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# Spatial memory predicts home range size and predation risk in pheasants

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1	Spatial memory predicts home range size and predation risk in pheasants
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#### 23 Abstract

24 Most animals confine their activities to a discrete home range, long assumed to reflect the fitness 25 benefits of obtaining spatial knowledge about the landscape. However, few empirical studies have 26 linked spatial memory to home range development, or determined how selection operates on spatial 27 memory via the latter's role in mediating space use. We assayed the cognitive ability of juvenile 28 pheasants (Phasianus colchicus) reared under identical conditions before releasing them into a novel 29 landscape and used high-throughput tracking to record their movements as they developed their home 30 ranges, and determine the location, timing and cause of any mortality events. Individuals with greater 31 spatial reference memory developed larger home ranges. Mortality risk from predators was highest at 32 the periphery of an individual's home range, in areas they had less experience and opportunity to 33 obtain spatial information. Importantly, predation risk was lower in individuals with greater spatial 34 memory and larger core home ranges, suggesting selection may operate on spatial memory by 35 increasing the ability to learn about predation risk across the landscape. Our results reveal that spatial 36 memory, determined from abstract cognitive assays, shapes home range development and variation, 37 and suggests predation risk selects for spatial memory via experience-dependent spatial variation in 38 mortality.

39

#### 40 Main text

## 41 Introduction

42 Most animals confine the majority of their activities to a relatively restricted spatial region: their 43 home range <sup>1-4</sup>. The patterns and dynamics of animal home ranges have fundamental consequences 44 across biology, from epidemiology <sup>5,6</sup> to population dynamics and predator-prey interactions <sup>6–9</sup>, yet 45 surprisingly little work demonstrates the processes by which selection can emerge and shape home 46 range variation. A long-standing explanation for why many animals form home ranges, supported by 47 a wealth of mechanistic modelling studies (e.g. <sup>1,10</sup>), is that it allows a resident to learn the location of, 48 and optimally exploit, patchily distributed resources such as food and shelter <sup>11–13</sup>. Consequently, an 49 animal's ability to learn, retain and update information on the spatial location of resources within its home range is assumed to be of great benefit <sup>12,14</sup>, and is a central tenet to many definitions of the 50 home range <sup>15,16</sup>. Decades of work incorporating proxies of resource-based spatial memory into 51 52 mechanistic random-walk models of home range development results in the emergence of more 53 realistic home ranges, or better statistical power in predicting the development of real animal home ranges, compared to alternative processes such as sensory detection of resources <sup>4,17–27</sup>. Whilst these 54 55 prior studies have produced convincing conclusions when incorporating artificial analogues of spatial 56 memory, they still require empirical validation in light of the extensive work from comparative and 57 classical psychology demonstrating animals often make use of multiple and distinct cognitive modalities that work synergistically to process different types of information during daily life <sup>28</sup>. 58 59 Indeed, the few studies that incorporate different types of spatial memory such as 'working memory' 60 (short-term memory of recently visited locations<sup>29</sup>) and 'reference memory' (longer-term memory of landmarks<sup>30</sup>) suggest they work in concert to maximise the efficiency of movement patterns within a 61 home range (e.g. <sup>21,22</sup>), and that working memory itself may be a particularly important determinant of 62 home range size <sup>21</sup>. However, we still lack empirical studies that test the predictions made by these 63 64 simulation studies to establish the role of cognitive ability (and specifically various types of spatial 65 memory) in real home range formation and size variation.

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67 Whilst a great deal of work has focused on the drivers of interspecific variation in home range size, for instance showing that it scales as a function of body size, metabolic rate and forage type <sup>31</sup> as well 68 as being heavily influenced by locomotion and foraging strategies <sup>32–34</sup>, less is known about the causes 69 and consequences of home range size at the intraspecific level, particularly with regards to how these 70 71 are affected by individual-level variation in different traits. Larger home ranges may infer a range of benefits such as increased access to mates, refuges and resources <sup>35</sup>, although the factors constraining 72 73 size are less well established empirically, particularly in species that do not suffer economic costs of aggressively defending spatially distributed and non-divisible resources <sup>22,36,37</sup>. Spatial memory 74 75 ability, as a major factor enabling animals to form stable home ranges in the first place, could be a

76 potential mechanistic component limiting home range size. If animals form home ranges to exploit 77 spatial knowledge about important resources or local variation in predation risk, then all else being 78 equal, better spatial memory should translate to animals being able to utilise larger home ranges 79 without suffering associated costs caused by poor knowledge about the landscape such as inefficient travel between resources <sup>21,22</sup> or suboptimal vigilance whilst travelling across regions of varying 80 predation risk <sup>38</sup>. Whilst some biologically-plausible mechanistic models incorporating multiple 81 82 cognitive modalities have indeed suggested that better spatial memory can translate into the emergence of larger home ranges <sup>21,22</sup>, this needs validation from field-based studies of real animals. 83

84

85 Here, we use a unique system in common pheasants (Phasianus colchicus) where we can rear large 86 numbers of individuals under standardised conditions, assay them across a battery of cognitive tasks, 87 tag all individuals with high spatial and temporal resolution radio tags, and release them into a novel 88 wild landscape to track their movements as they develop their home ranges. This approach reduces 89 variation in early life environmental effects on subsequent behavioural traits, allowing us to test key 90 predictions about the causes and consequences of movement ecology (although, as with any 91 experimental control, this comes at the potential expense of reducing ecological realism). Specifically, 92 we aimed to test how individual-level variation in three cognitive modalities (associative learning, 93 spatial working memory, and development of a spatial reference memory) obtained from 94 psychometric assays predict variation in the development of home range size.

