# The joint evolution of movement and competition strategies

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# **Abstract**

Competition typically takes place in a spatial context, but eco-evolutionary models rarely address the the joint evolution of movement and competition strategies. Here we investigate a spatially explicit producer-scrounger model where consumers can either forage on a heterogeneous resource landscape or steal prey from conspecifics (kleptoparasitism). We compare different scenarios for the interaction of movement and competition strategies. In all cases, movement strategies evolve rapidly and consistently across replicate simulations. At equilibrium, foragers do not match the prey input rate, contrary to 'ideal free' predictions. We show that this is related to the intrinsic difficulty of moving effectively on a depleted landscape with few reliable cues for movement. The evolved movement strategies of kleptoparasites differ markedly from those of foragers. Even within each competition strategy, polymorphisms emerge, corresponding to individual differences in movement rules. Our study emphasises the advantages of a mechanistic approach when studying competition in a spatial context.

# 1 Introduction

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Intraspecific competition is an important driver of population dynamics and the spatial dis-15 tribution of organisms (Krebs and Davies, 1978), and can be broadly classified into two main 16 types, 'exploitation' and 'interference'. In exploitation competition, individuals compete in-17 directly by depleting a common resource, while in interference competition, individuals com-18 pete directly by interacting with each other (Birch, 1957; Case and Gilpin, 1974; Keddy, 2001). 19 A special case of interference competition which is widespread among animal taxa is 'kleptoparasitism', in which an individual steals a resource from its owner (Iyengar, 2008). Experi-21 ments with foraging birds have shown that competition, including kleptoparasitism, can affect the spatial distribution of individuals across resource patches (Goss-Custard, 1980; Vahl et al., 23 2005a,b, 2007; Rutten et al., 2010a). The avoidance of competitive interactions can also affect the distribution and behaviour of animals foraging in groups (Rutten et al., 2010b; Bijleveld et al., 25 2012). At larger scales, competition among different behavioural types in a species can strongly influence species distributions and animal movement decisions (e.g. Duckworth and Badyaev, 27 2007; Schlägel et al., 2020). The fine-scale mechanisms and evolutionary consequences of com-28 petition are difficult to study in free-living animals, yet their knowledge is central to evolution-29 ary ecology. It is nearly impossible to study competition and its coevolution with movement strategies at evolutionary time-scales in most animals, due to a lack of long-term data (Clutton-31 Brock and Sheldon, 2010) — this necessitates the use of models. 32 Competition is key to paradigms of animal space-use, including the ideal free distribution 33 (IFD, Fretwell and Lucas, 1970), information-sharing based local enhancement (also called 34 conspecific aggregation Giraldeau and Beauchamp, 1999; Folmer et al., 2012), and producer-35 scrounger dynamics (Barnard and Sibly, 1981; Vickery et al., 1991; Beauchamp, 2008). Yet these paradigms often treat competition in highly simplified ways. Most IFD models, for instance, 37 assume that resource depletion is either negligible, or that resources have a constant influx 38 rate, where standing stock densities are effectively zero due to immediate consumption (con-39

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tinuous input/immediate consumption models, Tregenza, 1995; van der Meer and Ens, 1997).

Other IFD models that do include resource depletion make simplifying assumptions about the

importance of interference competition, considering it unimportant, or even modelling a ben-

efit of grouping (e.g. Amano et al., 2006). Producer-scrounger models are primarily concerned
with the benefits of choosing either a producer or scrounger strategy in relation with local
conditions, such as the number of conspecifics (Vickery et al., 1991), or the order of arrival on
a patch (Beauchamp, 2008). While models such as Beauchamp (2008) and Tania et al. (2012)
allow for individuals to either search for food or scrounge off exploited patches, they simplify
the mechanisms by which this decision is made (see also Holmgren, 1995; Garay et al., 2020).
Few foraging ecology models — either IFD or producer-scrounger — allow for realistic movement strategies, often assuming omniscient animals with no travel costs (Vickery et al., 1991;
Tregenza, 1995; van der Meer and Ens, 1997, but see Amano et al. 2006; Cressman and Křivan
2006).

On the contrary, competition occurs in a spatial context, and spatial structure is key to for-53 aging (competition) decisions (Beauchamp, 2008). Consequently, the abundance of resources and their depletion, as well as the presence of potential competitors is of obvious importance 55 to individuals' movement decisions (resource selection, sensu Manly et al., 2007). How an-56 imals are assumed to integrate the costs (and potential benefits) of competition into their 57 movement decisions has important consequences for theoretical expectations of population 58 distributions (van der Meer and Ens, 1997; Hamilton, 2002; Beauchamp, 2008). In addition to 59 short-term, ecological effects, competition should also have evolutionary consequences for in-60 dividual movement strategies, as it does for so many other aspects of behaviour (Baldauf et al., 61 2014), setting up feedback loops between ecology and evolution. Modelling competition and 62 movement decisions jointly is thus a major challenge. A number of models take an entirely 63 ecological view, assuming that individuals move or compete ideally, or according to some fixed strategies (Vickery et al., 1991; Holmgren, 1995; Tregenza, 1995; Amano et al., 2006, but 65 see Hamilton 2002). Models that include evolutionary dynamics in the movement (de Jager 66 et al., 2011, 2020) and foraging competition strategies (Beauchamp, 2008; Tania et al., 2012) are 67 more plausible, but they too make arbitrary assumptions about the functional importance of environmental cues to individual decisions. Ideally, both movement and competition strategies should be the joint outcomes of selection, allowing for different competition strategies to be associated with different movement rules (see the approach in Getz et al., 2015). 71

Here, we present a first mechanistic, model of intraspecific competition in a spatially explicit

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context, where competition is shaped by the joint evolution of foraging competition and move-73 ment strategies. As foraging and movement decisions are taken by individuals, we study the 74 joint evolution of both types of decision-making by means of a simulation model. Such mod-75 els are well suited to modelling the ecology and evolution of complex behaviours (Guttal and 76 Couzin, 2010; Kuijper et al., 2012; Getz et al., 2015, 2016; White et al., 2018; Long and Weiss-77 ing, 2020; Netz et al., 2020, for conceptual underpinnings see Huston et al. (1988); DeAngelis and Diaz (2019)). This allows us to both focus more closely on the interplay of exploitation 79 and interference competition, and to examine the feedback between movement and foraging 80 behaviour at ecological and evolutionary timescales. In our model, foraging individuals move 81 on a spatially fine-grained resource landscape with discrete, depletable food items that need 82 to be processed ('handled') before consumption. Foragers make movement decisions using an inherited (and evolvable) strategy which integrates local cues, such as the local resource and competitor densities. After each move, individuals choose between two foraging strategies: 85 whether to search for a food item or steal from another individual; the mechanism underly-86 ing this foraging choice is also inherited. We take lifetime resource consumption as a proxy 87 for fitness, such that more successful individuals produce more offspring, and thus are more 88 successful in transmitting their movement and foraging strategies to future generations (subject to small mutations). We consider three scenarios: in the first scenario, we examine only 90 exploitation competition. In the second scenario, we introduce kleptoparasitic interference as 91 an inherited strategy, fixed through an individual's life. In the third scenario, we model klep-92 toparasitism as a behavioural strategy conditioned on local environmental and social cues.

Our model allows us to examine the evolution of individual movement strategies, populationlevel resource intake, and the spatial structure of the resource landscape. The model enables
us to take ecological snapshots of consumer-resource dynamics (animal movement, resource
depletion, and competition) proceeding at evolutionary time-scales. Studying these snapshots
from all three scenarios allows us to check whether, when, and to what extent the spatial distribution of competitors resulting from the co-evolution of competition and movement strategies corresponds to standard IFD predictions. Using this model, we investigate three primary
questions: (1) Under what conditions does kleptoparasitic interference evolve and persist in
the population? How do the movement strategies of kleptoparasites differ from those of the

foragers? (2) What are the eco-evolutionary implications of conditional kleptoparasitism? Do
conditional strategies evolve under broader conditions than a polymorphism of fixed pure
strategies, and do they lead to a different spatial distribution of competitors? (3) To what extent does the spatial distribution of competitors and resources in space correspond to an ideal
free distribution?

