The Evolutionary Ecology of Individual Foraging Decisions

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1 Abstract

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Understanding the causes and consequences of animal movement is key to mechanistically linking individual behaviour with population-level patterns. Classical models of individual-3 to-population foraging distributions do not account for the complex and changeable resource landscapes animals must navigate. Neither are the rich behavioural repertoires addressed that animals may exhibit in a foraging context, and their evolution is almost entirely ignored. We take a spatially explicit, individual-based simulation approach to model the evolution of individual movement and foraging strategies, and its consequences for population distributions in three simple foraging scenarios of increasing behavioural complexity. We show that broad equilibrium 9 conditions are reached within only a few generations. This equilibrium obscures ever-changing 10 conditions including the constant redistribution of individuals as they harvest prey-items and steal from each other. Individual movement decisions evolve such that populations can track 12 resource gradients even in the absence of direct cues of landscape quality. We show that klep-13 toparasitism as a foraging strategy is able to rapidly invade a population due to pre-adaptations 14 in the movement mechanism. Phenomenological models of intake rate in relation to competitors 15 and prey-items misrepresent the underlying behavioural mechanisms, and are thus an unreli-16 able guide to evolutionary trajectories. However, these models show that kleptoparasitic foragers 17 are an endogeneously evolved third trophic level whose intake depends on harvesting primary 18 consumers and competition with other kleptoparasites. The behavioural complexity of multiple 19 foraging strategies, a handling time for prey-items, as well as the difficulty of detecting resource 20 gradients in the landscape, result in individuals undermatching their landscape. As a result, 21 populations with low behavioural complexity can still deviate easily from the ideal free distribution. 23

2 Introduction

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3 Methods: Simulation Model of Movement-Behaviour Co-Evolution

Our model is an individual-based evolutionary simulation whose most basic components the environment size and shape, its gridded structure and each cell's capacity to hold multiple 28 individuals, as well as the discrete conception of time within and between generations — is taken from Netz et al. in prep.. We conceptualised the model and the scenarios around the behaviour 30 of waders (Charadrii, and especially oystercatchers Haematopus sp.), which are extensively studied 31 in an optimal foraging context (e.g. Ens et al., 1990; Vahl et al., 2005a,b,c). We simulated a fixed population with a fixed size of 10,000 individuals moving on a landscape of 5122 grid cells, with 33 the landscape wrapped at the boundaries so that individuals passing beyond the bounds at one end re-appear on the diametrically opposite side. Individuals have a lifetime of T timesteps, with T set to 400 by default. After their lifetime, individuals reproduce and transmit their heritable 36 traits proportional to their fitness over their lifetime. The model code (in C++) can be found as 37 part of the Supplementary Material in the Zenodo repository at **Zenodo/other repository here**.

3.1 Flexibility in Foraging Strategies

Our model considers three main scenarios of flexibility in individual foraging strategies. The **first** scenario is an inflexible producer-only case, in which individuals move about on the landscape and probabilistically find and consume discrete prey food items. Between finding and consuming a food item, individuals must 'handle' the prey for a fixed handling time T_H which is constant across prey items. Prey handling time T_H is set at 5 timesteps by default. The handling time dynamic is well known from many systems; for instance, it could be the time required for a wader to break through a mussel shell, with the handling action obvious to nearby individuals,