95

96 The degree to which individuals can learn and process spatial information, particularly in relation to 97 resources, should assumedly affect fitness, with potential consequences on selection depending on the 98 heritability of these traits <sup>39</sup>. Indeed, a limited number of studies have uncovered fitness benefits 99 associated with variation in cognitive ability <sup>40,41</sup>, and some have suggested that spatially-explicit 100 processes are critical in driving the association <sup>42,43</sup>. Nevertheless, the pathway by which spatial 101 memory and other cognitive modalities drive variation in the movement of individuals, and how this

102 in turn affects how individuals interact with their environment to ultimately generate selection on 103 spatial memory, is still unclear. Whilst most studies addressing the formation and benefits of home 104 ranges have focused on the importance of exploiting patchily distributed resources (or restrictions in movement caused by aggressive defence by territorial neighbours <sup>7,10,27</sup>), another critical factor that 105 106 shapes natural selection and varies non-randomly across the landscape, is predation. Many predators, 107 particularly ambush hunters, display fine-scale temporal and spatial patterning in their hunting behaviour <sup>44</sup>, which prey can potentially learn and mitigate through strategies such as increased 108 vigilance, avoidance of high risk areas, or exploiting memorised escape routes <sup>38,44-47</sup>. As such, there 109 110 is growing evidence that predation risk may be particularly high in regions that prey are less familiar <sup>44,48,49</sup>. An intuitively appealing, but so far untested, pathway by which selection may operate on 111 spatial memory ability is therefore via variation in predation risk mediated by an individual's 112 113 familiarity (i.e. degree of prior experience) with high-risk areas as determined by the location and 114 structure of its home range.

115

116 The high spatial and temporal resolution of our tracking data allowed us to identify the precise 117 location and timing of predation events during our study (n=45), as well as identify the predator 118 species responsible (all caused by red foxes (Vulpes vulpes), an ambush hunter). These death locations 119 were non-randomly distributed across the landscape, constituting regions of high predation risk. By 120 coupling cognition, movement, and predation data, we were therefore able to test whether individuals 121 are more likely to be killed outside their core home range in regions they have less knowledge 122 regarding predation risk, the degree to which this is predicted by different cognitive abilities, and 123 whether this 'experience-dependent' predation risk has the potential to generate selection on cognition 124 and home range variation. A likely consequence of the hypothesised relationship between spatial 125 memory and home range size we detail above is that predation will also be higher in less familiar parts of the landscape (as found previously in deer <sup>44,49</sup>), a phenomenon we predict to be particularly 126 127 exaggerated in individuals with poor spatial memory since these would be less capable of memorising 128 the locations of non-lethal predator encounters and thus optimising antipredator behaviours across

varying regions of predation risk (e.g. <sup>38</sup>). Whilst prior work on the antipredator benefit of different
cognitive modalities is scarce, we predict that all three types of cognition we measured can affect
predation risk; reference memory for learning landscape features and specific locations associated
with regions of high risk; working memory in memorising and exploiting the locations of recently
visited refuges and escape routes when predators are encountered; and general associative memory
via the learning of non-spatial components associated with predation (e.g. heterospecific alarm calls <sup>50</sup>
and search images of camouflaged predators <sup>51</sup>).

136

We hatched and reared 126 pheasants and assayed their performance on three well-established 137 138 cognitive tasks between the ages of four and seven weeks old. The first cognitive assay, a binary 139 colour discrimination task, tested the ability of individuals to associate a particular colour with a food 140 reward, a proxy for general associative learning ability (i.e. the ability to learn simple associations between stimuli<sup>52</sup>), and was explicitly designed to not be solved using spatial cues (Fig. 1C). The 141 second, a radial arm maze task, was intended to test an individual's ability to use spatial cues to orient 142 around a maze and optimally exploit multiple food resources as they were depleted, a common 143 paradigm for testing spatial working memory <sup>53</sup> and one that is particularly ecologically relevant to 144 species such as pheasants that feed on patchily-distributed resources (e.g. <sup>54</sup>). The third, a task similar 145 to the classic 'Lashley maze' 55, was intended to test the ability of an individual to learn to navigate 146 147 through a complex maze ('complex maze task' from hereon; Fig. 1A). Previous studies have shown that the Lashley maze can be solved using allocentric-based visual spatial cues, and/or varying 148 degrees of egocentric-based learned turn sequences when these are absent or insufficient <sup>56</sup>. We have 149 150 previously shown that pheasants show individual-level variation in their preference for using allocentric and egocentric processes when solving this same complex maze task <sup>57</sup>, suggesting that 151 152 performance in this maze reflects at least some degree of spatial reference memory ability in this 153 species, regardless of navigation strategy deployed. Indeed, whilst multiple cognitive modalities can be used to solve even explicitly domain-targeted tasks (e.g. <sup>58</sup>), we chose these three tasks due to their 154

extensive prior validation from classical cognition testing, their suitability for testing with pheasants,
and specifically for their likely relevance to processes important in animal movement ecology <sup>21</sup>.

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158 After the completion of their cognitive tests, we simultaneously released the juvenile pheasants into a 159 heterogeneous rural landscape (mixture of agricultural, grassland and woodland habitats) at 10 weeks 160 old (the age wild birds start independence from their mother <sup>59</sup>). We followed their movements and fate using ATLAS, a recently-developed reverse-GPS automated radio-tracking system with high 161 spatial and temporal resolution <sup>60–63</sup>, detecting locations at up to eight times per minute, for up to 120 162 days as birds acquired their home ranges. This design ensured that all birds were equally naïve to a 163 164 shared landscape, allowing us to test whether the stabilized size of an individual's home range was predicted by their earlier performance in our three cognitive tests (accounting for prior experience; 165 difficult with studies of adult animals but critical when comparing developmental patterns of 166 behaviour between individuals<sup>14</sup>). 167

168

#### 169 *Results and Discussion*

170 We found that pheasants exhibiting superior performance in the complex maze task (designed to test 171 navigation-related spatial cognition, including spatial reference memory) subsequently developed 172 larger core home ranges (i.e. the area of the home range that birds spend a disproportionately large 173 amount of time<sup>11</sup>; see Methods) calculated from autocorrelated kernel density estimates (AKDEs, 174 which control for temporal autocorrelation in location data<sup>64</sup>) (-0.52  $\pm$  0.24 (coefficient  $\pm$  standard error); Fig. 1D; Table S1 and Table S3). This positive relationship provides direct empirical support 175 for the long-standing implicit home range hypothesis suggesting that greater cognitive ability, 176 particularly spatial memory, allows an individual to form a larger core home range <sup>15,65,66</sup>. In contrast, 177 178 we found no evidence that spatial working memory or associative learning ability predicted core 179 home range size, or that any cognitive metrics predicted the size of the peripheral home range (area of 180 the 85% AKDE isopleth subtracted from the area of the 99% AKDE isopleth; Table S2 – Table S6).

