1 Seal carrion is a predictable resource for coastal ecosystems

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13 Abstract

14 The timing, magnitude, and spatial distribution of resource inputs can have large 15 effects on dependent organisms. Few studies have examined the predictability of 16 such resources and no standard ecological measure of predictability exists. We 17 examined the potential predictability of carrion resources provided by one of the UK's 18 largest grey seal (Halichoerus grypus) colonies, on the Isle of May, Scotland. We 19 used aerial (11 years) and ground surveys (3 years) to quantify the variability in time, 20 space, quantity (kg), and quality (MJ) of seal carrion during the seal pupping season. 21 We then compared the potential predictability of seal carrion to other periodic 22 changes in food availability in nature. An average of 6893 kg of carrion ·yr⁻¹ 23 corresponding to 110.5 x 10³ MJ·yr⁻¹ was released for potential scavengers as 24 placentae and dead animals. A fifth of the total biomass from dead seals was 25 consumed by the end of the pupping season, mostly by avian scavengers. The 26 spatial distribution of carcasses was similar across years, and 28% of the area 27 containing >10 carcasses ha⁻¹ was shared among all years. Relative standard errors 28 (RSE) in space, time, quantity, and quality of carrion were all below 34%. This is 29 similar to other allochthonous-dependent ecosystems, such as those affected by 30 migratory salmon, and indicates high predictability of seal carrion as a resource. Our 31 study illustrates how to quantify predictability in carrion, which is of general relevance 32 to ecosystems that are dependent on this resource. We also highlight the importance 33 of carrion to marine coastal ecosystems, where it sustains avian scavengers thus 34 affecting ecosystem structure and function. 35 36 37

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- **Keywords:** Predictability, Resource Subsidy, Coastal, Carcass, Pinniped, Relative
- 46 standard error, Scavenger.

47 **1 Introduction**

Many ecosystems are spatially linked by flows of nutrients and energy (e.g. Polis et al. 1997; Power and Rainey 2000; Anderson and Polis 1998; Reiners and Driese 2001). Such flows shape the structure and function of donor and receiving ecosystems by regulating nutrient availability and the dynamics of consumers that depend on them (Polis and Hurd 1996b).

53 Coastal regions are often affected by marine-derived inputs transferred from 54 the ocean to the terrestrial ecosystem (Polis and Hurd 1996a, b; Polis et al. 1997; 55 Rose and Polis 1998; Stapp and Polis 2003; Barton 2015). A noticeable example of 56 this is the stranding of plant detritus and carrion on the shore (Polis et al. 1996). 57 Animals also play a key role in the transport of marine nutrients when they move 58 from one system to another. Salmon (Cederholm et al. 1999), sea turtles (Bouchard 59 and Bjorndal 2000) and penguins (Erskine et al. 1998) play a fundamental role in this 60 process by congregating at similar times of the year and in defined areas. The 61 periodic availability of carrion and other nutrients released at such animal 62 aggregations attract terrestrial foragers to the coast (e.g. Polis and Hurd 1995), locally increase the number of invertebrate consumers (e.g. Sánchez-Piñero and 63 64 Polis 2000; Janetski et al. 2009; Spiller et al. 2010), and nourish soil and plants 65 (Fariña et al. 2003). Despite the known effects of such inputs on both consumers and plants, few studies have examined the variation and predictability of these 66 67 resources in coastal ecosystems. Whether a periodic resource pulse is predictable or 68 not has important implications for how they are used by consumers and how they 69 affect ecosystem function.

70 A challenge to understanding resource predictability is its definition and 71 measurement. If similar food resources occur in similar places and at similar times 72 each year we might consider this to be "predictable", but how should "predictability" 73 be guantified? Colwell (1974) was one of the first to define the predictability of 74 periodic phenomena, by using their constancy (temporal uniformity) and contingency (consistency of timing between years). However, this method uses data categorised 75 76 into classes and not continuous measures of true totals. Alternatively, a pulsed 77 resource can be described as having low frequency, large magnitude and short 78 duration (Yang et al. 2008), but these attributes can characterise both unpredictable 79 and predictable resource pulses. Other attributes of predictability can be relevant

80 from a consumer perspective. This can include whether resource availability is 81 related to day length or the lunar cycle (Horning and Trillmich 1999), or whether it is 82 stable in space and time (Cama et al. 2012), thus permitting consumers to find it 83 (Weimerskirch 2007) and adapt their foraging behaviour to its availability (Overington 84 and Lefebvre 2011). Although there are many examples of animal responses to the 85 predictability of resource subsidies at both ecosystem and individual levels (e.g. 86 Davenport 1995; Graham et al. 2006; Furness et al. 2007; Hoogenboom et al. 2013; 87 Reid et al. 2012; Monsarrat et al. 2013; Oro et al. 2013), a clear definition of 88 predictability is often not given. This means that measures of predictability are 89 typically study-specific, and cannot be compared across ecosystems or resources.