# 108 2 The Model

We implement an individual-based evolutionary simulation model whose most basic compo-109 nents — the environment's gridded structure, and each cell's capacity to hold multiple in-110 dividuals, as well as the discrete conception of time within and between generations — are 111 inspired by the behaviour of waders (Charadrii). Waders are extensively studied in the context 112 of foraging competition, both empirically (e.g. Vahl et al., 2005a,b, 2007; Rutten et al., 2010a,b), and using IBMs (reviewed in Stillman and Goss-Custard, 2010) — but few, if any, models consider the interplay of ecology and evolution when studying wader movement and competition. We simulated a population with a fixed size moving on a landscape of 512<sup>2</sup> grid cells, 116 with the landscape wrapped at the boundaries so that individuals passing beyond the bounds 117 at one end re-appear on the diametrically opposite side. The model has two time scales, first, 118 an ecological (or behavioural) time scale of T timesteps, during which individuals move, make foraging decisions, and handle prey items they find or steal. Individuals are modelled as being 120 immobile while handling food items, creating the conditions for kleptoparasitism (Brockmann 121 and Barnard, 1979; Holmgren, 1995). On the second, evolutionary time scale, individuals re-122 produce and pass on their movement and foraging strategies to their offspring, the number of 123 which is proportional to their intake at the behavioural time scale. By default, we set T to 400, 124 and simulated 1,000 generations of a population of 10,000 individuals; thus there are about 26 125 times more grid cells than individuals. 126

### **2.1 Resource Landscape**

**Prey Abundance** We considered a resource landscape that is heterogeneous in its productivity of discrete resources, but with strong spatial clustering of grid cells of similar productivity

(see Fig. 1C; panel gen: 1, showing a subset of  $60^2$  grid cells). We assigned each cell a constant 130 probability of generating a new prey item per timestep, which we refer to as the cell-specific 131 growth rate r. We modelled clustering in landscape productivity by having the distribution of 132 r across the grid take the form of 1,024 resource peaks, placed at regular distances of 16 grid 133 cells from the peaks around them; r declines from the centre of each peak (called  $r_{max}$ ) to its pe-134 riphery (see Fig. 1C). Effectively, the cell at the centre of each cluster generates a prey item five 135 times more frequently than the cells at the edges. We ran all three scenarios at a default  $r_{max}$  of 136 0.01, and also across a range of  $r_{max}$  values between 0.001 and 0.05. For an  $r_{max} = 0.01$ , the most 137 productive cells (at the centres of a cluster) are likely to generate one item per 100 timesteps (or 138 four items per generation, for T = 400), while the least productive cells (at cluster peripheries) 139 are likely to generate one item every 500 timesteps (only about one item per generation, for T = 400). Since our model was conceived to represent foraging waders, we considered our resources to represent mussels, a common prey of many waders, whose abundances are largely 142 driven by external gradients; we refer to these resources as 'prey items' henceforth. Cells in 143 our landscape were modelled as having a uniform carrying capacity K of 5 prey items, and 144 while a cell is at carrying capacity its *r* is 0.

**Prey Acquisition by Foragers** Foragers can perceive a cue indicating the number of all prey items *P* in a cell, but do not know the exact locations of these prey. We model foragers as 147 having a probability q of failing to detect a prey item, and a probability  $q^P$  of not detecting any 148 of P prey items; foragers are thus successful in finding a prey item with a probability  $1 - q^P$ . 149 Individuals on a cell forage in a randomised sequence, and the probability of finding a prey item  $(1-q^P)$  is updated as individuals find prey, reducing P. Foragers that are assigned a prey 151 item in timestep t begin handling it, and are considered to be handlers from the end of the 152 timestep, i.e., for the movement and foraging decisions of other individuals. Foragers that are 153 not assigned a prey item are considered idle, and are counted as non-handlers. 154

### 2.2 Movement and Competition Strategies

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Movement Strategies We model movement as a fine-scale process comprised of small, discrete steps of fixed size. These steps are the outcome of short-term individual movement deci-

sions, which are made using evolved movement strategies. Movement decisions are modelled 158 as the individual selection of a destination cell, after assessing potential destinations based on 159 available cues (akin to the concept of step selection or resource selection Fortin et al., 2005; 160 Manly et al., 2007), an approach used previously by Getz et al. (2015, 2016) and White et al. 161 (2018). At the end of each timestep t, individuals scan the nine cells of their Moore neighbour-162 hood for three environmental cues, (1) an indication of the number of discrete prey items P, (2)the number of individuals handling prey H (referred to as 'handlers'), and (3) the number of 164 individuals not handling prey N (referred to as 'non-handlers'). Based on these cues, a 'suit-165 ability score' S is assigned to each cell as  $S = s_P P + s_H H + s_N N$ . At the start of timestep t + 1, 166 each individual moves to the cell to which it assigned the highest suitability. The weighing 167 factors for each cue,  $s_P$ ,  $s_H$ , and  $s_N$ , are genetically encoded and and transmitted from parents to their offspring. All individuals move simultaneously, and then implement their foraging or kleptoparasitic behaviour to acquire prey. Individuals move and forage on the resource 170 landscape for *T* timesteps per generation. 171

**Scenario 1: Exploitative Competition** The first scenario simulates only exploitative competition; individuals (henceforth called 'foragers') move about on the landscape and probabilistically find and consume prey items. A forager can be either in a 'searching' state or in 174 a 'handling' state, as we assume that between finding and consuming a prey item individu-175 als must handle each prey for a fixed handling time  $T_H$  (set at 5 timesteps by default). The 176 handling time dynamic is well known from many systems; for instance, it could be the time re-177 quired for an oystercatcher to break through a mussel shell, or the time between catching and subduing prey for raptors, with the handling action obvious to nearby individuals, and the 179 prey not fully under the control of the finder (Brockmann and Barnard, 1979). Handlers are as-180 sumed to be fully absorbed in their processing of prey, and do not make any movements until 181 they have fully handled and consumed their prey. In scenario 1, the only evolvable properties 182 are the cue weighing factors which determine the suitability scores ( $s_P$ ,  $s_H$  and  $s_N$ ). 183

Scenario 2: Fixed Interference Competition The second scenario builds on Scenario 1, but in addition to foragers there is a second category of individuals called 'kleptoparasites'.

Kleptoparasites do not search for food, but rather for handlers from which they can steal 186 prey (similar to Holmgren, 1995). Agents that steal are termed kleptoparasites. Kleptopar-187 asites are always successful in stealing from a handler; this may be thought of as the benefit of 188 the element of surprise, a common observation among birds (Brockmann and Barnard, 1979). 189 However, if multiple kleptoparasites target the same handler, only one of them, randomly 190 selected, is considered successful — thus kleptoparasites also compete exploitatively among themselves. Individuals that have been stolen from subsequently 'flee' and are moved to a 192 random cell within a Chebyshev distance of 5. Having acquired prey, a kleptoparasite con-193 verts into a handler, but need only handle prey for  $T_H - t_h$  timesteps, where  $t_h$  is the time 194 that the prey has already been handled by its previous owner; thus kleptoparasites save time 195 on handling compared to a forager. Unsuccessful kleptoparasites are considered idle, and are also counted as non-handlers. Handlers that finish processing their prey in timestep t return to the non-handler state and are assessed as such by other individuals when determining their 198 movements. In scenario 2, there are two fixed competition strategies that are inherited from 199 parents to offspring, and each of these strategies can evolve a (separate) movement strategy. 200

**Scenario 3: Conditional Interference Competition** In scenario 3, each individual can either act as a forager, searching for food, or as a kleptoparasite, dependent on its local circumstances. Individuals process the cell-specific environmental cues P, H, and N to determine their location in the next timestep (based on their inherited movement strategy). Additionally, individuals process the environmental cues to determine their strategy in the next timestep as

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strategy = 
$$\begin{cases} \text{forager,} & \text{if } w_P P + w_H H + w_N N \ge w_0 \\ \text{kleptoparasite,} & \text{otherwise} \end{cases}$$
 (1)

where the cue weights  $w_P$ ,  $w_H$  and  $w_N$ , and the threshold value  $w_0$  are also genetically encoded and heritable between generations. Apart from the ability to switch between foraging and kleptoparasitism, the foraging dynamics are the same as described in the fixed-strategy case.