181 Comparison of our results to two previous mechanistic studies that incorporated multiple spatial 182 memory systems show interesting discrepancies. Studies by both Van Moorter et al, and Riotte-183 Lambert et al suggest that home range size should strongly covary with working memory, and in the former study, be largely insensitive to variation in reference memory <sup>21,22</sup>, with both memory systems 184 185 working synergistically to result in the optimal exploitation of resources that causes the emergence of 186 realistic home range patterns. Our study suggests that variation in spatial reference memory, 187 independent of working memory, predicts the development of differing home range sizes (also suggested by Riotte-Lambert et al.<sup>22</sup>), with no evidence for an interaction between these two spatial 188 189 cognitive processes or any effect of working memory variation on home range size at all. We note that 190 comparisons between theoretical and empirically-derived measures of cognition should be made with 191 caution, but nonetheless a fascinating avenue for future mechanistic studies would be to investigate 192 the relative importance played by predators and resource utilisation, and how these in turn place 193 differing priorities on different memory systems, in driving home range development, particularly in systems that suffer high predation rates such as ours (as has been done in more nomadic species <sup>67</sup>). 194

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196 We recorded 45 separate predation events during our study, for which we could identify the precise 197 timing, location and predator species responsible due to the high temporal and spatial accuracy of our 198 ATLAS data. All predation events were caused by red foxes (see Methods for a separate validation 199 experiment that characterised untagged-predator and tagged-prey movement patterns). By comparing 200 the whereabouts within the predated bird's home range of these real death locations to a null 201 distribution of simulated death locations, we find that pheasants were disproportionately likely to be 202 predated on the periphery of their core home range, the area with which they had least experience (deaths occurring on  $85\% \pm 25\%$  (mean  $\pm$  SD) isopleth of AKDE; Omnibus test:  $\chi = 242.61$ , df= 70, 203 204 P < 0.001; Fig. 2A and Fig. 2B). This suggests that familiarity with the landscape was an important determinant of an individual's predation risk, supporting previous work in deer <sup>44,49</sup>. However, a 205 206 potential alternative (but not mutually exclusive) explanation for this spatial distribution of predation, 207 not discounted in previous studies, is that prev individuals may simply establish their home ranges

208 outside of high-risk areas - mortality might be particularly high outside of the core home range simply 209 because this is the area with highest intrinsic risk. Indeed, many prey species are known to form a 'landscape of fear' by learning the temporal and spatial patterns of predation risk in their habitat <sup>47,68</sup>, 210 211 providing a possible avenue by which cognitive ability may facilitate predator avoidance. We directly 212 tested for this possibility by calculating areas of high predation risk based on the predation events we 213 recorded during our study. Whilst these were non-randomly distributed across the landscape, we 214 found no indication that cognitive ability played a role in pheasants' ability to avoid these high-215 mortality areas (Tables S7 – S10), or indeed any evidence that pheasants avoided these areas at all 216 (Extended Data Figure 1). Instead, the home ranges of surviving birds overlapped substantially more 217 with the high-mortality regions than we would expect by chance  $(89.1\% \pm 22.2 \text{ (mean} \pm \text{standard}))$ deviation) of a bird's core home range; Permutation test: P=0.010; Extended Data Figure 1), possibly 218 due to predators being attracted to areas of high prey abundance <sup>69</sup>. Crucially, the locations of each 219 220 death commonly fell within the home ranges of multiple individuals. We were therefore able to test, at 221 a fine scale, whether it was an individual's own experience (as determined by the percentage kernel 222 isopleth of their home range) of the location that predicted their death rather than the intrinsic risk of 223 the location itself. For the bird killed at a particular location, this was situated more peripherally 224 within their home ranges compared to the other birds whose home ranges encompassed the location 225 yet were not killed by predators during the study ( $65\% \pm 34\%$  (mean  $\pm$  SD) isopleth of AKDE; 226 Kolmogorov-Smirnov test; D=0.32, P=0.001; Extended Data Figure 2). Taken together, these results 227 suggest that predation risk is not simply a product of the probability of interacting with predators, but 228 rather the level of experience an individual has with the landscape when it encounters those predators. 229 Ignorance of local risk at the home range edge may manifest as a failure to optimally resolve the 230 trade-off between vigilance and resource use, which may develop with age and experience as 231 individuals obtain knowledge about the spatial and temporal patterns of predation risk, refuges and escape routes within their environment <sup>47,68,70</sup>, rendering juveniles (or otherwise naïve individuals) in 232 233 rarely visited areas at greatest risk of predation.

235 Further corroborating the antipredatory importance of spatial knowledge, the familiarity-dependent 236 patterning of predation risk we found also created differential survival in pheasants contingent on 237 their spatial memory. Survival was highest in pheasants whose performance in the complex maze task 238 (assessing spatial reference memory) was correlated with their performance in the working memory 239 task; birds had the highest survival if they combined good spatial working memory abilities with good 240 spatial reference memory, and the lowest survival was found in birds with mismatching spatial 241 abilities (i.e. they performed well in one but not the other) (Cox proportional hazards (Cph) model; 242 reference memory x working memory interaction:  $-2.98 \pm 0.90$  (coefficient  $\pm$  SE); Fig. 2C; Table 243 S11- Table S13). When controlling for this interaction, birds with better performance in the complex 244 maze task also had an overall lower probability of being predated (Cph:  $-1.02 \pm 0.44$  (coefficient  $\pm$ 245 SE); Table S13), although there was no important effect of working memory as a main effect (Cph: -246  $0.45 \pm 0.59$  (coefficient  $\pm$  SE); Table S13). We found no support for an effect of general associative 247 learning on survival, where our two best models included only sex or an intercept-only model (Table S14). Given that spatial cognition in pheasants has a (albeit low) heritable component <sup>71</sup>, the 248 249 familiarity-dependent spatial variation in predation risk we detail here has potential to generate 250 selection on these two forms of spatial memory. An important prediction from our results is that, since 251 spatial memory positively predicts home range size, with predation risk being highest just outside the 252 core home range, and that this predation risk in turn generates selection on spatial memory, then core 253 home range itself should be expected to be under directional, positive selection. Confirming this, we 254 find that indeed the size of the core home range itself also predicted mortality, with predation risk 255 being lowest in the birds with the largest home ranges (Cph:  $0.12 \pm 0.07$  (coefficient  $\pm$  SE); Fig. 2D).