90 When a subsidy is temporally and spatially constrained, consumers can 91 predict where and when it will occur using prior knowledge and learning (Stephens 92 and Krebs 1986), or programmed and evolved responses to signals (Berthold 1996). 93 Female savanna elephants (Loxodonta africana), for instance, use past experience 94 to adjust their timing of reproduction according to seasonal pulses in vegetation 95 productivity (Wittemver et al. 2007). Seabirds arrive to certain areas at times of high 96 food availability, even if this means beginning to move before the food itself is 97 detectable (Weimerskirch 2007). In these cases, both elephants and seabirds can 98 predict resource availability, and respond to the location and timing of a resource, as 99 well as its quantity and quality. While it is clear that some animals predict timing, 100 spatial distribution, quantity, and quality of a resource, no study to our knowledge 101 has examined all these aspects of predictability at once. A method to quantify the 102 different aspects of predictability in common units will greatly help to understand how 103 the characteristics of resources affect the response of populations dependent upon 104 them and their success.

105 In this study, we examined resource predictability in a coastal island 106 ecosystem that experiences seal carrion pulses. Like many other animal 107 aggregations, the inputs of nutrients released by pinnipeds when forming breeding 108 colonies are likely to be substantial for the coastal systems. In sea lion colonies, for 109 instance, onshore mortality together with defecation are major sources of nutrients 110 (Fariña et al. 2003). Seal colonies represent an important system for studying the 111 predictability of a resource as there is an annual pulse of carrion which is used either 112 directly by above ground secondary consumers, or indirectly as a source of nutrients 113 to primary producers (Anderson and Polis 1998).

114 Here we asked the question: Is seal carrion a predictable resource for 115 scavengers on seal colonies? We investigated the seal carrion (placentae and dead 116 seals) produced during the pupping season in one of the largest grey seal 117 (Halichoerus grypus) colonies in the UK (Isle of May, Scotland). We evaluated 118 variability in the timing, spatial distribution, biomass, and energy content of this 119 carrion by calculating their relative standard errors (RSE). This provides a common 120 metric to evaluate variability across different measurement units. We hypothesised 121 that the RSEs of seal carrion production would be similar to those of resources that 122 animal consumers are known to predict (e.g. salmon runs). The flux of carrion to the 123 local avian scavenging community was estimated as the biomass consumed at the 124 end of the seal pupping season. Finally, we discuss the potential importance of seal 125 carrion for the ecosystem as a whole.

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127 **2 Methods**

128 **2.1 Study area**

129 This study was undertaken on the Isle of May (56°11'19"N, 2°33'27"W), 130 situated at the entrance to the Firth of Forth on the east coast of Scotland. The island (1.8 x 0.5 km), covers an area of 45 ha, with the long axis extending in a northwest-131 132 southeast direction (Fig. 1). The Isle of May is a designated Special Area of Conservation (SAC) because it hosts a breeding colony of grey seals. This colony 133 134 contributed ca. 4.3% to the annual UK pup production in 2010 (2153 pups born) and 135 appears to be increasing since then (SCOS 2016). 136 The main scavenger occurring on the island is the great black-backed gull

(*Larus marinus*): 40 pairs were counted nesting on the Isle of May during summer
 2012 (SNH 2012). Between October and March the Scottish resident gull population
 receives immigrants from Scandinavia and Russia (Forrester et al. 2007).

140 **2.2 Data collection**

We used a combination of aerial surveys and ground visual censuses of pups
and carcasses to quantify the characteristics of the carrion resource (placentae and
dead seals) available at the seal colony.