### 210 2.3 Reproduction and Inheritance

For simplicity, our model considers a population of fixed size (10,000 individuals) with discrete, non-overlapping generations. Individuals are haploid and reproduction is asexual. Each individual has 7 gene loci that encode the decision making weights; only the weights in control of individual movement ( $s_P$ ,  $s_H$ ,  $s_N$ ) are active in scenarios 1 and 2. In scenario 3, the weights for foraging decisions ( $w_P$ ,  $w_H$ ,  $w_N$ ,  $w_0$ ) are also active, and are transmitted from parent individuals to offspring. Hence the alleles at these loci correspond to real numbers that are transmitted from parent individuals to their offspring.

We assume that the expected number of offspring per individual is proportional to the individual's total lifetime intake of resources (hence resource intake is used as a proxy for fitness). This is implemented as a weighted lottery (with weights proportional to lifetime resource intake) that selects a parent for each offspring in the subsequent generation (see prior implementation in Tania et al., 2012; Netz et al., 2020). In all scenarios, the movement decision-making weights are subject to independent random mutations with a probability of 0.001. The mutational step size (either positive or negative) is drawn from a Cauchy distribution with a scale of 0.01 centred on zero. This allows for a small number of very large mutations while the majority of mutations are small. In scenarios 1 and 2, the foraging-decision weights are not relevant. However, in scenario 2 we allow a forager to mutate into a kleptoparasite (or *vice versa*) with a probability of 0.001. In scenario 3, the foraging weights mutate in the same way as the movement weights, described above. We intialised each offspring at a random location on the landscape; this potentially forces individuals to contend with conditions very different from those of their direct parent.

### 2.4 Simulation Output and Analysis

Population Activities and Intake We counted the number of times foragers were searching for prey, kleptoparasites were searching for handlers, and the number of timesteps that individuals of either strategy were handling a prey item. We refer to the ratio of time spent foraging, stealing, and handling as the population's 'activity budget'. We examined how the population activity budget developed over evolutionary time, and whether a stable ecological

equilibrium was reached. Furthermore, we counted the population's mean per capita intake
per generation as a measure of population productivity.

Resource Landscape and Individual Distribution Snapshots To visualise the effect of different foraging strategies on the resource landscape, we exported snapshots of the entire simulation landscape at the mid-point of each generation (t = 200). This snapshot contained data on (1) the number of prey items, (2) the number of handling individuals, and the number of individuals using either a (3) searching strategy or (4) kleptoparasitic strategy, on each grid cell. We used only a subset of the total landscape ( $60^2$  of  $512^2$  cells) for further analyses to speed up computation.

Testing the Input Matching Rule To examine whether foragers in our model achieved an IFD, we used the snapshots to test a basic prediction of the IFD and the related matching rule: that the number of individuals on occupied patches should be proportional with patch productivity (Fretwell and Lucas, 1970; Parker, 1978; Houston, 2008). In real world systems, patch productivity is challenging to measure, but not so in simulation models, where it is among the model's building blocks. We calculated the correlation coefficient between the number of individuals (excluding handlers) and the cell-specific productivity r, and plotted it over generations for each replicate.

Informativeness of Resource Landscapes Another measure of whether foragers have achieved
the IFD on their resource landscape is whether individuals can improve their intake by moving to another location; in our model, this would be to a neighbouring cell. We calculated the
cell-specific item gradient for each landscape snapshot, as the difference in item counts between each cell and its neighbouring cells. We then calculated the proportion of grid cells from
which it was possible to sense differences in prey items, i.e., a cell with either more of fewer
items. When it is impossible to sense such differences, movement informed by relative prey
item abundances is difficult, and achieving the IFD becomes a challenge.

Visualising Decision-Making Weights To understand the evolutionary consequences of our simulation on the individual decision making weights, we exported the weights of each indi-

vidual in every generation of the simulation. To depict as much as possible of the typical variation of weights, which could take arbitrarily large values and therefore vary by orders of magnitude, we multiplied each weight by 20 and applied a hyperbolic tangent transform. This scaled the weights between -1 and +1.

Model Code, Analysis Code, and Data Availability The simulation model code can be found on Github: github.com/pratikunterwegs/Kleptomove and archived on Zenodo: zenodo.org/record/4905476. Simulation data used in this study are available on the DataverseNL repository: doi.org/10.34894/JFSC41. Code for data analysis and preparing the figures in the main text and Supplementary Material is available on Github: github.com/pratikunterwegs/kleptomovems and archived on Zenodo: doi.org/10.5281/zenodo.4904497.

### 3 Results

## 3.1 Scenario 1: No Kleptoparasitism

When only foragers are present in the population, individuals deplete prey items faster than 277 they can be replenished, and the overall number of prey items is drastically reduced within 278 only a few generations (Fig. 1A). The population's activity budget is split between searching 279 and handling (Fig. 1B). The proportion of time spent handling and the population's mean 280 per capita intake are both initially low, but then peak within ten generations (Fig. 1C). This 281 is because individuals can easily acquire prey items from the fully stocked landscape in the 282 first few generations. As the number of prey items reduces, handling as a share of the activ-283 ity budget declines to a stable  $\sim 45\%$  within 50 generations; this is because fewer searching 284 foragers find a prey item. This leads to a similar stabilisation in mean per capita intake (Fig. 285 1C). Though in early generations, foragers are attracted to resource peaks and tend to accu-286 mulate on resource-rich locations (Fig. 1A), the correlation between the number of foragers 287 on a cell and the productivity of that cell is only slightly positive (Fig. 1D). This is in contrast 288 with the perfect correspondence between resource input rate and forager density (the 'input 289 matching rule'), which is a defining property of the IFD (Parker, 1978; Houston, 2008). Con-290 trary to standard IFD assumptions, foragers cannot directly "sense" the local cell productivity

r; instead they can only use the (small) number of prey items available in a cell as a cue for local productivity ("cell quality").

### 294 3.2 Scenario 2: Co-existence of Foragers and Kleptoparasites

When both foraging and kleptoparasitism is allowed, the spatial distribution of consumers and 295 prey items at equilibrium are very different from a forager-only population. Consumers gather on resource peaks in early generations, grazing them down until very few prey items remain 297 on the landscape and consumers are dispersed; however, within 50 generations the resource 298 landscape recovers with prey abundances higher than in the earliest generations (Fig. 2A). 299 This is explained by the presence of kleptoparasites (Fig. 2B), which reduces the prey intake 300 rate of the population, and accordingly lead to a higher prey abundance. In early generations, kleptoparasites are very rare, and the activity budget, the population mean per capita intake, 302 and the distribution of consumers over the landscape show the same patterns as in Fig. 1. 303 Mean per capita intake also spikes in early generations as individuals successfully acquire 304 prey items from the fully stocked prey landscape (Fig. 2C). At this stage, it becomes more 305 likely for a kleptoparasite to encounter a handler than for a searching forager to find a prey 306 item, and from generation 30 onwards, as kleptoparasitism becomes the dominant strategy 307 (a stable  $\sim$ 70% of the population; see Fig. 2B), searching for handlers from whom to steal 308 becomes the dominant activity. However, the high frequency of this activity, and the low 309 frequency of handling, indicate that many kleptoparasites are actually unsuccessful at finding 310 handlers. With few searching foragers, fewer prey items are extracted from the landscape, 311 which recovers beyond its initial prey abundance within 50 generations (Fig. 2A).

Despite the strong spatial structure of the resource landscape within 50 generations, the correlation between consumer abundance and cell productivity remains weak or zero across generations (Fig. 2D). This may be explained by the dynamics of kleptoparasitism: foragers fleeing a kleptoparasitic attack are displaced far from their original location, and kleptoparasites must track these foragers if they are to acquire resources. As kleptoparasites, the numerically dominant strategy, seek to move towards handlers (rather than prey), they are not strongly influenced by prey item abundances.

The increase of kleptoparasitism from a negligible fraction to the dominant strategy (Fig.