256

A small number of previous studies have demonstrated survival or mate choice benefits that are associated with cognitive abilities <sup>40,42,43,72</sup>. However, the process by which cognition actually conveys these benefits by mediating how animals interact with their environment is unclear. Our results reveal that spatial variation in predation risk, arising from a prey individual's own familiarity within its home range, may generate selection on spatial cognitive abilities that in turn predict home range size.

262 An intriguing result from our data is that survival was also relatively high in birds that performed 263 poorly in the tasks that test both spatial working memory as well as navigational-based spatial 264 cognition (spatial reference memory) (Fig. 2C). This result requires further exploration to determine 265 the overall effect on selection experienced across multiple generations, although the overall effect of 266 highest survival being found in the birds that performed best in both, in combination with the highest 267 mortality being found in birds with mismatching performance, should result in overall positive 268 selection in both cognitive abilities (as suggested by our result showing that birds with the largest 269 home ranges also had highest survival). Previous modelling work has highlighted the importance of a 270 synergistic mechanistic role of working and reference memory for an animal to create a home range that optimises foraging efficiency <sup>21,22</sup>; our results suggest that a mismatch between these two facets 271 272 of spatial memory may also have ultimate consequences in terms of survival. Whilst spatial reference 273 memory is known to be important in allowing prey to efficiently exploit safe refuges, particularly when encountering predators <sup>73–75</sup>, the role of working memory in relation to predation risk is poorly 274 275 known so we can only speculate on this intriguing result. Perhaps birds with greater reference but 276 poor working memory traverse larger home ranges, with correspondingly greater exposure to high-277 risk areas, but their poor working memory means they are less capable of exploiting recently visited 278 escape routes should they encounter a predator. Regardless of the explanatory factor for this result, it 279 suggests that interactions between complementary modalities may be required for selection on some 280 cognitive abilities to arise. The nature of our study site, with the simultaneous introduction of same-281 aged, equally naïve birds at a single location, in the virtual absence of resident competitors and 282 abundant natural and provisioned food mean that we cannot explore the role of resource distribution 283 on home range development, maintenance and benefits in this analysis. Fluctuations or heterogeneity 284 in resources or competition likely affect home ranges with larger ranges expected in low-productivity 285 areas with patchily distributed resources. Under such circumstances, the relative values of working 286 and reference memories may differ. Thus, we encourage future studies to explore the interplay 287 between risk or resource distributions and spatial memory ability to better understand how selection 288 may act more generally on cognition under varying ecological conditions.

289	
290	Our findings provide empirical support for the long-assumed links proposed between movement
291	ecology and spatial cognition as revealed in abstract cognitive tasks dependent on cues of space and
292	movement. They suggest that spatially-contingent predation risk arises through familiarity and
293	experience with the landscape, a potentially critical factor with important implications for how
294	cognition shapes movement and space use. Importantly, they provide a plausible pathway by which
295	selection acts on spatial learning and memory through an individual's acquisition of, and familiarity
296	with, their home range, and their life or death within it.
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298	
299	Methods
300	
301	Housing and rearing conditions

302 On the 24th May 2018 we hatched 126 pheasant chicks from eggs obtained from commercial breeders 303 (which sell large numbers of pheasant eggs each year) and allocated these approximately equally 304 across four indoor enclosures (each 2m x 2m). Chicks were provided water and fed commercial chick 305 crumbs (Sportsman<sup>©</sup> Game Feed) ad libitum, provided a brooder heat lamp for thermoregulation, and 306 given various perches. At eight days old, we tagged all birds with small patagial wing tags (Roxan 307 Ltd, Selkirk, UK) to allow individual identification. Chicks were given access to a sheltered outdoor 308 enclosure (1.5 x 2.4m) at two weeks old and an outdoor run (4m x 12m) at four weeks old. At 10 309 weeks old, we weighed and measured the tarsus length of all individuals (a common proxy for skeletal body size in birds<sup>76</sup>) before releasing all birds into a large outdoor pen allowing them to 310 311 disperse into the surrounding rural landscape (mixed farmland and woodland; North Wyke Farm, Okehampton, Devon, UK). Prior to the release of our study birds, all wild pheasants were captured for 312 313 a captive breeding programme. Pre-release visual surveys, as well as ongoing observations during the 314 study, confirmed that less than 10 wild birds were ever present in the entire field site during the study.

315

# **316** *Cognitive testing*

All cognitive tasks took place within the same 75cm x 75cm testing chamber adjacent to their indoor enclosure. For the first three weeks we trained chicks to enter this chamber via a remotely controlled sliding door using live mealworms as rewards. An additional remotely controlled door with access to the outdoor enclosure enabled us to ensure birds entered and were tested within the testing chamber individually. Trial and session number represent a compromise between maximising the collection of cognitive data that our and previous work has suggested should allow the collection of learning slopes (e.g. <sup>57,71</sup>) against the time and logistical constraints of running this multi-stage study.

324

325 Colour discrimination task

326 At 25 days old, we tested the chick's associative learning ability on a binary-choice colour

327 discrimination task that required birds to pierce tissue paper-covered wells to reach a mealworm food

328 reward (Fig. 1C). One week prior to testing, birds were trained to peck through the tissue paper via 329 repeated exposure to the apparatus in the testing chamber in small groups until all birds could 330 consistently perform this behaviour. Each bird completed 10 trials within each of eight sessions (80 binary choices) between the 19<sup>th</sup> and 22<sup>nd</sup> June. In each session, a bird would enter the testing 331 332 chamber where they then completed a total of 10 trials. Each 2cm diameter well was encircled with 333 either a blue or green coloured ring; wells encircled with a blue ring contained three mealworms, 334 while those encircled with a green ring could not be pierced due to a layer of undetectable cardboard 335 immediately underneath the tissue. The two wells were oriented vertically in relation to the pheasant's 336 perspective, with a 1cm gap between them. We placed a dead mealworm between the two wells to 337 attract birds to the apparatus and ensure they had a standardised orientation for each set of trials. Birds were only allowed to make one choice per trial. After each choice, the apparatus was replaced before 338 339 starting the next trial. The location of the rewarded well was pseudorandomised so that it did not 340 appear on the top or bottom more than 3 times in a row and thus removing any spatial associations 341 that may form.