and the prey not fully under the control of the finder. We refer to such individuals as 'handlers' for convenience. Handlers are assumed to be fully absorbed in their processing of prey, and do not make any movements until they have fully handled and consumed their prey. The second scenario is a fixed-strategy case which adds some flexibility. Individuals at the start of their lifetime each choose between two foraging strategies, which are then fixed through life. The strategy choice is based on local environmental cues, and is covered in "Movement and Foraging Decisions". The two strategies are to produce, i.e., to probabilistically find, handle, and consume discrete prey (as in the producer-only case), or to scrounge as a kleptoparasite, i.e., to steal a found prey item from the individual handling it. We refer to such scroungers as 'kleptoparasites' 55 from here onwards. Kleptoparasites can steal from any handler, regardless of whether that handler acquired its prey by searching or theft. Kleptoparasites are always successful in stealing from the handler they target; this may be thought of as the benefit of the element of surprise, a common observation in nature. Having acquired prey, a kleptoparasite need only handle it for $T_H - t_h$ timesteps, where t_h is the time that the prey has already been handled by its previous handler. The targeted handler deprived of its prey is assumed to flee from the area, and does not make a further movement decision. Thus kleptoparasites clearly save time on handling compared to a producer, and the time saved increases with the handling time T_H of the prey. The **third** scenario is a flexible-strategy case, and individuals are allowed to be plastic in their foraging strategies, and choose between producing and scrounging strategies in each timestep. Apart from the frequency of the choice, the actual foraging dynamics are the same as described in the fixed-strategy case. Individuals move about on the environment, and each foraging strategy choice is based on local environmental cues (see "Movement and Foraging Decisions").

3.2 Movement and Foraging Decisions

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Individuals essentially use cues available in timestep t to predict their best move for the next timestep t + 1, and the strategy associated with that move (when this is allowed). The movement decision is based on three local environmental cues: (1) the number of discrete prey items G,

(2) the number of individuals handling prey H (referred to as 'handlers'), and (3) the number of individuals not handling prey P (referred to as 'non-handlers'). The notation is chosen in keeping with Netz et al. *in prep.*. These cues are available to individuals in all three model scenarios. Individuals occupy a single grid cell on the environment at a time, and assign a suitability score S incorporating S, S, and S per cell to the nine cells in their Moore neighbourhood (including their current cell). Following Netz et al. *in prep.*, individuals calculate the cell-specific S as

$$S = m_{\mathfrak{g}}G + m_{h}H + m_{\mathfrak{p}}P + m_{h} \tag{1}$$

where the weighing factors for each cue m_g , m_h and m_p , and the bias m_b are genetically encoded and heritable between generations. Individuals rank their Moore neighbourhood by S in timestep t and move to the highest ranked cell in timestep t+1.

Individuals in the producers-only case make no foraging decisions and find food items probabilistically (see "Prey Environment and Ecological Dynamics"). In the fixed-strategy case, individuals pick a lifelong foraging strategy in their first timestep (t_0), while in the flexible-strategy case, individuals pick a strategy in each timestep t to be deployed in t+1. Individuals in these latter two cases process the cell-specific environmental cues G, H, and P to determine their foraging strategy F for life (fixed strategy), or in the grid cell into which they have chosen to move in t+1 (flexible strategy). F is determined as

$$F = \begin{cases} producer, & \text{if } f_gG + f_hH + f_pP + f_b \ge 0\\ scrounger, & \text{otherwise} \end{cases}$$
 (2)

where the cue weights f_g , f_h and f_p , and the bias f_b are also genetically encoded and heritable between generations.

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In both latter cases that allow for kleptoparasitism, individuals make their foraging strategy choice for the next timestep after they have passed through the ecological dynamics of their current location. This excludes individuals that have been stolen from are an important exception; these fleeing agents are moved to a random cell within a Chebyshev distance of 5, and do not make a foraging decision there. Thus kleptoparasitism not only gains individuals prey items

while depriving the targeted individual, it also displaces a potential competitor. All individuals move simultaneously, and attempt to implement the foraging strategy chosen for their new location (see below).

3.3 Prey Environment and Ecological Dynamics

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Since our model was initially conceived to represent foraging waders, we developed a resource landscape based on mussels (family *Mytilidae*) that are commonly found in inter-tidal systems. Mussels beds share some important characteristics with other discrete prey items. Firstly, mussels are immobile relative to their consumers, and their abundances are largely driven by extrinsic environmental gradients and very small-scale interactions (de Jager et al., 2020, 2011). Secondly, in common with many ecological systems (Levin, 1992), mussels are not uniformly distributed across the inter-tidal mudflats, and are instead strongly spatially patterned into clusters ('beds') (de Jager et al., 2020, 2011). Thirdly, while prey or their signs in an area are often visible to consumers, consumers are not always certain of obtaining one of these prey, since prey can show small-scale anti-predator avoidance responses.

