in all categories, but this was most pronounced in forage fish specialists. Conversely, female discard specialists travelled substantially smaller maximum distances from the colony than other females. Males changed little in maximum displacement distance regardless of dietary type.



Fig. 1. Maximum distance travelled from the colony by northern gannets on foraging trips (averaged across all colonies) depends on gannet dietary type and sex (females [F] = grey bars, males [M] = white bars; boxes represent interquartile range and median; dots represent outliers). Number of individuals within each dietary type: generalist: F = 24, M = 30; forage fish specialist: F = 12, M = 3; discard specialist: F = 6, M = 12

#### **Consequences of foraging strategies**

#### Habitat selection

Based on *K*-fold cross-validation, the best predictive HSF was one that incorporated separate competition smoothers for each foraging specialisation category on a colony-by-colony basis (Tables S2 & S3 in Supplement 1). This indicates that the relationship between foraging specialisation and the density of conspecifics encountered at sea varied both among strategies and colonies, despite the fact that, within a colony, similar total ranges of competition were experienced (Fig. 2). This result was also reflected spatially, with individuals pursuing different foraging strategies often diverging in geographical locations visited (Fig. 2).

At the 2 largest colonies at which discard use was recorded (Ailsa Craig and Grassholm; Fig. 2), discard specialists showed greater usage of foraging areas with higher levels of competition, with usage rapidly reducing in areas of lower competitive pressure. In contrast, forage fish specialists showed a reversal of this trend. While central-place foraging necessarily means they experience the highest levels of competition, peak predicted usage rose above that of other dietary types at lower levels of competition, indicating that forage fish specialists spent more foraging effort in areas with low conspecific densities. Generalist foragers showed a similar pattern to discard specialists, but with a weaker selective response to areas of high competition. Similar results were also observed at the largest colony (Bass Rock; Fig. 2) where generalist foragers were predicted to make greater use of areas with higher conspecific competition than forage fish specialists, with usage reversed at the lowest levels of competition (no discard regurgitates were identified here in 2011). However, such differentiation between strategies was not apparent on the west coast of Ireland, where colonies showed little spatial differentiation and strategies followed similar trajectories across the competition gradient. Lastly, at the smallest colony (Great Saltee; Fig. 2) neither specialist type extensively foraged under the higher levels of competition experienced by generalists.

#### Body condition

Females were significantly heavier than males at most colonies, but dietary type had no impact on scaled mass (Figs. S2 & S3, Table 3).

Table 2. Comparison of mixed models examining the effect of dietary specialisation on northern gannet foraging movement metrics. The top model determined by Akaike's information criterion (AIC) ranking for each metric is presented in **bold**, and the most parsimonious model is marked with \* (see 'Materials and methods' for more details). The variance explained by the top model for each metric (and the most parsimonious where relevant) is also presented

Model	ΔAIC compared to top model for each foraging met		
	Trip distance	Max. distance	Departure angle
	(km)	from colony (km)	(°)
Sex + colony	0.00*	5.17	24.00
Sex + dietary type	33.62	38.73	72.08
Colony + dietary type	4.61	7.01	16.39
Sex + colony + dietary type	0.86	5.21	15.28
$Sex + colony + sex \times colony$	1.56	3.31	0.36*
$Sex + colony + dietary type + sex \times dietary type$	0.38	0.72*	17.00
$Sex + colony + dietary type + sex \times colony$	2.25	4.39	0.00
$Sex + colony + dietary type + sex \times colony + sex \times dietary type$	1.70	0.00	3.65
Null	35.89	39.10	86.84
Goodness-of-fit <b>top</b>	0.199	0.295	0.475
Goodness-of-fit most parsimonious*	-	0.284	0.471

## DISCUSSION

Our results demonstrate how, in an unconstrained system across multiple populations and environmental conditions at large spatial scales, variation in dietary strategy can have consequences for spatial separation in, and the competitive environments experienced by, an apex predator. We demonstrated that individuals specialising on forage fish showed greater usage of areas of reduced competitive pressure (i.e. lower densities of conspecifics) compared to either discard specialists or generalists (Fig. 2). However, dietary specialisations were also present in some instances without broad spatial separation in foraging locations, highlighting the degree to which environmental variation is important in facilitating the realisation of specialisations. We also showed that there is significant variation in foraging movements between females, but not males, pursuing different strategies (Fig. 1). However, these individual differences had limited consequences for our measured fitness correlate (body condition), suggesting that different strategies may represent alternative successful solutions to cope with interspecific competitive effects in this species.