342

#### 343 *Complex maze task*

Between  $3^{rd} - 5^{th}$  July 2018 (once the birds reached five weeks old), the testing chamber was 344 converted into a 3 x 3 chambered maze using 35cm walls of opaque black plastic to test the 345 346 pheasants' reference learning ability, recalling a route after an interval of several hours (Fig. 1A). We 347 first habituated birds to this arena by repeatedly feeding them within the maze when all the doors to 348 each chamber were open. During the complex maze task, birds were required to make at least seven 349 turns to navigate to the exit, at which point they would receive a mealworm reward and could exit to 350 the outdoor enclosure. We ensured that all cues external to the maze (e.g. observer position) were 351 identical in all trials. Each bird first underwent a total of eight trials. In each trial, we counted the 352 number of chambers that a bird entered erroneously, with an error being defined as a chamber that did 353 not lie on the most direct route to the exit. A bird was considered to have entered a chamber if their 354 head passed over the chamber threshold. A bird could repeatedly re-enter a wrong chamber so the

number of (potential) errors was not capped. In a perfect performance with no errors, the bird wouldprogress directly through the maze via seven chambers to the exit.

357

### 358 Radial arm maze

359 Once chicks had reached seven weeks old, we tested their working memory based on spatial cues over 360 a period of a couple of minutes, by converting the testing chamber into a radial arm-style maze with 361 four arms rewarded with a single large mealworm (Fig. 1B). The reward was concealed at the end of 362 each arm by an opaque black plastic wall which the birds had to navigate around. On entry to the 363 testing chamber, birds were lured to approach a central platform using a single mealworm as bait, 364 after which they could enter one of the arms and so make their decision. In each trial, we recorded the 365 number of errors a bird made (number of times a bird stepped off the central platform into an arm 366 they had already depleted the reward from) and a trial continued until a bird found all the mealworm 367 rewards, at which point it was released into the outdoor enclosure. All birds received a total of 12 368 trials. A perfect performance would see a bird visiting and depleting each location just once, with no 369 visits to previously visited locations.

370

# 371 *Cognitive performance analyses*

372 We quantified an individual's learning performance in the radial arm and complex maze tasks by using the slope that describes their reduction in numbers of errors across trials, and for the colour 373 374 discrimination task by using the slope that describes the number of correct choices out of the 10 trials 375 across the eight sessions. For the colour discrimination task, we obtained the slope from running a 376 Poisson regression between the number of correct choices out of the 10 trials across the eight sessions 377 of the task. For the complex maze and radial arm maze tasks, we fitted a linear relationship to the 378 pattern of errors across the 8 and 12 trials respectively. We derived the equation for the relationship 379 and used the coefficient slope value. Steeper negative slopes for the radial arm and complex mazes,

and steeper positive slopes for the colour discrimination task, indicate that their learning of the taskwas faster (although these are inversed in our figures to aid interpretation; Fig. 1D and Fig. 2C).

We only included birds that had completed at least eight of the ten trials in the radial arm maze and
colour discrimination tasks, and seven of the eight trials in the complex maze for further analysis.
Different birds reached criterion on different tests, which is why sample sizes for relationships with

385 particular tests vary, and also why not all birds that contributed death location data also contributed

386 spatial ability data (and vice-versa if birds did not establish a stable home range; see below). None of

387 our cognitive variables were significantly correlated (linear model (lm) of spatial reference memory

**388** predicted by spatial reference memory:  $-0.01 \pm 0.26$  (coefficient  $\pm$  standard error),  $F_{1,42} = <0.01$ ,

**389** P=0.999; lm of spatial reference memory predicted by colour discrimination:  $-0.08 \pm 3.91$  (coefficient

 $\pm$  standard error), F<sub>1,31</sub>=<0.01, P=0.983; lm of spatial working memory predicted by colour

391 discrimination:  $2.66 \pm 2.18$  (coefficient  $\pm$  standard error),  $F_{1,31}=1.49$ , P=0.232.

392

### **393** *Release into the wild and ATLAS system*

394 All birds were allowed to naturally disperse into the wild by being first released into an outdoor release pen (~4000 m<sup>2</sup> in area) within woodland at North Wyke Farm, North Wyke, UK, once they 395 396 reached ten weeks old. Before release, they were fitted with radio tags, sexed, and their mass was 397 recorded. Within the pen, birds could enter and leave as necessary by flying or via a system of wire 398 tunnels that were too small to admit terrestrial predators. We provided birds with supplementary food 399 supplied in 43 artificial feeders situated inside and within 2 km of the release pen (see Fig. 2A for 400 map of the release site). To ensure the attached radio tags had no adverse effects, tagged birds were 401 monitored in captivity for seven days before being released. Specifically, we monitored behavioural 402 (lethargy, isolation) and physiological (breathing, vocalisation) indicators of welfare as well as 403 visually monitoring the pheasants for sites on the body that could be susceptible to abrasion. We tracked birds using a recently developed reverse-GPS system (ATLAS)<sup>60,61,63</sup>. This system uses 404 405 fixed-position receiver stations to detect and collect the time of arrival data from tag-derived radio406 signals. These data are then collated at a database on a central server where location is calculated.

407 Location data, accurate to ~4-6m relative to GPS measurements <sup>63</sup>, can be accessed in real-time

408 through an internet connection which allows for continuous monitoring.