We captured these essential aspects of prey dynamics when implementing the resource landscape on which our individuals move. We modelled relative prey immobility and extrinsically driven abundance by assigning each grid cell of the resource landscape a constant probability of generating a new prey item per timestep, which we refer to as the growth rate r. We modelled clustering in the abundance of prey by having the distribution of r across the grid cells take the form of 1,024 uniformly distributed resource peaks with r declining from the centre of each peak to its periphery (Figure X). Effectively, the cell at the centre of each patch generates a prey item five times more frequently than the cells at the edges. Thus for a simulation-specific baseline r_{base} = 0.03, the central cell of a resource peak would have an r_{centre} = 0.03, and generate 3 items every 100 timesteps, compared with r_{edge} = 0.006, or 0.6 items generated in 100 timesteps. We ran the simulation with r_{base} values of 0.001, 0.01, 0.03, and 0.05, which we considered a sufficiently broad range. Cells in our landscape were modelled as being able to hold a maximum of K prey items, with the default K = 5. While a cell is at carrying capacity its r is 0. We modelled nearperfect intermediate-range perception but uncertain short-range acquisition of prey by allowing
individuals to perceive all prey items G in a cell, but giving individuals which choose a producer
strategy only a probability of finding one of these prey. The probability of finding a prey item p(success) is given as the probability of not finding any of G prey

$$p(success) = 1 - (1 - p_i)^G \tag{3}$$

where p_i is the detection probability of each of G items, which is uniformly set to 0.2 by default for all items.

Since we model foraging events as occurring simultaneously, it is possible for more producers to be considered successful in finding prey than there are discrete items in that cell. We resolve this simple case of exploitation competition by assigning G prey among some N successful finders at random. Producers that are assigned a prey item in timestep t begin handling it, and are considered to be handlers for the purposes of timestep t+1 (primarily movement and foraging decisions of other individuals). It is important to note that a producer that has converted into a handler in timestep t is not an available target for kleptoparasites until timestep t+1. Producers that are not assigned a prey item are considered idle during timestep t, and are counted as non-handlers for t+1.

Kleptoparasites in the fixed- or flexible-strategy case face a slightly different challenge. All kleptoparasites in a cell successfully steal from a handler, contingent on the number of handlers matching or exceeding the number of kleptoparasites in timestep t. When the number of kleptoparasites exceeds handlers, handlers are assigned among kleptoparasites at random. Successful kleptoparasites convert into handlers, and similar to producer-handlers are unavailable as targets to other kleptoparasites until the next timestep. Unsuccessful kleptoparasites are considered idle, and are also counted as non-handlers for timestep t + 1. A handler that finishes processing its prey in timestep t returns to the non-handler state and is assessed as such by other agents when determining movements for t + 1.

Individuals move and forage on the resource landscape for T timesteps per generation, and T is set at 400 by default. Handling a food item requires a maximum of T_H timesteps, during which the handler is immobile.

3.4 Reproduction and the Evolution of Decision Making

At the end of each generation, the population is replaced by its offspring, maintaining the fixed 154 population size, and the decision-making weights which determine individual movement (m_g , 155 m_h , m_p , m_b) and foraging strategy choice (f_g, f_h, f_p, f_b) are transmitted from parent individuals 156 to offspring. The number of offspring of each parent is proportional to the parent's share of the 157 population fitness, and this is implemented as a weighted lottery that selects a parent for each 158 offspring. The total lifetime intake of individuals is used as a proxy of fitness, and the popu-159 lation's total fitness is its total intake. The decision-making weights are subject to independent 160 random mutations with a probability of 0.001. The size of the mutation (either positive or negative) is drawn from a Cauchy distribution with a scale of 0.01 centred on the current value of the 162 weight to be mutated. This allows for a small number of very large mutations while the majority 163 of mutations are small. We recognised that spatial autocorrelation in the landscape coupled with 164 limited natal dispersal can lead to spatial heterogeneity in evolved movement rules, as lineages 165 adapt to local conditions. Furthermore, limited natal dispersal could lead to population-level 166 movements due to differential reproduction that mirror shifts in resource abundance, rather than 167 individual movement. To ensure that global individual movement rules evolved, we intialised each offspring at a random location on the landscape, and also reset its total intake to zero.