When considering links between dietary specialisations and foraging movements, we only found support for differences in maximum displacement from the colony. Females tended to travel farther than males within all strategies (Fig. 1) and, within females, forage fish specialists reached significantly more distant points than discard specialists. This movement metric reflects a degree of both the effort involved and the use of both public and private knowledge, and suggests that individuals pursuing all strategies have favoured search localities or environmental triggers that they will repeatedly target (Dall et al. 2012, Masello et al. 2013, Patrick et al. 2014, Wakefield et al. 2015). However, we found no significant relationships between the pursuit of different dietary strategies and either trip length or departure angle. This in turn suggests that diverse localities and patch types were available within all colonies' foraging ranges, and that, for individuals pursuing all strategies, time to locate food patches varied between trips in this dynamic environment (Scales et al. 2014, Wakefield et al. 2015). This lack of commonality between the extent of specialisation in prey selection and in multiple foraging movements suggests that these 2 components may not form a behavioural syndrome in this species (Sih et al. 2012). Behaviours may simply be linked across time periods (Wakefield et al. 2015), or certain foraging techniques and locations may be best suited to certain individual phenotypes (Lewis et al. 2002, Dall et al. 2012).

However, we did find that birds exhibiting different dietary strategies (generalists, forage fish or discard specialists) frequently experienced different competitive regimes while foraging (Fig. 2), and while sample sizes at any one colony could be relatively small, this pattern was repeated at several of our study colonies. This suggests that an interaction between foraging preference and the degree of competition experienced at a location may well affect the foraging decisions of individuals and thus explain repeatable displacement distances from the colony (Corman et al. 2016). Forage fish specialists, particu-





Table 3. Comparison of general linear models examining the effect of dietary specialisation on scaled mass of adult gannets. The top model determined by Akaike's information criterion (AIC) ranking is presented in **bold**, and the most parsimonious model is marked with \* (see 'Materials and methods' for more details). The variance explained by the top model is also presented

Model	ΔAIC compared to top model (scaled mass)
Colony + sex	0.00*
$Colony + sex + colony \times sex$	4.14
Colony + sex + dietary type	2.72
$Colony + sex + dietary type + colony \times sex$	6.64
Colony + sex + dietary type + sex × dietary type	4.04
Sex	5.49
Dietary type	11.17
Sex + dietary type	7.18
Colony	5.85
Null	10.15
Goodness-of-fit for top model	0.143

larly females, tended to fly further (Figs. 1 & 2), and Bartumeus et al. (2010) demonstrated that such foraging on natural prey tends to create a super-diffusive movement process characterised by longer flights. This suggests an alternative strategy that may be employed by females in particular as a result of competitive exclusion by more aggressive males at discarding opportunities (Nelson 2001, Lewis et al. 2002, Stauss et al. 2012). Alternatively, it may reflect certainty of parentage and a willingness to 'work harder' at chick provisioning (Kokko & Jennions 2008), or differences in nutritional demands, particularly post-egg production, between the sexes (Machovsky-Capuska et al. 2016a, Botha & Pistorius 2018). Contrastingly, we found that discard specialists traveled shorter distances and experienced higher competition, supporting a sub-diffusive movement pattern for discard specialists (Bartumeus et al. 2010) (Fig. 2). Becoming a discard specialist has been suggested to provide large volumes of food with reduced flying (and therefore energetic) costs for adults, although with additional costs in terms of nutritional quality (Grémillet et al. 2008, van Donk et al. 2017). However, remaining closer to the colony will naturally lead to individuals foraging in areas where greater numbers of conspecifics are present. Our results suggest that any energetic benefits of exploiting discards through reduced commuting costs may be offset by greater conspecific competitive pressures and the potential for conflicts this can produce at a spatially concentrated resource. This potential cost-benefit scenario for the exploitation of

discards should be explored further with respect to its potential to affect population growth at individual colonies.