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3A) is associated with the evolutionary divergence of the movement strategies in foragers and 321 kleptoparasites. While foragers and kleptoparasites respond to prey density and non-handler 322 density in a similar way (see Supplementary Figure S5), preferring higher prey density while 323 avoiding high non-handler density, the two types of competition strategy differ substantially 324 in their response to handlers (Fig. 3B, 3C). Kleptoparasites very rapidly (within 3 generations) 325 evolve a strong preference for moving towards handlers, which are their primary resource 326 (Fig. 3B). In the absence of kleptoparasites, foragers would evolve a preference for moving 327 towards handlers (see Supplementary Figure S4), because the presence of handlers is an indi-328 rect indicator of productive cells. However, with kleptoparasites common in the population, 329 searching foragers both avoid and prefer handlers in about equal proportions (Fig. 3C). While 330 all kleptoparasites evolve to prefer moving towards handlers, the strength of the attraction to 331 handlers shows multiple, distinct values or 'morphs', which are remarkably persistent across 332 generations (Fig. 3B). In replicate 3, for example, the dominant movement strategy exhibits 333 only a weak attraction to handlers, but this strategy coexists with various strategies that all 334 show a much stronger attraction to handlers (Fig. 3B). The movement strategies of foragers 335 show an even higher degree of polymorphism (Fig. 3C). Typically, there are no predominant 336 movement strategies. Instead, a wide range of coexisting handler attraction/repulsion values 337 emerges. In other words, some foragers are strongly attracted by handlers, others are strongly 338 repelled by handlers, and yet others are neutral to the presence of handlers. 339

### 3.3 Scenario 3: Condition-dependent Kleptoparasitism

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When individuals are allowed to choose their competition strategy (foraging or kleptopara-341 sitism) based on local environmental cues, the distribution of individuals and prey items is 342 substantially different from the two previous scenarios (Fig. 4A). Initially, as in scenario 1, 343 depletion of prey items results in the degradation of the resource landscape within ten gener-344 ations. By generation 50, the resource landscape recovers some of the spatial structure of the 345 first generation, but prey-item abundances do not reach the level of the similar recovery seen in scenario 2. This too is explained by the observation that by generation 30, all individuals 347 have a propensity to steal from handlers, i.e., when handlers are present in the vicinity, con-348 sumers will choose to target handlers for prey items, rather than forage for prey themselves 349

("opportunistic kleptoparasitism"; Fig. 4B; orange line). However, unlike scenario 2, individu-350 als search for prey more often and steal less (at or below 25%; compare with Fig. 2B), leading 351 to a sustained high mean per capita intake rate and the depletion of the resource landscape. 352 Consequently, mean per capita intake stabilises (after an initial spike, as in scenarios 1 and 2) 353 354 within ten generations to a level similar to scenario 1 (Fig. 4C). The reduced depletion following the evolution and persistence of kleptoparasitism leads to a resource landscape recovery 355 intermediate between scenarios 1 and 2 within 50 generations (Fig. 4A). As individuals move 356 and choose foraging strategies conditionally, they are able to better choose between the payoff 357 of more prey items, more handling foragers from which to steal, and the risk of falling victim to 358 kleptoparasites. Thus, while not as strong as predicted by IFD theory, the correlations between 359 consumer abundance and cell productivity are weakly positive (Fig. 4D).

## 3.4 Evolution, the Ideal Free Distribution, and Alternative Movement Cues

Orienting movement towards resources (Nathan et al., 2008, ; where to move) can be a chal-362 lenge in a system with low densities of discrete prey items, because the local prey density may 363 provide very limited information about local productivity. In fact, major parts of the resource 364 landscape may be a 'clueless plateau' (Perkins, 1992); on most cells, foragers cannot make di-365 rected movement decisions based on prey-item abundances alone, as all neighbouring items 366 abundances are identical (see white areas in Fig. 5A; A1: scenario 1, A2: scenario 2, A3: sce-367 nario 3). At the beginning of all three scenarios, about 75% of landscape cells have a different 368 number of prey-items from the cells around them; these are primarily the cells with an in-369 termediate r, which have more prey than the lowest productivity cells at the peripheries of resource peaks, but fewer prey than the peaks themselves. This proportion rapidly declines to 371 a much lower value within 10 generations in all three scenarios. 372

The scenarios differ, however, regarding the 'cluelessness' of the landscape on evolutionary timescales (Fig. 5B). In scenario 1, the proportion of cells from which it is possible to move to a neighbouring cell with more items is initially very high (Fig. 5A1). This proportion rapidly declines to  $\sim$ 25% within 10 generations, as foragers deplete most cells on the landscape until they are devoid of prey items, and the majority of the landscape is a clueless plateau. In this context, foragers evolve to move towards handlers — with > 75% of individuals showing a

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preference for handlers within 100 generations (Fig. 5B1 – blue line). Forager preference for handlers (which do not represent a direct resource in scenario 1), may be explained as the sensing of a long-term cue of local productivity. Since handlers are immobilised on the cell where they find a prey item, cells with handlers are (1) certain to have a non-zero rate of prey item generation, and (2) likely to border cells which can generate prey items, due to the autocorrelation (spatial patterning) of the landscape.

Scenario 2 landscapes follow a similar pattern to scenario 1 in the first 10 generations (Fig. 385 5A2). However, within 50 generations, the reduction in prey item extraction allows most cells 386 to regenerate prey items, with differences among cells according to their productivity r (see 387 also Fig. 2A). Thus it is possible to move to a neighbouring cell with more items from > 75% 388 of the landscape (Fig. 5A2 – panel gen: 50, 5B2). However, since the majority of foragers are 389 kleptoparasites, which seek out handlers, this recovery of the resource landscape provides only 390 indirect movement cues, as searching foragers are more likely to convert to handlers on cells 391 with more prey items. Unlike scenario 1, the rapid increase in handler preference is driven 392 by correlational selection on kleptoparasites (see subsection above). Scenario 3 is similar to 393 scenario 2, except that only about half the landscape allows movement to a neighbouring cell 394 with more prey items (Fig. 5A3, 5B3). Here, the rapid evolution of a handler preference in 395 movement decisions cannot be assigned a clear cause, since handlers are both a potential direct 396 resource as well as indirect cues to the location of productive cells. 397

### 3.5 Effect of Landscape Productivity on Evolutionary Outcomes

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The prey-item regrowth rate that characterises the peaks of the resource landscape ( $r_{max}$ ) is a measure of the productivity of the resource landscape overall. Having thus far focused on scenarios with  $r_{max} = 0.01$  (corresponding to a peak production of 4 food times per consumer lifetime), we find that, not unexpectedly, the value of  $r_{max}$  has a marked effect on evolved population activity budgets, mean per capita intake, and even evolved strategies. The frequency of foraging reduces with  $r_{max}$  in scenarios 1 and 3; this is caused by more frequent acquisition of prey items (as regrowth keeps pace with depletion), which results in a greater frequency of handling rather than foraging.

In scenario 2 however, the frequency of handling is relatively unaffected by increasing  $r_{max}$ 

(Fig. 6A). The difference between scenarios 2 and 3 has to do with the change in the frequency 408 of kleptoparasitism (Fig. 6B). In scenario 2, kleptoparasitism forms > 75% of all activities at 409 very low  $r_{max}$ , and is much more common than in scenario 3 populations at the same regrowth 410 rate. However, at relatively high  $r_{max}$  (0.03), the fixed kleptoparasitic strategy goes extinct. 411 This is because at high  $r_{max}$ , the probability of a forager finding prey is much greater than the probability of a kleptoparasite finding a handler, in both initial (< 10) and later generations (> 50). Consequently, kleptoparasites are responsible for only a very small fraction of the total 414 population intake, have relatively much lower fitness, and do not proliferate. Thus at these 415 regrowth rates, a scenario 2 population is nearly identical to a scenario 1 population; while 416 some kleptoparasites may be seen in later generations, these are more likely due to mutation 417 in the forager strategy, rather than longstanding lineages of kleptoparasites.

In scenario 3, kleptoparasitism persists at low frequencies even at the highest regrowth rates (Fig. 6B); thus some foragers lose time in extracting items which are then stolen from them. Consequently, while populations in all three scenarios achieve very similar mean per capita intakes at low  $r_{max}$ , at intermediate regrowth rates (0.01 – 0.025), conditionally kleptoparasitic populations achieve a higher mean per capita intake than populations using fixed strategies. Only at high regrowth rates, when fixed strategy populations (scenario 2) effectively convert to purely forager populations (scenario 1), do they achieve a higher intake than scenario 3 populations (Fig. 6C).