3.5 Simulation Output and Analysis

3.5.1 Decision Making Weights

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To understand the evolutionary consequences of our simulation, we exported the the decisionmaking weights which determine individual movement (m_g, m_h, m_p, m_b) and foraging strategy

choice (f_g, f_h, f_p, f_b) of each individual in every generation of the simulation. We examined how 174 the frequency of these weights changed over the simulation, i.e., how the weights evolved. We 175 visualised weights' evolution after scaling them between -1 and +1 using a hyperbolic tangent 176 function, and binning the scaled values into intervals of 0.1. We refer to these scaled and binned 177 values as phenotypes for convenience. Weights at or near -1 would represent the maximum 178 evolved avoidance of an environmental cue (in relation to a movement weight) or the greatest 179 evolved negative effect of a cue on choosing the foraging strategy (in relation to a strategy choice 180 weight). Similarly, weights at or near +1 represent the greatest evovled preference for or positive 181 effect of a cue on the movement and strategy choice mechanism of an individual. 182

183 3.5.2 Functional and Aggregative Response

In our simulation, individuals perceive and respond to the standing stock of prey items on a 184 cell rather than its productivity, which they cannot sense directly. This standing stock is un-185 predictable due to consumption by other individuals, and the movement (and consumption) of 186 individuals is also unpredictable. To understand the consequences of evolved movement rules, 187 we must investigate how individual intake varies with the presence of items and other indi-188 viduals. Determining the functional response of intake to competitors, and the distribution of 189 predators relative to prey sensu Meer and Ens (1997) is a prevalent method in spatial ecology. Adopting this method, we sought to fit the widely used Beddington model of functional and 191 aggregative response to pseudo-ecological data generated by our simulation. Over the final ten 192 generations of each simulation run, we summed the number of individuals and items on each 193 cell, as well as the total intake on the cell. We were able to record the number of individuals following a forager and kleptoparasite strategy, as well as intake due to foraging or stealing, 195 separately. This allowed us to determine the average per-capita, per-strategy intake on each cell, 196 which we plotted against the number of competing agents on the cell (Figure X). Additionally, 197 we plotted the average number of individuals following each strategy against the number of 198 prey items on the cell (Figure Y). In both cases, we used data only from the second half of each 199

generation so as to capture the system in a state of ecological equilibrium.

3.5.3 Distribution Along Productivity Gradients

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While recognising that individuals move in response to their rapidly-changing prey landscape, 202 it is useful to determine how individuals distribute along more slowly-changing productivity 203 gradients; this is because these may often be easier to measure in the real world. The ideal 204 free distribution (IFD) and the matching rule robustly predict that individuals should distribute 205 themselves such that intake rates are equalised over patches of similar productivity. The large 206 volume of pseduo-ecological data generated by our simulation allowed us to test whether intake 207 rates were indeed equalised over the productivity gradient. Having previously calculated the 208 average numbers of each strategy, and the average per-captia intake for each strategy on each cell, we plotted both against the productivity of the cell (Figure Z). Here too, we used data only 210 from the second half of each generation to approximate ecological equilibrium. 211

Data Availability. Simulation data used in this study are available on the Dryad/IRODS/Zenodo repository REPOSITORY LINK HERE; simulation code is available on Github and archived on Zenodo at ZENODO LINK HERE; data analysis and figure code is available on Github and archived on Zenodo at ZENODO LINK HERE.