144 Aerial survey data

The Sea Mammal Research Unit (SMRU, University of St. Andrews) has 145 146 carried out annual aerial surveys of the Isle of May (and all other major Scottish grey 147 seal breeding colonies) since the early 1990s, in order to estimate seal pup 148 production (number of pups born per year) and the mean pupping date. Surveys 149 were carried out annually up until 2010, when the frequency was reduced to every 150 two years. The number of white coated pups and moulted pups are counted from a 151 series of 3-6 aerial photo surveys carried out at approximately 10-12 day intervals 152 throughout the breeding season. The pup counts are used together with estimates of 153 'time to moult' and 'time to leave' (Wyile, 1988) to model the birth curve and obtain 154 estimates of total pup production (with 95% confidence limits) and mean pupping 155 date. Dead pups were also counted for each survey, but pup mortality was not 156 estimated. For further details of the methods see SCOS (1996). 157 The number of dead pups counted in aerial surveys was used here to approximate 158 the number of carcasses released into the ecosystem every year for the decade 159 2000-2010 and the year 2012. The highest number of dead pups counted each year 160 (among all the aerial surveys) was considered to be the most accurate, even though 161 it is still likely to be an underestimation. To minimise this underestimation, data from 162 ground visual census of carcasses conducted in 2008 and 2012 were used (223 and 163 226 carcasses, respectively; see below). Both censuses showed a greater number of 164 dead pups than the highest count obtained by aerial surveys performed in both 165 years. Therefore an error of underestimation was calculated from the percentage of 166 dead pups missing in the temporally closest aerial survey count when the ground 167 visual census was completed in 2008 and 2012 (35.0% and 42.0% of extra 168 carcasses were found in the two years, respectively). As the underestimation was 169 reasonably consistent between the two years, the mean error (38.5%, SD = 5.0%)170 was then used to adjust counts for all other years.

171 Ground visual census data

Ground visual census of carcasses was carried out at the end of the breeding season (late November to early December) in 2008, 2012 and 2013. Carcasses were detected by a team (3-6 people) systematically searching the seal breeding areas of the island. Sex and developmental stage (from 1 to 5, according to Kovacs and Lavigne, 1986) were determined for each carcass. However, those that appeared starved, scavenged, or in late state of decay could not be scored for sex

- 178 and/or developmental stage (coded N/A). Starvelings (pups starved to death whose
- 179 carcass lacked the natural layer of blubber) and dead adults were also recorded.
- 180 GPS fixes (Garmin eTrex Summit; accuracy: <15m RMS) or marks on aerial
- 181 photographs (in 2012), were made for all dead animals.
- 182 Carcasses recorded during the ground visual census of 2012 were scored for
- 183 consumption state as follow: A = intact, B = lacked both eyes and/or showed one
- opening on the body, C = showed multiple openings, D = body looked flat and lacked
- some internal organs, head and/or some bones, E = remains (only bones, fur and
- 186 skin). To assign an estimated mass loss to each consumption state, during the 2013
- 187 pupping season 11 experimental carcasses were deployed and weighed at regular
- 188 time intervals to monitor changes in mass due to scavenging activity. The mean
- 189 cumulative mass loss (in %) for consecutive consumption states was then
- 190 calculated.

191 2.3 Data analysis

We calculated several attributes of the carrion resources on the Isle of May from
aerial and ground surveys. These included the timing, quantity, quality, spatial
distribution, and consumption of carrion.

195 **Timing of pupping**

196 Mean pupping dates for the decade 2000-2010 and 2012 were compared and 197 the inter-annual variability in timing calculated.

198 **Quantity of seal carrion**

199 Seal carrion was divided into two sources: placentae and dead seals. The 200 annual total estimated pup production was used to calculate the mean annual 201 biomass of placentae produced during the period 2000 – 2010 and 2012. For this 202 purpose average biomass of placentae was acquired by analysing placenta samples 203 collected in the field during the pupping season 2013: six whole placentae (including 204 amnion) were collected, weighed $(\pm 10q)$ and sampled in fresh condition. Biomass 205 (kg) of placentae available to the ecosystem for each year was then estimated by 206 multiplying the pup production by the average wet mass of a grey seal placenta. 207 Because pup carcasses were not weighed during the ground visual census, 208 the discrepancy between the mass of alive and dead pups estimated by using raw

209 data collected by Baily (2014) was then used to adjust mass values according to

210 Kovacs and Lavigne (1986). Differences in mass of alive (n = 80) and dead (n = 37)pups belonging to the 2nd-3rd-4th developmental stages were analysed. The 1st and 211 212 5th stages were excluded from this analysis because of low sample size. For this 213 reason, mass (kg) of pup carcasses was first calculated by using the equations 214 provided by Kovacs and Lavigne (1986) and the resulting values were finally 215 adjusted by subtracting the difference in mass found between alive and dead pups. 216 Biomass of pups was estimated considering sex and developmental stage of 217 carcasses; mass of pups of non-identifiable sex was approximated by averaging the 218 mass of male and female pups for the different developmental stages. Mass of dead 219 female pups belonging to the first developmental stage was given as a conservative 220 value for starvelings and pups where developmental stage was not known. The 221 mean maternal mass at weaning according to Pomeroy and Fedak (1999) was given 222 for female adult carcasses (117 \pm 18kg). A value of 234 kg was assigned to dead 223 male adults, as breeding adult males are typically twice the mass of females when 224 animals mate (Pomeroy and Fedak 1999). Mass of dead juvenile seals (called 225 yearlings) was assumed as 56.4 ±6.3 kg (Addison and Stobo 1993).