# 4 Discussion

Summary of Results Our spatially-explicit individual-based model implements the ecology and evolution of movement and foraging decisions, as well as resource dynamics, in biologically plausible ways, and offers a new perspective about the distribution of animals in relation to their resources under different scenarios of competition. First, we show that when moving with a limited perception range and competing only by exploitation, individuals evolve movement strategies for both direct and indirect resource cues (prey items and handlers, respectively). Regardless, on a resource landscape with discrete prey items, large areas may become devoid of any movement cues, leading to a mismatch between individual distribu-

tion, prey item distribution, and landscape productivity. Second, we show that when inter-436 ference competition in the form of kleptoparasitism is allowed as a fixed strategy, it rapidly 437 establishes itself on landscapes where stealing is more time-efficient than searching for prey. 438 This rapid increase in kleptoparasitism as a strategy is accompanied by the evolution of move-439 ment strategies that favour moving towards handlers, which are the primary resource of the kleptoparasites. In this sense, obligate kleptoparasites may be thought of as forming a higher trophic level, with any handling consumers as their prey. Third, we show that when foraging 442 strategy is allowed to be conditional on local cues, (1) the population's mean per capita in-443 take is significantly higher than that of a population with fixed strategies, and (2) unlike fixed 444 strategy populations, kleptoparasitism as a strategy does not go extinct on high-productivity 445 landscapes. However, across scenarios, individuals are broadly unable to match the productivity of the resource landscape, contrary to the predictions of IFD based models, which predict input matching for some (Parker and Sutherland, 1986; Holmgren, 1995; Hamilton, 2002), or 448 all of the competitive types Korona (1989). 449

**Comparison with Previous Models** Existing models of competition and movement impose fixed movement rules on individuals to mimic either ideal or non-ideal individuals (Vickery et al., 1991; Cressman and Křivan, 2006; Amano et al., 2006; Beauchamp, 2008; Stillman and 452 Goss-Custard, 2010; White et al., 2018). When individual competitive strategies are included, 453 models consider them to represent differences in competitive ability (e.g. Parker and Suther-454 land, 1986; Holmgren, 1995; Hamilton, 2002), or as a probabilistic switch between producing 455 and scrounging (Beauchamp, 2008). In contrast, our model allows individuals' movement (and competition) decisions to be adaptive responses to local environmental cues. Similar to Getz 457 et al. (2015, 2016) and White et al. (2018), our individuals choose from among the available 458 movement options after weighing the local environmental cues; this is analogous to the move-459 ment ecology concepts of resource or step selection functions (Fortin et al., 2005; Manly et al., 460 2007; White et al., 2018). The local environmental cues in our model are constantly changing, 461 as we model discrete, depletable prey items on the resource landscape, contrasting with many IFD models (Tregenza, 1995; Amano et al., 2006). This allows for a more plausible, fine-scale 463 consideration of exploitation competition, which is often neglected, and allows the cues sensed 464

by individuals to strongly structure the distribution of competitors (see below).

Adaptive responses must have an explicit evolutionary context, and consider multiple gen-466 erations of the population. We follow Beauchamp (2008) and Getz et al. (2015) in allowing 467 the decision making weights for movement, and variation thereof, to be the outcomes of nat-468 ural selection. However, instead of using 'evolutionary algorithms' (Beauchamp, 2008; Getz 469 et al., 2015, 2016) to 'optimise' individual movement rules, we consider a more plausible evolutionary process: Instead of allowing the fittest 50% of the population to replicate, the number 471 of offspring are proportional to individual fitness (Hofbauer and Sigmund, 2003). The weight 472 loci are subject to mutations independently, rather than subjecting all loci of an individual to si-473 multaneous mutation. Finally, we avoided the unrealistic assumption of 'simulated annealing', 474 which adapts the mutation rate or the mutational step sizes to the rate of evolutionary change. Instead we drew mutation sizes from a Cauchy distribution, so that most mutations are very 476 small, but large-effect mutations do occur throughout the simulation. Similarly, rather than 477 determining foraging style (searching or stealing) probabilistically or ideally (Vickery et al., 478 1991; Beauchamp, 2008; Tania et al., 2012), our individuals' foraging decisions are also shaped 479 by selection (in scenarios 2 and 3). 480

Movement Rules on Depleted Landscapes In our scenario 1, depletion of discrete prey can leave many areas empty of prey items: in such areas, movement informed by a resource gradient is impossible, and individuals move randomly ('clueless regions': Perkins, 1992). Our work suggests that this lack of movement cues and the resulting random movement might be among the mechanisms by which unsuitable 'matrix' habitats modify animal movement on heterogeneous landscapes (Kuefler et al., 2010).

When individuals do not sense resource gradients, the presence of successful consumers

(handlers) may indicate a suitable foraging spot (local enhancement; see Giraldeau and Beauchamp,

1999; Beauchamp, 2008; Cortés-Avizanda et al., 2014), or direction of movement (Guttal and

Couzin, 2010). The presence of unsuccessful individuals, meanwhile, may signal potential

costs from exploitation or interference competition. This selects for movement strategies in
corporating the presence and condition of competitors into individual movement decisions

('social information': Danchin et al., 2004; Dall et al., 2005). While external costs such as pre-

dation have been invoked to explain consumer aggregation (Krause and Ruxton, 2002; Folmer et al., 2012), information on the location of productive areas alone may be a sufficient benefit to promote the evolution of social information use, and consequently aggregation (Folmer et al., 2010; Cortés-Avizanda et al., 2014).

While individuals broadly prefer prey and handlers, and avoid non-handlers, there is substantial individual variation in the strength of each weight within populations, as expected
from heterogeneous landscapes (see Supplementary Material; see Wolf and Weissing 2010 for
background). The persistence of multiple 'movement morphs' (see Getz et al., 2015) across
multiple generations indicates that they are alternative movement strategies of equal fitness.

Indeed, polymorphism in movement rules may help reduce competition as individuals make
subtly different movement decisions when presented with the same cues (see Wolf and Weissing, 2012, for a review; see also Laskowski and Bell 2013).

Movement Rules and Competitive Interactions IFD models predict that individual movement should result in 'input matching': consumer distributions should track the profitability of resource patches (Fretwell and Lucas, 1970; Parker, 1978), and dominant competitive types (including kleptoparasites) should monopolise the best patches (Parker and Sutherland, 1986; Holmgren, 1995; Hamilton, 2002, but see Korona 1989). In our scenarios 2 and 3, individuals seeking to steal prey from competitors unsurprisingly and rapidly evolve to track handlers (a direct resource), while avoiding non-handlers (potential competitors). However, these evolved rules do not lead kleptoparasites to occupy the best cells as predicted (Parker and Sutherland, 1986; Holmgren, 1995; Hamilton, 2002). Across our scenarios (including sc. 1), individual density is only weakly correlated with cell productivity.

In scenario 2, this departure from predictions is driven by the contrasting movement rules of foragers, which evolve to *avoid* handlers as well as non-handlers, both of which might be kleptoparasites (Folmer et al. 2010; Bijleveld et al. 2012; see Supplementary Material). Thus, foragers (the subordinate competitive type) undermatch the resource landscape, as resource peaks are more likely to have handlers (Parker and Sutherland, 1986; Holmgren, 1995; Hamilton, 2002). Fixed kleptoparasites cannot extract prey themselves, and must move off resource peaks to track and rob handlers (similar to Parker and Sutherland, 1986), breaking the link

between individual density and productivity.

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This shows the pitfalls of simplistically linking current ecological conditions with evolved 524 behavioural responses (and emergent distributions), and shows why understanding the evo-525 lutionary history of competitive interactions is key. In this vein, a hypothetical experiment 526 would wrongly expect that foragers from resource-rich landscapes (sc. 2) better tolerate com-527 petitors due to less exploitation competition, while foragers from resource scarce landscapes 528 (sc. 1) might be competitor-avoidant. Furthermore, ecological conditions may not be a reliable 529 guide to the presence of individual variation, which is often expected as an outcome of en-530 hanced competition. Indeed, scenario 2 shows significant within-strategy individual variation 531 in movement weights, which might help reduce within-strategy exploitation competition, or 532 help foragers better avoid kleptoparasites (Wolf and Weissing, 2012; Laskowski and Bell, 2013). Interestingly, scenario 3 has the least individual variation in movement rules, potentially be-534 cause behavioural plasticity in foraging strategy dampens individual variation (Pfennig et al., 535 2010). Here, the non-handler avoidance shows the most morphs, but it is unclear whether this 536 variation is linked to the frequency with which individuals use either foraging strategy — po-537 tentially leading to subtle, emergent behavioural differences that are conditioned on the local 538 environment (Wolf and Weissing, 2010, 2012).