4 Results: Simulation Model Outcomes

4.1 Emergence of an Evolutionary Equilibrium

Work in progress: Dynamic equilibrium emerges within 1000 generations — equilibria are identical across replicates — kleptoparasites emerge in most replicates when allowed — kleptoparasites
establish as a stable and significant proportion of individuals, or individual strategies.

4.2 Movement-Behaviour Co-Evolution

222 4.2.1 Movement-Behaviour Co-Evolution with Exploitative Competition

23 4.2.2 Movement-Behaviour Co-Evolution with Kleptoparasitism

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4.3 Evolution of Kleptoparasitism Requires Movement Pre-Adaptation

The scrounging kleptoparasitic strategy evolves and is established in populations in some tens of 225 generations, and emerges relatively quickly in the evolutionary history of populations (see Fig-226 ure X) This rapid emergence and invasion is made possible by the pre-adaptation of individuals to use the kleptoparasitic strategy successfully. Scenarios 2 and 3 prior to the emergence and 228 establishment of kleptoparasitism are identical to scenario 1, and all individuals are producers. 229 Producers evolve to move towards both items and handlers at most regrowth rates (Figure X), 230 since these are cues to the immediate benefit, and the regrowth rate of a cell, respectively. For the kleptoparasitic strategy, the mapping of cues is reversed but the direction of preference re-232 mains the same. To kleptoparasites, the number of handlers indicates the immediate resource 233 abundance, while the number of items indicates the probability of resource generation, i.e., indi-234 viduals converting into handlers. This coincidental alignment of movement decisions with either behavioural strategy is essential to the persistence of kleptoparasitism. 236

The initial evolution of kleptoparasitism is then only conditional on the mutation of any one of the strategy weights to a sufficiently negative value such that the individual attempts to steal rather than search for prey. At very high regrowth ($r_{base} = 0.25$), the landscape is saturated with prey-items, and individuals can ignore the presence of handlers and evolve to move only in response to prey-items ('socially naive producers'). Under such circumstances in **scenario** 2, though strategy weight mutations lead to some few individuals using a fixed kleptoparasitic strategy, they do not move optimally for their strategy. Thus kleptoparasitism as a fixed-strategy repeatedly evolves and goes extinct in high-productivity environments, as these individuals find themselves in a 'desert of plenty'. Under the same conditions in **scenario** 3 however, a mixed

foraging strategy allows individuals to be producers when appropriate, and yet steal a march on pure-producers when kleptoparasitism is possible.

4.4 Functional Response of Intake

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Understanding the consequences of aggregating with other individuals is key to explaining why 249 animals form groups. We examined the functional response of individual intake of each be-250 havioural strategy on a cell to the number of individuals on that cell. The foragers-only case 251 presents a useful starting point: forager intake is invariant with individual (forager) density, and 252 only declines at very high or low densities (Fig. 4.a,d). Similarly, the functional response of 253 both foragers and kleptoparasite strategies in the fixed-strategy and conditional-strategy case is 254 hump-shaped, with an apparently 'optimal' competitor density at which individual intake rates 255 are maximised (Fig. 4.b,c). Furthermore, the kleptoparasitic strategy's per-capita intake is always 256 greater than that of the forager strategy (Fig. 4.b,c). However on separating potential competi-257 tors by strategy, we find that the individual intake of both strategies increases with increasing 258 forager density (Fig. 4.e,f), but decreases with increasing kleptoparasite density (Fig. 4.g,h). 259 These consistently opposite responses to foragers and kleptoparasites explain why the overall 260 functional response to all competitors appears hump-shaped. With growing forager densities, 261 exploitative, scramble competition for prey items is increased, but foragers also accumulate on 262 high-productivity cells, increasing average per-capita intake overall. When kleptoparasites ac-263 cumulate, however, interference competition results in both lower extraction (as only foragers extract prey) as well as lower intake, as the same item is repeatedly passed between agents 265 in stealing interactions. Thus we show that taking the type of competition, and behavioural 266 variation among individuals more generally into account is crucial to correctly understand the 267 consequences of multiple competing individuals foraging on the same patch (or in the same group/in proximity to each other). 260