409

410 Determination of death locations

411 Because the major predator of pheasants in our field sites, red foxes (Vulpes vulpes), will frequently 412 transport their captured prey before eating or caching them, we combined three methods to determine 413 the exact death location of each of our predated pheasants. Firstly, we ran an observational study using ATLAS-tagged fresh pheasant carcasses to determine the range of movement patterns displayed 414 415 by predators carrying a dead pheasant (details of experiment below). Secondly, informed by the 416 filtered movement tracks of scavenged dead pheasants, we visually assessed movement patterns of 417 our live birds to subjectively determine putative kill sites where the bird's movement path underwent 418 an unusual substantial change (e.g. when the bird rapidly covered a large distance in an 419 uncharacteristically straight line before suddenly stopping) and matched those displayed by the 420 movement patterns of scavenged carcasses. Thirdly, we confirmed our putative kill sites by running a 421 state change analysis on the tracked paths of each predated bird to determine whether locations that 422 immediately preceded a substantial change in movement highlighted by our subjective assessment 423 matched those determined using our state change analysis. Specifically, we used the segmentation approach by Lavielle<sup>77</sup>, where segment numbers (allowing us to identify change-points) were 424 425 identified based on graphically examining the relationship between segment number and the 426 decreasing contrast function, produced using the 'lavielle' function in the 'adehabitatLT' R package  $(version 0.3.25)^{78}$ . 427

In our carcass tracking study, we placed 44 carcasses of ATLAS tagged adult pheasants (22 male and
22 female) on 5<sup>th</sup> February 2019 in locations around the field site frequented by our live tracked birds.
Two camera traps (Bushnell Trophy) were setup at each location to identify the predator that moved
the carcass. After eight days (once all carcasses had been moved), we then located carcasses based on

their ATLAS coordinates. Camera trap footage confirmed that all carcasses moved from the
introduction site were caused by red foxes. ATLAS coordinates for each bird were then filtered using
the same method for our real birds, and the resulting movement patterns were used to help denote
death locations as described above.

436 In all instances of predation from our main release experiment, the paths of live birds that were 437 predated were typified by regular movement patterns before a sudden and uncharacteristically long-438 distance and rapid movement in a straight line before immediately becoming stationary. This rapid, 439 single straight-line movement was recorded for all red fox scavenging instances we recorded in the 440 carcass tracking study above, representing a fox moving its prey a long distance from the site it was 441 captured to a new location to process and cache. This first 'stop' location was usually the spot where 442 we recovered the predated bird via radio tracking, and the location immediately preceding the first 443 long-distance movement (i.e. the last point of normal pheasant movement) was denoted as the 444 location the bird was predated. We confirmed the species of predator using field signs around 445 carcasses such as whether the carcass was buried, teeth marks on the radio-tags, and feather remains 446 (e.g. characteristic field signs of a fox kill are large wing and tail feathers cleanly severed close to the 447 base of the feather shaft, whereas raptors pluck feathers by grasping the feather further up the shaft, 448 leaving a characteristic V-shaped kink). All the birds killed outside their pens in this study were 449 predated by red foxes.

450

## 451 *Location data filtering*

452 All ATLAS coordinates were first filtered to increase the accuracy of locations used to calculate the 453 home ranges. We first removed all coordinates where the variance in signal strength between base 454 stations was greater than 75%. Each day was then split into five-minute blocks, and the median 455 location during each five-minute block was calculated (so that every bird in the study had one location 456 assigned within the same time periods within each day). This filtering process excluded anomalous 457 locations caused by standard issues with radiotracking such as reflection <sup>79,80</sup>. Because the release pens which birds were kept in before dispersing effectively excluded foxes, we only included thecoordinates of birds that were outside the pen in our home range estimations.

460

#### 461 *Home range estimation*

To estimate home ranges of each pheasant we fitted a continuous-time stochastic movement model 462 463 (ctmm) followed by AIC-based model selection to calculate autocorrelated kernel density estimates (AKDEs) for each of our birds using the *ctmm* package <sup>64</sup> in R (version 3.5.3) <sup>81</sup>. Parameters for our 464 465 ctmms were chosen based on visually assessing variograms of time-series movement patterns for each 466 individual bird. The variograms also allowed us to identify and discard from the analysis a further 39 467 birds who had not formed a stable home range because of either predation or death soon after release, 468 tracker malfunction, or because they quickly dispersed from the study site. Core home ranges were estimated as the 85% AKDE isopleth based on the mean point of asymptote between the home range 469 470 size and AKDE isopleth, indicating the region within which birds were concentrating their movement 471 patterns <sup>11</sup>. To determine the region of the home range where birds were killed, we then calculated the 472 minimum AKDE isopleth that encompassed the coordinates of the bird's predation location.

473

## 474 Do pheasants avoid high-risk areas?

475 One explanation for why pheasants are killed more frequently on the outskirts of their home range is because they avoid riskier areas - death locations on the home range periphery may simply reflect 476 477 cases where pheasants ventured into high-risk areas instead of occurring due to a lack of intrinsic 478 knowledge about local predation risk. To test whether our mortality patterns could therefore be driven 479 by birds avoiding areas of high risk, we quantified the home ranges of non-predated pheasants and 480 determined the proportion of their core home range (85% AKDE) that overlapped with a kernel 481 density estimate (80% kernel estimate with no autocorrelated time estimate) created from the death 482 locations of our predated birds. The 80% kernel density estimate from the actual death locations was 483 chosen as a suitable 'high risk zone' as it represented the start of the plateau between the kernel

484 density percentage and size (surface area) of the home range, thus representing an area of485 concentrated predation risk.

486

487 To determine whether the home range-risk area overlap differed from what we would expect by 488 chance, based on the null hypothesis that pheasants can be predated with equal probability at any 489 previously visited location within their home range, we compared the overlaps of our real birds to 490 5000 simulated high-risk zones. Each simulated high-risk zone was created by selecting a single relocation coordinate (randomly selected from any of the coordinates recorded during a bird's 491 492 lifetime) from each of our predated birds and using this as a simulated predation location, before re-493 calculating a new high-risk zone and subsequently calculating how much this overlapped with the real 494 core home range of the surviving birds. Only those surviving birds that overlapped with some part of 495 the home range of at least one of the predated birds were included in this analysis. The proportion of 496 randomised survivor-predation home range overlaps that were lower or higher than the real survivor-497 predation home range overlaps was used as our two-tailed statistical significance value (Extended 498 Data Figure 2).