Ecological Constraints and Evolved Foraging Strategies Foraging strategies in which animals specialise on a resource type are expected to be constrained by the availability of that resource; thus kleptoparasitism should constrained by the density of targets (Spencer et al., 2017; Spencer and Broom, 2018). In our scenarios 2 and 3, we would expect to see more kleptoparasitism with increasing  $r_{max}$ , as prey and consequently, handlers, are more abundant. We find instead that kleptoparasitism declines with increasing  $r_{max}$ , in line with Emlen (1966), who predicted that the commoner food type (prey) rather than the more efficiently exploited one (handlers) should be preferred.

This effect is especially stark in scenario 2, where kleptoparasites go extinct when prey are very common at high  $r_{max}$ . At stable population densities (due to external constraints such as nest-site availability), the persistence of fixed kleptoparasitism depends on their intake *relative* to foragers. Since intake is an outcome of movement rules, and population movement rules are

not well adapted to their environment in early generations, foragers obtain, as a clade, more 552 intake than kleptoparasites. Modelling discrete prey items and individuals in a spatial context, 553 then, leads to the finding that obligate kleptoparasitism is only a viable strategy when the 554 probability of encountering prey is much less than the probability of encountering handlers. 555 While few taxa show such a marked foraging specialisation within populations, this might 556 explain why kleptoparasitism is frequent among seabirds, whose communal roosts are much 557 more easily targeted than unpredictable shoals of fish out at sea (Brockmann and Barnard, 558 1979). 559 Finally, comparing across regrowth rates shows why predicting the long-term effect of envi-560 ronmental change on populations must keep unanticipated behavioural complexity in mind. 561 While both scenario 1 and 2 populations appear identical at high  $r_{max}$ , even a small decrease 562 in environmental productivity could lead to an abrupt drop in per-capita intake — and po-563 tentially, strongly reduced growth or survival — for fixed strategy populations due to (unex-564

# 566 5 Acknowledgments

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pected) emergent kleptoparasitism.

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## References

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Amano, T., K. Ushiyama, S. Moriguchi, G. Fujita, and H. Higuchi. 2006. Decision-Making in Group Foragers with Incomplete Information: Test of Individual-Based Model in Geese. Ecological Monographs 76:601–616.

Baldauf, S. A., L. Engqvist, and F. J. Weissing. 2014. Diversifying evolution of competitiveness.
Nature Communications 5:5233.

- Barnard, C. J., and R. M. Sibly. 1981. Producers and scroungers: A general model and its application to captive flocks of house sparrows. Animal behaviour 29:543–550.
- Beauchamp, G. 2008. A spatial model of producing and scrounging. Animal Behaviour 76:1935–1942.
- Bijleveld, A. I., E. O. Folmer, and T. Piersma. 2012. Experimental evidence for cryptic interfer ence among socially foraging shorebirds. Behavioral Ecology 23:806–814.
- Birch, L. C. 1957. The Meanings of Competition. The American Naturalist 91:5–18.
- Brockmann, H., and C. Barnard. 1979. Kleptoparasitism in birds. Animal Behaviour 27:487–589 514.
- Case, T. J., and M. E. Gilpin. 1974. Interference Competition and Niche Theory. Proceedings of
   the National Academy of Sciences 71:3073–3077.
- Clutton-Brock, T., and B. C. Sheldon. 2010. Individuals and populations: The role of long-term,
   individual-based studies of animals in ecology and evolutionary biology. Trends in Ecology
   & Evolution 25:562–573.
- Cortés-Avizanda, A., R. Jovani, J. A. Donázar, and V. Grimm. 2014. Bird sky networks: How do avian scavengers use social information to find carrion? Ecology 95:1799–1808.
- Cressman, R., and V. Křivan. 2006. Migration Dynamics for the Ideal Free Distribution. The
  American Naturalist 168:384–397.
- Dall, S. R. X., L.-A. Giraldeau, O. Olsson, J. M. McNamara, and D. W. Stephens. 2005. Information and its use by animals in evolutionary ecology. Trends in Ecology & Evolution
   20:187–193.
- Danchin, É., L.-A. Giraldeau, T. J. Valone, and R. H. Wagner. 2004. Public Information: From Nosy Neighbors to Cultural Evolution. Science 305:487–491.
- 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.
- DeAngelis, D. L., and S. G. Diaz. 2019. Decision-Making in Agent-Based Modeling: A Current
   Review and Future Prospectus. Frontiers in Ecology and Evolution 6.
- Duckworth, R. A., and A. V. Badyaev. 2007. Coupling of dispersal and aggression facilitates the rapid range expansion of a passerine bird. Proceedings of the National Academy of Sciences of the United States of America 104:15017–22.
- Emlen, J. M. 1966. The Role of Time and Energy in Food Preference. The American Naturalist 100:611–617.

- Folmer, E. O., H. Olff, and T. Piersma. 2010. How well do food distributions predict spatial distributions of shorebirds with different degrees of self-organization? Journal of Animal Ecology 79:747–756.
- 2012. The spatial distribution of flocking foragers: Disentangling the effects of food
   availability, interference and conspecific attraction by means of spatial autoregressive modeling. Oikos 121:551–561.
- Fortin, D., H. L. Beyer, M. S. Boyce, D. W. Smith, T. Duchesne, and J. S. Mao. 2005. Wolves Influence Elk Movements: Behavior Shapes a Trophic Cascade in Yellowstone National Park. Ecology 86:1320–1330.
- Fretwell, S. D., and H. L. Lucas. 1970. On territorial behavior and other factors influencing habitat distribution in birds. Acta Biotheoretica 19:16–36.
- Garay, J., R. Cressman, F. Xu, M. Broom, V. Csiszár, and T. F. Móri. 2020. When optimal foragers
   meet in a game theoretical conflict: A model of kleptoparasitism. Journal of Theoretical
   Biology 502:110306.
- Getz, W. M., R. Salter, A. J. Lyons, and N. Sippl-Swezey. 2015. Panmictic and Clonal Evolution on a Single Patchy Resource Produces Polymorphic Foraging Guilds. PLOS ONE 10:e0133732–e0133732.
- Getz, W. M., R. Salter, D. P. Seidel, and P. van Hooft. 2016. Sympatric speciation in structureless
   environments. BMC Evolutionary Biology 16:50–50.
- Giraldeau, L.-A., and G. Beauchamp. 1999. Food exploitation: Searching for the optimal joining
   policy. Trends in Ecology & Evolution 14:102–106.
- Goss-Custard, J. D. 1980. Competition for food and interference among waders. Ardea 55:31–53.
- Guttal, V., and I. D. Couzin. 2010. Social interactions, information use, and the evolution of
   collective migration. Proceedings of the National Academy of Sciences 107:16172.
- Hamilton, I. M. 2002. Kleptoparasitism and the distribution of unequal competitors. Behav ioral Ecology 13:260–267.
- Hofbauer, J., and K. Sigmund. 2003. Evolutionary game dynamics. Bulletin of the American
   mathematical society 40:479–519.
- Holmgren, N. 1995. The Ideal Free Distribution of Unequal Competitors: Predictions from a
   Behaviour-Based Functional Response. Journal of Animal Ecology 64:197–212.
- Houston, A. I. 2008. Matching and ideal free distributions. Oikos 117:978–983.
- Huston, M., D. DeAngelis, and W. Post. 1988. New Computer Models Unify Ecological The oryComputer simulations show that many ecological patterns can be explained by interactions among individual organisms. BioScience 38:682–691.
- Iyengar, E. V. 2008. Kleptoparasitic interactions throughout the animal kingdom and a re evaluation, based on participant mobility, of the conditions promoting the evolution of klep toparasitism. Biological Journal of the Linnean Society 93:745–762.