4.5 Population Distribution

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Predicting the distribution of a population on a heterogeneous landscape is among the key goals of spatial ecology. This is often achieved by finding the aggregative response of individuals to a 272 relatively static indicator of landscape quality; i.e., how many individuals are present on patches 273 or areas with different resource levels. We examined the aggregative response of individuals in 274 our simulations in relation to landscape productivity, to compare and contrast with predictions from optimal foraging theory (Meer and Ens, 1997, ; see Fig. 5). We find that in the foragers-276 only case, individuals follows the matching rule in relation to grid-cell productivity, with more 277 foragers on higher productivity cells (Fig. 5.a). In the fixed-strategy case, the aggregative re-278 sponse of foragers and kleptoparasites differs; forager counts peak on lower productivity cells and declines with further increases in productivity. Kleptoparasite counts initially increase with 280 cell productivity and then stabilise (Fig. 5.b). In the conditional-strategy case, both foragers 281 and kleptoparasite counts peak on intermediate productivity cells and then begin to decline (Fig. 282 5.c). In the spatial context of our simulation, this translates to three distinct patterns with (1) 283 individuals clustered on productivity peaks in scenario 1, (2) kleptoparasites dominating pro-284 ductivity peaks with fewer foragers in scenario 2, and (3) individuals using a forager strategy 285 more frequently than a kleptoparasitic strategy on productivity peaks in scenario 3. We further 286 find that there are appreciable differences between average and median counts of individuals of 287 each strategy on cells, with the mean typically higher than the median. 288

4.6 Kleptoparasitism and Predation Have Similar Landscape Effects

Work in progress: Kleptoparasite intake depends only on the number of individuals — kleptoparasite intake is highest when cells have few kleptoparasites and many producers — kleptoparasite emergence is associated with a reduction in item depletion — add also landscape metrics etc

5 Discussion

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6 Conclusion

7 Acknowledgments

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Literature Cited

- de Jager, M., J. van de Koppel, E. J. Weerman, and F. J. Weissing. 2020. Patterning in Mussel Beds
 Explained by the Interplay of Multi-Level Selection and Spatial Self-Organization. Frontiers in
 Ecology and Evolution 8.
- de Jager, M., F. J. Weissing, P. M. J. Herman, B. A. Nolet, and J. van de Koppel. 2011. Lévy
 Walks Evolve Through Interaction Between Movement and Environmental Complexity. Science
 332:1551–1553.
- Ens, B. J., P. Esselink, and L. Zwarts. 1990. Kleptoparasitism as a problem of prey choice: A study on mudflat-feeding curlews, Numenius arquata. Animal Behaviour 39:219–230.
- Levin, S. A. 1992. The Problem of Pattern and Scale in Ecology: The Robert H. MacArthur Award

 Lecture. Ecology 73:1943–1967.
- Meer, J. V. D., and B. J. Ens. 1997. Models of Interference and Their Consequences for the Spatial
 Distribution of Ideal and Free Predators. The Journal of Animal Ecology 66:846.
- ³¹³ Vahl, W. K., T. Lok, J. van der Meer, T. Piersma, and F. J. Weissing. 2005a. Spatial clumping of

- food and social dominance affect interference competition among ruddy turnstones. Behavioral
- Ecology 16:834–844.
- Vahl, W. K., J. van der Meer, F. J. Weissing, D. van Dullemen, and T. Piersma. 2005b. The
- mechanisms of interference competition: Two experiments on foraging waders. Behavioral
- Ecology 16:845–855.
- 319 . 2005c. The mechanisms of interference competition: Two experiments on foraging
- waders. Behavioral Ecology 16:845–855.

- 8 Appendix A: Supplementary Figures
- 8.1 Fox-dog encounters through the ages

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9 Appendix B: Additional Methods

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9.1 Measuring the height of fox jumps without a meterstick

Tables 10 Tables

11 Figure legends

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11.1 Online figure legends