Interestingly, while most apparent at larger colonies, clearer spatial separation of different strategies was not consistently achieved with increasing colony size, although such spatial divergence between different strategies has been demonstrated theoretically and on smaller mesocosm scales (Svanbäck & Bolnick 2005, 2007, Bolnick et al. 2010). For example, dietary specialisation was achieved by some individuals at the most western colonies (Bull Rock and Little Skellig) despite almost complete overlap in foraging space and competitive environments experienced (Fig. 2), and a substantial difference in these colony sizes. Breeding gannets are almost exclusively foragers in neritic waters (Nelson 2001), and the closer proximity of the shelf break to these colonies compresses both natural and anthro-

pogenic foraging opportunities into a smaller area, such that variation in ecological opportunities may be maintained despite spatial restrictions. Contrastingly, at the smallest colony (Great Saltee), there was clear spatial separation between forage fish specialists and discard specialists, likely reflecting the high levels of discards available in the southern Irish Sea (Anonymous 2011). These results highlight alternative ways in which ecological opportunities can facilitate the maintenance of dietary specialisations, and emphasise the necessity of considering the interactions between intraspecific competition and ecological opportunity in order to understand when and how individuals are able to achieve foraging differentiation (Roughgarden 1974, Parent & Crespi 2009, Araújo et al. 2011).

Although proportions were neither consistent across colonies, nor scaled with colony size, we found far more individuals followed generalist than specialist strategies amongst those sampled. While gannets are capable of taking a wider range of prey than many other sympatric seabirds (Nelson 2001), and thus may seem to have a greater potential for developing individual specialisations, their foraging opportunities are often constrained by conspecific interference competition (Garthe & Huppop 1998, Lewis et al. 2001, Votier et al. 2013). When combined with inter-annual changes in prey availability and environmental parameters (Hamer et al. 2007), this may preclude high degrees of specialisation and ensure individuals are able to respond to changeable conditions (Hamer et al. 2007, Dall et al. 2012) while

meeting their nutritional requirements (Machovsky-Capuska et al. 2016a). This potential for flexibility may also explain why, despite variation in habitat usage and distances covered in response to competitive and environmental pressures, different foraging strategies did not affect adult scaled mass. Although specialisation on forage fish and discards has previously been linked to better and poorer body condition respectively at one of these colonies (Grassholm; Votier et al. 2010), a similar result was not found when examining the relationship across multiple colonies (with the exception of Great Saltee; see Fig. S3 in Supplement 1). However, as outlined above, this relationship may vary across years as a consequence of changes in prey field availability and nutritional composition (Hamer et al. 2001, Scales et al. 2014, Tait et al. 2014, Wakefield et al. 2015, Machovsky-Capuska et al. 2016a), and may also be affected by sample sizes. The only other clear distinction was that, at the largest colonies, the scaled mass of individuals tended to be lower. This could be due to competition-driven increases in foraging range impacting on body condition (Lewis et al. 2001), or it may be a strategic decision to reduce wing loading to facilitate longer flights. Whether this has any important effects on longevity or reproductive output remains unknown, particularly as differences may become apparent only under especially unfavourable conditions or when individuals are followed over many years (Annett & Pierotti 1999, Hamer et al. 2007, Lescroel et al. 2010). This is especially likely as long-lived adults maintain a wide safety margin in body mass, prioritising self-maintenance over current provisioning, potentially requiring much longerterm individual based studies to determine fitness effects (Lecomte et al. 2010).

Our findings demonstrate that dietary specialisations can have important consequences for the competitive regimes that individual gannets experience and, at several colonies, although sample sizes were relatively small, this can result in spatial separation of individuals of specialist and generalist foraging strategies. This pattern was seen at both small and large colonies that were located away from shelf breaks, suggesting that intraspecific competitive effects are not the sole contributor to these patterns. For example, interspecific effects may mirror intraspecific interactions at multi-species aggregations, leading to disruption of feeding opportunities, with such interspecific competitive regimes often important in affecting species foraging distributions (Ballance et al. 1997, Ronconi & Burger 2011, Dhondt 2012). The interactions between foraging specialisations and competition are nuanced, and the consequences found here highlight the complexity of examining interacting consequences at large spatial scales.

*Data archive*. Tracking data are available at http://seabird tracking.org. Data for the models presented in Table 2 are included in Supplement 2 at www.int-res.com/articles/suppl/m604p251\_supp2.xlsx.