- Keddy, P. A. 2001. Studying competition. Pages 1–59 in P. A. Keddy, ed. Competition, Popula tion and Community Biology Series. Springer Netherlands, Dordrecht.
- Korona, R. 1989. Ideal free distribution of unequal competitors can be determined by the form of competition. Journal of Theoretical Biology 138:347–352.
- 659 Krause, J., and G. D. Ruxton. 2002. Living in Groups. Oxford University Press.
- 660 Krebs, J., and N. Davies. 1978. Behavioural Ecology: An Evolutionary Approach.
- Kuefler, D., B. Hudgens, N. M. Haddad, W. F. Morris, and N. Thurgate. 2010. The conflicting
   role of matrix habitats as conduits and barriers for dispersal. Ecology 91:944–950.
- Kuijper, B., I. Pen, and F. J. Weissing. 2012. A guide to sexual selection theory. Annual Review
   of Ecology, Evolution, and Systematics 43:287–311.
- Laskowski, K. L., and A. M. Bell. 2013. Competition avoidance drives individual differences in response to a changing food resource in sticklebacks. Ecology Letters 16:746–753.
- Long, X., and F. J. Weissing. 2020. Individual variation in parental care drives divergence of sex roles. bioRxiv page 2020.10.18.344218.
- Manly, B., L. McDonald, D. L. Thomas, T. L. McDonald, and W. P. Erickson. 2007. Resource
   Selection by Animals: Statistical Design and Analysis for Field Studies. Springer Science &
   Business Media.
- Nathan, R., W. M. Getz, E. Revilla, M. Holyoak, R. Kadmon, D. Saltz, and P. E. Smouse. 2008.

  A movement ecology paradigm for unifying organismal movement research. Proceedings of the National Academy of Sciences 105:19052–19059.
- Netz, C., H. Hildenbrandt, and F. J. Weissing. 2020. Complex eco-evolutionary dynamics induced by the coevolution of predator-prey movement strategies. bioRxiv page 2020.12.14.422657.
- Parker, G. 1978. Searching for mates.[In: Behavioural ecology. An evolutionary approach. JR Krebs and NB Davies, eds] .
- Parker, G. A., and W. J. Sutherland. 1986. Ideal free distributions when individuals differ in competitive ability: Phenotype-limited ideal free models. Animal Behaviour 34:1222–1242.
- Perkins, D. N. 1992. Topography of Invention. Page 238 in Inventive Minds: Creativity in
   Technology, vol. 10. Oxford University Press, New York, NY, USA.
- Pfennig, D. W., M. A. Wund, E. C. Snell-Rood, T. Cruickshank, C. D. Schlichting, and A. P.
   Moczek. 2010. Phenotypic plasticity's impacts on diversification and speciation. Trends in
   Ecology & Evolution 25:459–467.
- Rutten, A. L., K. Oosterbeek, J. van der Meer, S. Verhulst, and B. J. Ens. 2010a. Experimental evidence for interference competition in oystercatchers, Haematopus ostralegus. I. Captive birds. Behavioral Ecology 21:1251–1260.
- Rutten, A. L., K. Oosterbeek, S. Verhulst, N. J. Dingemanse, and B. J. Ens. 2010b. Experimental evidence for interference competition in oystercatchers, Haematopus ostralegus. II. Free-living birds. Behavioral Ecology 21:1261–1270.

- Schlägel, U. E., V. Grimm, N. Blaum, P. Colangeli, M. Dammhahn, J. A. Eccard, S. L. Hausmann, A. Herde, H. Hofer, J. Joshi, S. Kramer-Schadt, M. Litwin, S. D. Lozada-Gobilard,
- M. E. H. Müller, T. Müller, R. Nathan, J. S. Petermann, K. Pirhofer-Walzl, V. Radchuk, M. C.
- Rillig, M. Roeleke, M. Schäfer, C. Scherer, G. Schiro, C. Scholz, L. Teckentrup, R. Tiedemann,
- W. Ullmann, C. C. Voigt, G. Weithoff, and F. Jeltsch. 2020. Movement-mediated community assembly and coexistence. Biological Reviews.
- Spencer, R., and M. Broom. 2018. A game-theoretical model of kleptoparasitic behavior in an urban gull (Laridae) population. Behavioral Ecology 29:60–78.
- Spencer, R., Y. I. Russell, B. J. A. Dickins, and T. E. Dickins. 2017. Kleptoparasitism in gulls Laridae at an urban and a coastal foraging environment: An assessment of ecological predictors.
   Bird Study 64:12–19.
- Stillman, R. A., and J. D. Goss-Custard. 2010. Individual-based ecology of coastal birds. Bio logical Reviews 85:413–434.
- Tania, N., B. Vanderlei, J. P. Heath, and L. Edelstein-Keshet. 2012. Role of social interactions in
   dynamic patterns of resource patches and forager aggregation. Proceedings of the National
   Academy of Sciences 109:11228–11233.
- Tregenza, T. 1995. Building on the Ideal Free Distribution. Pages 253–307 *in* Advances in Ecological Research, vol. 26. Elsevier.
- 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, K. Meijer, T. Piersma, and F. J. Weissing. 2007. Interference
   competition, the spatial distribution of food and free-living foragers. Animal Behaviour
   74:1493–1503.
- 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.
- van der Meer, J., 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.
- Vickery, W. L., L.-A. Giraldeau, J. J. Templeton, D. L. Kramer, and C. A. Chapman. 1991. Producers, Scroungers, and Group Foraging. The American Naturalist 137:847–863.
- White, L. A., J. D. Forester, and M. E. Craft. 2018. Disease outbreak thresholds emerge from interactions between movement behavior, landscape structure, and epidemiology. Proceedings of the National Academy of Sciences 115:7374–7379.
- Wolf, M., and F. J. Weissing. 2010. An explanatory framework for adaptive personality differences. Philosophical Transactions of the Royal Society B: Biological Sciences 365:3959–3968.

# 731 6 Figure legends

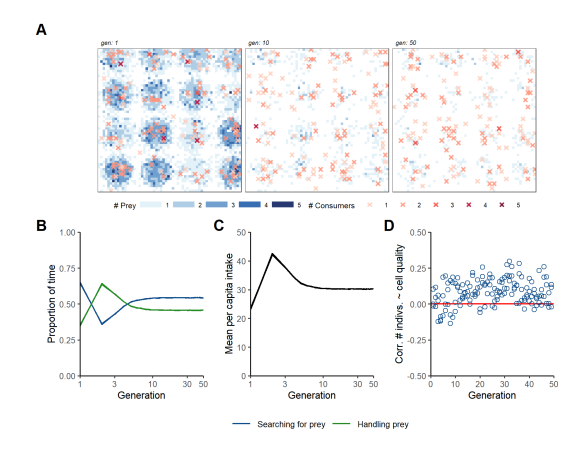


Figure 1: Eco-evolutionary implications of pure exploitation competition (scenario 1). (A) When a population is comprised solely of foragers seeking prey on a resource landscape, the initially well-stocked resource landscape is swiftly depleted within 10 generations (out of 1,000 simulated). This sparsity in prey-item abundance is maintained throughout the remaining generations of the simulation. Individuals, whose local density is shown by coloured crosses, are scattered over the landscape. These dynamics are explained by the fact that (B) within 20 generations of evolution, the population reaches an equilibrium in the relative proportion of time spent on searching prey and handling prey, and in (C) the total intake of the population. (D) In a departure from the intake matching rule of IFD theory, cell occupancy (number of foragers per cell) is only weakly correlated with cell productivity r. Panel A shows a single replicate, while panels B, C and D show three replicate simulations (lines overlap almost perfectly); all panels are for  $r_{max} = 0.01$ . NB: Both B, C show a log-scaled X axis to more clearly show dynamics in early generations.