499

500 In addition to the above, we also ran an analysis to determine whether the AKDE isopleth containing 501 a death location was higher in the bird that actually died compared to the same death location of a 502 neighbouring bird that did not die. This is directly testing whether the actual death location itself was 503 avoided at a more fine-scale level than can be determined by comparing home range overlaps with 504 regions of high risk. We therefore calculated the minimum AKDE isopleth that contained a death 505 location for birds that were killed and birds that were not recorded as being killed (surviving birds; 506 n=67). Surviving birds were only included if the 100% minimum convex polygon of their home range 507 encompassed the death location of the killed bird (to avoid including birds that never encountered the 508 death location). We then compared the distribution of minimum isopleths that contained death

509 locations between killed and surviving birds using a Kolmogorov-Smirnov test (Extended Data Figure510 2).

511

512 *Statistical analysis* 

513 To determine the drivers of home range variation, we ran linear models (LMs) with core and 514 peripheral home range size in hectares of the eventual stable home range of each pheasant as the 515 response variables. We define the peripheral home range as the area covered by the core (85% kernel 516 isopleth) subtracted from the area of the total (99% kernel isopleth) home range. We used this 517 measure rather than the 99% kernel to detect any effects of total home range size exclusive of the role 518 played by the underlying core home range (which is of course a composite of the total home range). 519 Both core home range size and peripheral home range size were strongly left skewed and so were log-520 transformed to obtain normally-distributed residuals from our models. To test the effect of different 521 cognitive modalities on these home range parameters we created a model to test performance in the 522 colour discrimination task, and a model that tested for the role of performance in both the spatial 523 cognition tasks, across both of the response variables detailed above. To be included in any analysis, 524 pheasants of course had to reach criterion in the relative cognitive tests (two tests in the case of the 525 spatial analyses), as well as form a stable home range before dying/suffering a transmitter 526 malfunction/dispersing out of the field site. For the survival analysis, we also only include birds 527 where the known cause of death was predation (as determined from above), or to include birds that 528 'survived', we only included their location data up to the point that they were still moving freely and 529 were producing high-accuracy data (i.e. hadn't dispersed outside of the range of the ATLAS 530 receivers). As such, there was rather extensive data loss from our original 126 birds (see specific 531 numbers in each statistical section below).

532

533 In the spatial cognition models, performance in the radial arm maze and complex maze were included534 as main effects to test for the general hypothesis that spatial reference memory is important in the

development and utilisation of a home range (e.g. <sup>25,26</sup>), as well as in a two-way interaction to 535 536 explicitly test the hypothesis that home range development and size arise and vary due to the synergistic effect of working and reference memory <sup>21</sup>. To control for the amount of time that a bird 537 538 could familiarise itself with the landscape, we also included a bird's lifespan (in days) as a covariate 539 in all models. In addition, pheasants show sex differences in home range size, with females having 540 larger home ranges than males when the latter become territorial during the breeding season (although 541 our study did not encompass this period), and so we therefore include sex in our models as a fixed term <sup>82,83</sup>. Our code syntax for the structure of our starting maximal model for the core home range 542 543 spatial analysis (n=44) was therefore:  $lm(log(Core \ HR \ hr) \sim Slope \ LM * Slope \ RAM + Sex +$ 544 Lifespan, data = akde data main)', and we returned support for a single model in our model selection 545 procedure (Table S1). To test whether body mass is a confound in this analysis (since home range is 546 known to scale with body mass at the intraspecific level in some species <sup>84</sup>, although see <sup>36</sup>), we also 547 ran a final model that included only lifespan and mass as predictors. Model comparison using AICc (154.1 for the reference memory model and 156.3 for the mass model) confirmed that mass is not a 548 549 confound. We intentionally excluded mass in our starting model to avoid issues with collinearity since 550 it is strongly correlated with sex (female mass:  $575g \pm 65$ ; male mass:  $717g \pm 100$  (mean  $\pm$  standard deviation); t-test: t = -5.65, df = 37.2, P<0.001). 551

552

553 Our code syntax for the structure of the starting maximal model for the peripheral home range spatial 554 analysis (n=44) was:  $lm(log(Peripheral_HR_H) \sim Slope_LM * Slope_RAM + Sex + Lifespan, data=$  $555 akde_data_main)'. Our model selection procedure produced two minimum adequate models (Table$ 556 S4). As with the core home range model above, we also tested for any confounding effect of mass by557 comparing our two final model AICcs with a model that only contained mass and lifespan as558 predictors, confirming again that it is not a confound.

560 For the associative learning analyses based on the colour discrimination task, we included sex, colour 561 discrimination performance and lifespan as the fixed effects. The syntax for our starting maximal 562 model for the core home range colour discrimination analysis (n=55) was therefore:  $lm(log(core \ HR \ hr) \sim CD \ Slope + Sex + Lifespan, data = akde \ CD)'$ , returning a single model 563 564 with only sex and lifespan as covariates. The syntax for our starting maximal model for the peripheral 565 home range colour discrimination analysis (n=55) was:  $\ln(\log(Peripheral HR hr) \sim CD Slope +$ 566 Sex + Lifespan, data = akde CD)'. This produced three separate minimum adequate models (Table 567 S5).

568

569 A central question in this study was to test whether animals were more likely to be predated on the 570 outskirts of their home range, in areas they were less familiar. However, the null expectation of where 571 predation locations should occur across a home range density estimate such as AKDE are unclear. As 572 such, we took a simulation approach, comparing the AKDE isopleth that a real death occurred on to a null distribution of 'simulated death isopleths', where every location a bird was recorded was 573 574 assigned as a simulated death location and the accompanying minimum AKDE isopleth that 575 encompassed it. To calculate our total P value for each bird, we then calculated the proportion of 576 simulated death isopleths that were higher or lower than the observed death isopleth, and tested for 577 overall significance of all birds by combining the P values using Fisher's combination (omnibus) test 85. 578

579

We also ran two binomial regression analyses investigating whether cognitive ability predicted the proportion of a pheasant's core home range that overlapped with the region of high predation risk. We again separated spatial memory and associative learning into separate models to maximise our sample sizes and statistical power, using a conditional log-link function in our models to control for the unequal distribution of our proportion data. We controlled for the size of the core home range in driving these patterns by including this variable as a covariate in all models. The syntax for our

586 maximal starting model for spatial memory (n=29) was: ' $glm(HR \ death \ overlap \ prop \sim Slope \ LM *$ 

587 Slope\_RAM + core\_HR + Sex, family= binomial(link= "cloglog"), data=

588 *akde\_cognition\_HR\_overlap)*' and for associative learning (n=29): ' $glm(HR\_death\_overlap\_prop \sim$ 

589 *core\_HR* + *CD\_Slope* + *Sex, family= binomial, data= HR\_overlap\_CD*)'. We identified three

- 590 candidate models for the spatial memory analysis and a single model for associative learning (Table
- **591** S7 and Table S8).