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#### LITERATURE CITED

- Aarts G, MacKenzie M, McConnell B, Fedak M, Matthiopoulos J (2008) Estimating space-use and habitat preference from wildlife telemetry data. Ecography 31:140–160
- Annett CA, Pierotti R (1999) Long-term reproductive output in western gulls: consequences of alternate tactics in diet choice. Ecology 80:288–297
- Anonymous (2011) Atlas of demersal discarding, scientific observations and potential solutions. Marine Institute, Bord Iascaigh Mhara, Dún Laoghaire. https://oar.marine. ie/handle/10793/666
- Araújo MS, Bolnick DI, Layman CA (2011) The ecological causes of individual specialisation. Ecol Lett 14:948–958
- Arnold TW (2010) Uninformative parameters and model selection using Akaike's information criterion. J Wildl Manag 74:1175–1178
- Ballance LT, Pitman RL, Reilly SB (1997) Seabird community structure along a productivity gradient: importance of competition and energetic constraint. Ecology 78: 1502–1518
- Bartumeus F, Giuggioli L, Louzao M, Bretagnolle V, Oro D, Levin SA (2010) Fishery discards impact on seabird movement patterns at regional scales. Curr Biol 20: 215–222
- Bearhop S, Adams CE, Waldron S, Fuller RA, Macleod H (2004) Determining trophic niche width: a novel approach using stable isotope analysis. J Anim Ecol 73: 1007–1012
- Bicknell AWJ, Oro D, Camphuysen KCJ, Votier SC (2013) Potential consequences of discard reform for seabird communities. J Appl Ecol 50:649–658
- Bodey TW, Jessopp MJ, Votier SC, Gerritsen HD and others (2014) Seabird movement reveals the ecological footprint of fishing vessels. Curr Biol 24:R514–R515
  - Bodey TW, Cleasby IR, Bell F, Parr N, Schultz A, Votier SC, Bearhop S (2018) A phylogenetically controlled metaanalysis of biologging device effects on birds: deleterious effects and a call for more standardized reporting of study data. Meth Ecol Evol 9:946–955

- Bolnick DI, Svänback R, Fordyce JA, Yang LH, Davis JM, Hulsey CD, Forister ML (2003) The ecology of individuals: incidence and implications of individual specialization. Am Nat 161:1–28
- Bolnick DI, Ingram T, Stutz WE, Snowberg LK, Lau OL, Paull JS (2010) Ecological release from interspecific competition leads to decoupled changes in population and individual niche width. Proc R Soc B 277:1789–1797
- Bolnick DI, Amarasekare P, Araújo MS, Bürger R and others (2011) Why intraspecific trait variation matters in community ecology. Trends Ecol Evol 26:183–192
- Botha JA, Pistorius PA (2018) Variability in the foraging distribution and diet of cape gannets between the guard and post-guard phases of the breeding cycle. Front Mar Sci 5:15
- Calenge C (2007) Exploring habitat selection by wildlife with adehabitat. J Stat Softw 22:1–19
- Carter MID, Cox SL, Scales KL, Bicknell AWJ and others (2016) GPS tracking reveals rafting behaviour of northern gannets (*Morus bassanus*): implications for foraging ecology and conservation. Bird Study 63:83–95
- Ceia FR, Ramos JA (2015) Individual specialization in the foraging and feeding strategies of seabirds: a review. Mar Biol 162:1923–1938
- Cleasby IR, Wakefield ED, Bodey TW, Davies RD and others (2015a) Sexual segregation in a wide-ranging marine predator is a consequence of habitat selection. Mar Ecol Prog Ser 518:1–12
- Cleasby IR, Wakefield ED, Bearhop S, Bodey TW, Votier SC, Hamer KC (2015b) Three-dimensional tracking of a wide-ranging marine predator: flight heights and vulnerability to offshore wind farms. J Appl Ecol 52:1474–1482
- Corman AM, Mendel B, Voigt CC, Garthe S (2016) Varying foraging patterns in response to competition? A multicolony approach in a generalist seabird. Ecol Evol 6: 974–986
- Dall SRX, Bell AM, Bolnick DI, Ratnieks FLW (2012) An evolutionary ecology of individual differences. Ecol Lett 15: 1189–1198
- Dhondt AA (2012) Interspecific competition in birds. Oxford University Press, Oxford
- Durell SEA Le V dit (2000) Individual feeding specialisation in shorebirds: population consequences and conservation implications. Biol Rev Camb Philos Soc 75:503–518
- Garthe S, Hüppop O (1998) Foraging success, kleptoparasitism and feeding techniques in scavenging seabirds: Does crime pay? Helgol Meeresunters 52:187–196
- Grémillet D, Pichegru L, Kuntz G, Woakes AG, Wilkinson S, Crawford RJM, Ryan PG (2008) A junk-food hypothesis for gannets feeding on fishery waste. Proc R Soc B 275: 1149–1156
- Hamer KC, Humphreys EM, Garthe S, Hennicke J and others (2007) Annual variation in diets, feeding locations and foraging behaviour of gannets in the North Sea: flexibility, consistency and constraint. Mar Ecol Prog Ser 338: 295–305
- Hobson KA, Clark RG (1992) Assessing avian diets using stable isotopes I: Turnover of <sup>13</sup>C in tissues. Condor 94: 181–188
- Ingram T, Stutz WE, Bolnick DI (2011) Does intraspecific size variation in a predator affect its diet diversity and topdown control of prey? PLOS ONE 6:e20782
- Kernaléguen L, Arnould JPY, Guinet C, Cherel Y (2015) Determinants of individual foraging specialisation in large marine vertebrates, the Antarctic and Subantarctic