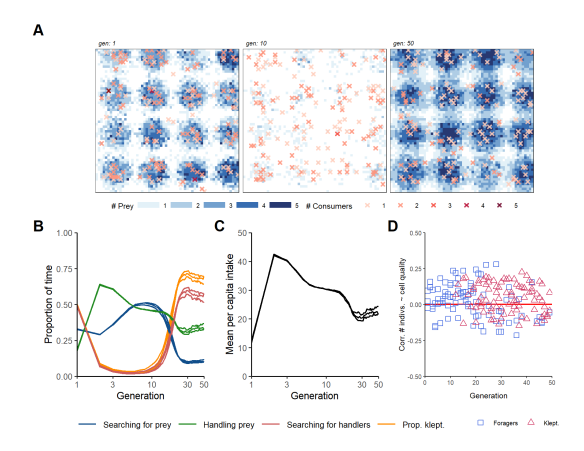


Figure 2: Eco-evolutionary implications of the coexistence of foragers and kleptoparasites (scenario 2). In populations with both foragers and kleptoparasites, (A) the initially wellstocked resource landscape is drastically depleted by generation 10; however, prey densities recover strongly by generation 50, even beyond the densities in generation 1. The local density of individuals on occupied cells is shown as coloured crosses. (B) An equilibrium between the strategies is reached within 30 generations, with the relative frequency of kleptoparasites (orange line) first dropping to very low levels but later recovering to reach a high level ( $\sim 70\%$ ) in all three replicates. The activity budget parallels the relative frequency of kleptoparasites, and at equilibrium, about 10% of the individuals are foragers searching for prey, 50% are kleptoparasites searching for handlers, and 40% are handlers (either foragers or kleptoparasites). (C) In early generations, when kleptoparasites are rare, the population intake rate exhibits the same pattern as in Fig. 1B, dropping to a lower level with the emergence of kleptoparasites. This is accompanied by an increase in the proportion of time spent on stealing attempts (red line − B), and a corresponding decrease in prey seeking (by searching foragers; blue line − B), and handling (green line – C). (D) Cell occupancy (local density of foragers per cell) is only weakly correlated with cell productivity r, dropping to zero at equilibrium. Panel A shows a single replicate, while **B**, **C** and **D** show three replicates; all panels are for  $r_{max} = 0.01$ .

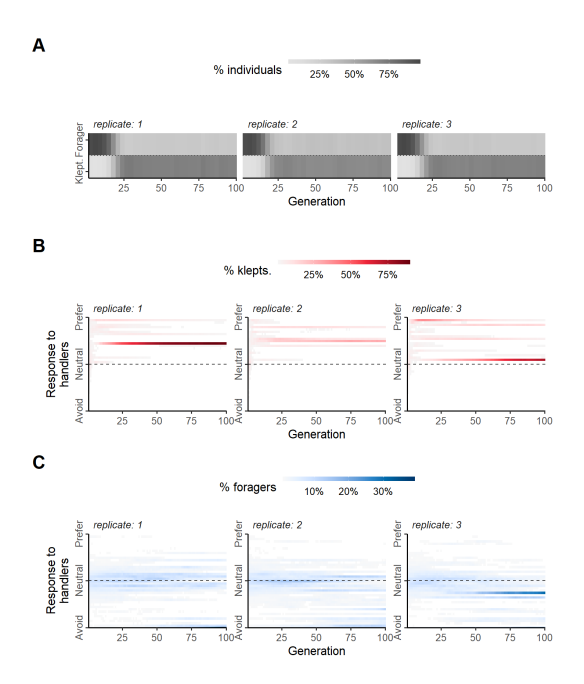
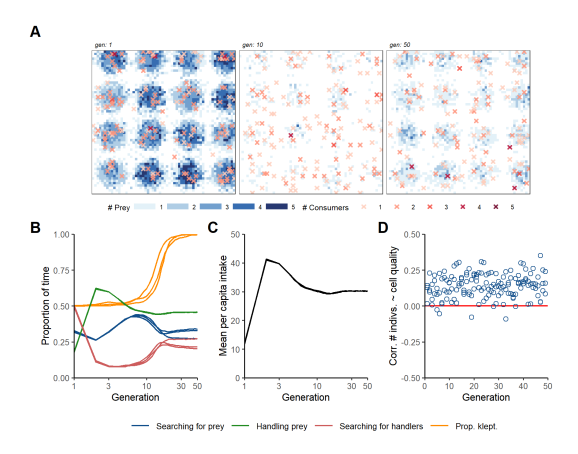
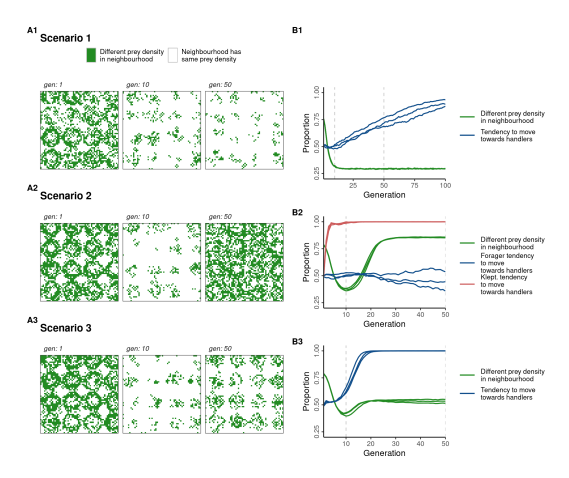


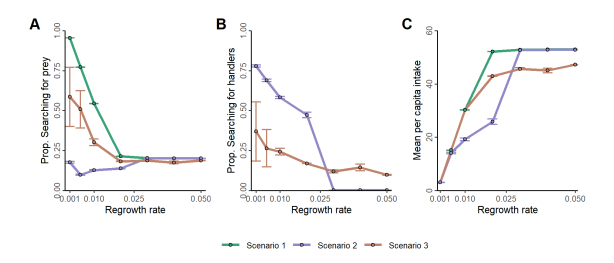
Figure 3: Divergence of movement strategies between foragers and kleptoparasites (scenario 2). (A) Kleptoparasitism rapidly becomes the more frequent strategy in scenario 2 populations for the parameters considered, with no differences across replicates. However, replicates differ considerably in the evolved movement strategies. This is illustrated by the distribution of the weighing factor  $s_H$  (describing the effect of local handler density on the movement decision) in kleptoparasites (B) and foragers (C), respectively. In kleptoparasites, the weights  $s_H$  are generally positive, indicating that kleptoparasites are attracted by handlers. However, different  $s_H$  values stably coexist, indicating that kleptoparasites are polymorphic in their movement strategy. Foragers are also polymorphic in their handler responses: foragers attracted by handlers (positive  $s_H$ ) coexist with foragers repelled by handlers (negative  $s_H$ ). All panels show three replicates at  $r_{max} = 0.01$ .



**Figure 4: Eco-evolutionary implications of conditional foraging strategies (scenario 3). (A)** The initially well-stocked resource landscape is rapidly depleted within 10 generations, yet within 50 generations, prey abundances recover on many cells, though not to the extent of scenario 2. The local density of individuals on occupied cells is shown as coloured crosses. **(B)** By generation 30, all individuals encountering handlers will choose to steal prey rather than search for prey themselves. The proportion of time spent searching (blue line), handling (green line), and stealing prey (red line) also reach an equilibrium that differs somewhat across replicates. **(C)** Yet, the total intake of the population reaches the same equilibrium value in all three replicates. **(D)** The correlation between the local density of individuals on a cell, and its productivity r is stronger than in scenario 2. Panel **A** shows a single replicate, while **B**, **C** and **D** show three replicates; all panels are for  $r_{max} = 0.01$ .



**Figure 5: Uninformative prey densities and the evolution of alternative movement cues. (A1, A2, A3)** On cells coloured green, local prey densities are informative for movement, as the central and neighbouring cells have different prey densities. While differences in local prey densities provide informative cues for 'adaptive' movement in early generations, this is much less true once the resource landscape is depleted of prey items (depending on the scenario). **(B1, B2, B3)** The proportion of cells where differences in local prey densities provide informative movement cues (green line), and the proportion of individuals preferring to move towards handlers (blue line), whose presence may be used as an alternative cue for movement towards higher-productivity areas of the landscape. In **(B2)** representing scenario 2, this proportion is shown separately for foragers (blue line) and kleptoparasites (red line). While panels in **(A)** show a single representative replicate for  $r_{max} = 0.01$ , panels in **(B)** show three replicates.



**Figure 6: Landscape productivity strongly affects scenario outcomes. (A)** The proportion of time spent searching for food decreases with increasing  $r_{max}$  in scenarios 1 and 3 but remains relatively stable within scenarios. This is partly due to a higher proportion of time spent handling at higher prey densities. **(B)** The proportion of time spent searching for handlers (in order to steal prey from them) also decreases with increasing  $r_{max}$ . In scenario 2, kleptoparasites go extinct for  $r_{max}$  values above 0.025. **(C)** At low productivity, the average intake is similar in all three scenarios. For higher  $r_{max}$  values the average intake rate is lowest in scenario, until  $r_{max}$  is larger than 0.025 and kleptoparasites go extinct (leading to the same kind of population as in scenario 1). At high  $r_{max}$ , the average intake rate in populations with conditional kleptoparasites (scenario 3) is substantially lower than in populations without kleptoparasitism.