592

593 To determine how spatial cognitive performance and home range size affected a pheasant's 594 probability of being predated, we ran three Cox-proportional hazards models to determine the effect 595 on lifespan and general probability of death. In the first survival model we included performance in 596 the complex and radial arm maze tasks as covariates, sex, and the size of the core home range 597 (hectares) as predictors. We also directly test whether an interaction between working and reference 598 spatial memory are important for the ultimate consequences of forming a home range, as suggested by mechanistic models (e.g. <sup>21</sup>) by including an interaction between performance in our complex maze 599 600 and in our radial arm maze tasks. We also included sex within a three-way interaction that also 601 includes working and reference memory. Our model syntax for this starting model (n=51) was 602 therefore: coxph(Surv(Lifespan, State bin) ~ Sex + Slope LM\*Slope RAM, data=spatial data). Our 603 AICc based model selection returned support for a single model (Table S11). As with our home range 604 models, we also investigated whether body size was a confound by adding it to our final model 605 (where sex and was not included and could therefore not cause any collinearity issues).

606

To test for the importance of associative learning performance on survival, our second model included
performance in the colour discrimination task, sex, and tarsus size, with a single two-way interaction
between colour discrimination and sex. Our syntax for this starting model (n=61) was therefore:

610 *coxph(Surv(Lifespan, State bin) ~ CD Slope\*Sex+Tarsus, data=CD data)*). Our AICc based model

611 selection returned support for a single, intercept-only model (Table S12). Because different birds did

612 not reach criterion in the different cognitive tests and were thus removed (see above), we also ran a 613 third survival model that included all birds for which we had stable home range sizes to maximise 614 statistical power (resulting in n=68) to test if core home range size itself, independent of cognitive 615 variables, affected survival. In this model, we included three predictors: core home range size in 616 hectares, sex, and a two-way interaction between these variables. Our model syntax for this starting 617 model (n=68) was therefore:  $coxph(formula = Surv(Lifespan, State bin) \sim core HR H*Sex, data =$ 618 HR surv data), and our our AICc based model selection returned support for a single model with 619 core home range size as the sole term (see main text).

620

621 Diagnostics were performed on all models to check they met assumptions. We used a stepwise AICc (AIC corrected for small sample size) model selection approach to find our best models. Specifically, 622 623 after constructing our starting model that allowed us to test our most complex hypotheses, we 624 performed a stepwise simplification procedure based on the "AICc" function from the R package 625 'MuMin'<sup>86</sup> to compare nested models until a best set of plausible models were obtained (minimum 626 adequate models), where models with  $\Delta AICc \leq 2$  were deemed more plausible. If this process created 627 more than a single best model, we used the *model.avg* function in the 'MuMin' package to average 628 the model estimates and standard errors. In addition, we present P values, uncorrected for multiple testing (as per <sup>87</sup>) for each of our best models in the supplementary material, which were obtained 629 630 using likelihood-ratio tests for change in deviance between nested models with and without the term 631 of interest.

632

All animal work in this study was conducted under Home Office license PPL 30/3204 and approvedby the University of Exeter Animal Welfare Ethical Review Board.

635

636 Data availability

- 637 Data required to rerun the statistical analyses of this study are available online
- 638 (https://data.mendeley.com/datasets/m89226xg6p)<sup>88</sup>. Animal AKDE models and GPS coordinates are
- available from the corresponding author upon request.
- 640

	641	Code	availa	bility
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- 642 R code used to run the simulation analyses of this study are available online
- 643 (https://data.mendeley.com/datasets/m89226xg6p)<sup>88</sup>.
- 644

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654

# 655 Author contributions

656 RJPH and JRM conceived the idea for the manuscript. CEB, PRL, MAW, JOvH and JRM collected

the cognition data. MAW, CEB and JRM collected the movement data. RJPH and MAW carried out

- the carcass tracking study. RJPH conducted the analyses and led the writing. RN, YO and ST
- 659 developed the reverse-GPS system and provided support throughout data collection. All authors
- 660 contributed critically to the drafts.

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# 662 Competing interests

663 The authors declare that they have no competing interests

664

# 666 Figure legends

Fig. 1. Cognitive predictors of home range size in pheasants. A: The complex maze task (A) radial
arm maze task (B) and colour discrimination tasks (C) used to assess spatial memory (A & B) and
associative learning (C), respectively, in juvenile pheasants. D: Performance in the complex maze task
positively predicted the size of the core home range in pheasants. Error bars denote the 95% CI. Xaxis in D represents the inverted coefficients taken from a regression between number of trial errors
over time for ease of interpretation, i.e. a positive slope equals increased learning rates.

Fig. 2. Spatial patterns of pheasant mortality and consequences for selection on spatial memory and 674 675 home range. A. Map of the release site showing an example pheasant home range showing 85% (dark 676 blue) and 99% (pale blue) kernel estimates in relation to where a bird was predated (red dot). B. Histograms showing the percentage kernel isopleths that contained the observed death locations (left) 677 678 compared to the kernel isopleths that contained simulated death locations (right) taken from the same killed individuals. C. Mortality risk from predators depends on the interaction between working and 679 680 reference memory. Complex maze and radial arm maze performance measures represent inverted 681 coefficients taken from a regression between trial errors over time for ease of interpretation. D. 682 Predation risk in pheasants is predicted by core home range size. Error bars denote the 95% CI. Map 683 layer Imagery ©2022 Google Maps, CNES / Airbus, Getmapping pic, Infoterra Ltd & Bluesky, 684 Maxar Technologies, Map data @2022. 685

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