fur seals. J Anim Ecol 84:1081-1091

- Kokko H, Jennions MD (2008) Parental investment, sexual selection and sex ratios. J Evol Biol 21:919–948
- Lecomte VJ, Sorci G, Cornet S, Jaeger A and others (2010) Patterns of aging in the long-lived wandering albatross. Proc Natl Acad Sci USA 107:6370–6375
- Lescroël A, Ballard G, Toniolo V, Barton KJ, Wilson PR, Lyver PO'B, Ainley DG (2010) Working less to gain more: when breeding quality relates to foraging efficiency. Ecology 91:2044–2055
- Lewis S, Sherratt TN, Hamer KC, Wanless S (2001) Evidence of intra-specific competition for food in a pelagic seabird. Nature 412:816–819
- Lewis S, Benvenuti S, Dall'Antonia L, Griffiths R and others (2002) Sex-specific foraging behaviour in a monomorphic seabird. Proc R Soc B 269:1687–1693
- Machovsky-Capuska GE, Senior AM, Benn EC, Tait AH and others (2016a) Sex-specific macronutrient foraging strategies in a highly successful marine predator: the Australasian gannet. Mar Biol 163:75–89
- Machovsky-Capuska GE, Senior AM, Simpson SJ, Raubenheimer D (2016b) The multidimensional nutritional niche. Trends Ecol Evol 31:355–365
- Masello JF, Wikelski M, Voigt CC, Quillfeldt P (2013) Distribution patterns predict individual specialization in the diet of dolphin gulls. PLOS ONE 8:e67714
- Nakagawa S, Schielzeth H (2013) A general and simple method for obtaining R<sup>2</sup> from generalized linear mixedeffects models. Methods Ecol Evol 4:133–142
  - Nelson B (2001) The Atlantic gannet, 2nd edn. Fenix Books, Great Yarmouth
- Oro D, Jover L, Ruiz X (1996) Influence of trawling activity on the breeding ecology of a threatened seabird, Audouin's gull Larus audouinii. Mar Ecol Prog Ser 139:19–29
- Parent CE, Crespi BJ (2009) Ecological opportunity in adaptive radiation of Galápagos endemic land snails. Am Nat 174:898–905
- Parnell AC, Inger R, Bearhop S, Jackson AL (2010) Source partitioning using stable isotopes: coping with too much variation. PLOS ONE 5:e9672
- Patrick SC, Bearhop S, Grémillet D, Lescroël A and others (2014) Individual differences in searching behaviour and spatial foraging consistency in a central place marine predator. Oikos 123:33–40
- Patrick SC, Bearhop S, Bodey TW, Grecian WJ, Hamer KC, Lee J, Votier SC (2015) Individual seabirds show consistent foraging strategies in response to predictable fisheries discards. J Avian Biol 46:431–440
- Peig J, Green AJ (2009) New perspectives for estimating body condition from mass/length data: the scaled mass index as an alternative method. Oikos 118:1883–1891
- Pirotta E, Edwards EWJ, New L, Thompson PM (2018) Central place foragers and moving stimuli: a hidden-state model to discriminate the processes affecting movement. J Anim Ecol 87:1116–1125
  - R Core Team (2017) R: a language and environment for statistical computing. R Foundation for Statistical Computing, Vienna. www.r-project.org
- Ronconi RA, Burger AE (2011) Foraging space as a limited resource: inter- and intra-specific competition among sympatric pursuit-diving seabirds. Can J Zool 89: 356–368
- Roughgarden J (1974) The fundamental and realised niche of a solitary population. Am Nat 108:232–235
- 🔎 Scales KL, Miller PI, Embling CB, Ingram SN, Pirotta E,

Votier SC (2014) Mesoscale fronts as foraging habitats: composite front mapping reveals oceanographic drivers of habitat use for a pelagic seabird. J R Soc Interface 11: 20140679

- Sih A, Cote J, Evans M, Fogarty S, Pruitt J (2012) Ecological implications of behavioural syndromes. Ecol Lett 15: 278–289
- Stauss C, Bearhop S, Bodey TW, Garthe S and others (2012) Sex-specific foraging behaviour in northern gannets *Morus bassanus*: incidence and implications. Mar Ecol Prog Ser 457:151–162
  - Svanbäck R, Bolnick DI (2005) Intraspecific competition affects the strength of individual specialization: an optimal diet theory method. Evol Ecol Res 7:993–1012
- Svanbäck R, Bolnick DI (2007) Intraspecific competition drives increased resource use diversity within a natural population. Proc R Soc B 274:839–844
- Tait A, Raubenheimer D, Stockin KA, Merriman M, Machovsky-Capuska GE (2014) Nutritional geometry of gannets and the challenges in field studies. Mar Biol 161: 2791–2801
- Tinker MT, Guimarães PR Jr, Novak M, Marquitti FMD and others (2012) Structure and mechanism of diet specialisation: testing models of individual variation in resource use with sea otters. Ecol Lett 15:475–483
- van den Hout PJ, van Gils JA, Robin F, van der Geest M, Dekinga A, Piersma T (2014) Interference from adults forces young red knots to forage for longer and in dangerous places. Anim Behav 88:137–146
- Van Donk S, Camphuysen KCJ, Shamoun-Baranes J, van der Meer J (2017) The most common diet results in low reproduction in a generalist seabird. Ecol Evol 7: 4620–4629
- Van Valen L (1965) Morphological variation and width of ecological niche. Am Nat 99:377–390

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- Villegas-Amtmann S, Jeglinski JWE, Costa DP, Robinson PW, Trillmich F (2013) Individual foraging strategies reveal niche overlap between endangered Galapagos pinnipeds. PLOS ONE 8:e70748
- Votier SC, Bearhop S, Witt MJ, Inger R, Thompson D, Newton J (2010) Individual responses of seabirds to commercial fisheries revealed using GPS tracking, stable isotopes and vessel monitoring systems. J Appl Ecol 47: 487–497
- Votier SC, Bicknell A, Cox SL, Scales KL, Patrick SC (2013) A bird's eye view of discard reforms: bird-borne cameras reveal seabird/fishery interactions. PLOS ONE 8:e57376
- Wagner EL, Boersma PD (2011) Effects of fisheries on seabird community ecology. Rev Fish Sci 19:157–167
- Wakefield ED, Bodey TW, Bearhop S, Blackburn J and others (2013) Space partitioning without territoriality in gannets. Science 341:68–70
- Wakefield ED, Cleasby IR, Bearhop S, Bodey TW and others (2015) Long-term individual foraging site fidelity — why some gannets don't change their spots. Ecology 96: 3058–3074
  - Wakefield ED, Owen E, Baer J, Carroll MJ and others (2017) Breeding density, fine-scale tracking, and large-scale modeling reveal the regional distribution of four seabird species. Ecol Appl 27:2074–2091
- Wennersten L, Forsman A (2012) Population-level consequences of polymorphism, plasticity and randomized phenotype switching: a review of predictions. Biol Rev Camb Philos Soc 87:756–767
- Woo KJ, Elliott KH, Davidson M, Gaston AJ, Davoren GK (2008) Individual specialization in diet by a generalist marine predator reflects specialization in foraging behaviour. J Anim Ecol 77:1082–1091
  - Wood SN (2006) Generalized additive models: an introduction with R. CRC Press, Boca Raton, FL

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