226 **Quality of seal carrion**

Energy content of placenta produced each year during the period 2000 – 2010 and 2012 was estimated by analysing the gross energy density (MJ kg⁻¹) of placenta samples collected in 2013 by bomb calorimetry (Sciantec Analytical Service, UK). The energy content (MJ) of the total biomass of placentae was estimated by multiplying the total biomass with the energy density of grey seal placenta.

233 The body composition of pup carcasses was calculated according to Lang et 234 al. (2011), in which percentages of water, protein and fat of grey seal pups at day 3 235 and 12 post-partum and at weaning are given. Percentages belonging to pups 236 nursed by primiparous grey seal mothers, instead of multiparous, were used as a 237 conservative approximation. Day 3 post-partum percentage of each body component 238 was given to first and second stage carcasses, but also to starvelings and pups 239 where developmental stage was not known. Day 12 post-partum percentages were 240 assigned to third stage carcasses, whereas fourth and the fifth stage carcasses were 241 assigned percentages calculated at weaning (Table 1). Body composition of female 242 adult carcasses was also estimated from Lang et al. (2011), considering percentages 243 of body components for primiparous grey seal mothers at day 12 post-partum.

244 Values provided by Lidgard et al. (2005) were used for male adult and juvenile 245 carcasses. Percentages of each body component were then transformed in terms of 246 biomass (kg) before estimating their energy content (MJ). Similar to body mass, 247 body composition might also differ between dead pups and healthy living pups. 248 Reduction or lack of blubber is evident in starvelings (Trites and Jonker 2000; Baily 249 2014). However, there is no documentation comparing alive and dead starveling 250 pups. The energy content of protein and fat was quantified based on values of 251 energy densities of 39.33 kJ g⁻¹ and 17.99 kJ g⁻¹ for fat and protein, respectively 252 (Schmidt-Nielsen 1997). Zero energy was attributed to the water component. Results 253 from this analysis represent an approximation of biomass and energy released into 254 the ecosystem at the moment when carcasses become immediately available to the 255 ecosystem. This analysis does not take into account possible variation in biomass 256 and energy content of carrion over time caused by decomposition.

257 Spatial distribution of seal carrion

To identify the areas of the island impacted by carcasses, the locations of 258 259 dead seals recorded during ground visual censuses in 2008, 2012 and 2013 were 260 used to produce maps showing the kernel smoothed intensity of the point pattern 261 created by the carcasses. From here onwards, the term *intensity* is used only when 262 referring to kernel smoothed intensity of the point process for carcasses, while the 263 term *density* is used in the strict sense in all other cases. Both measurements have the same units (n carcasses or placentae m⁻²). Kernel estimation is a tool used in 264 265 spatial ecology for detection of hotspots within the landscape (areas of high 266 abundance or biomass) (Nelson and Boots 2008). Here, the spatial pattern of dead 267 seals was analysed in order to produce maps showing the kernel smoothed intensity 268 of the point pattern created by the carcasses. Boundaries of the island and locations 269 of dead pups were projected onto a map using the British National Grid Reference 270 System based on the OSGB36 datum (Ordnance Survey Great Britain 1936). 271 The intensity of carcasses was calculated for the area of the island within the mean 272 low water boundary (697102 m²). Likelihood cross-validation was used to select the 273 proper smoothing bandwidth for each year as it assumes an inhomogeneous 274 Poisson process. The edge effect was set as 'false' as carcasses were not observed 275 outside the window represented by the island (20 x 20 m pixels were used). 276 Polygons of areas delimited by a minimum intensity of 0.001 carcasses m⁻² were 277 used as spatial thresholds of presence of carcasses and identified for each year. The

- 278 union and the intersection of these polygons was plotted to highlight areas of
- 279 persistent availability of carrion from year to year. Simple biomass density (kg m⁻²)
- was also calculated for the whole island and for the total area formed by the above-
- 281 mentioned polygons. Mean density (n placentae m⁻²) and biomass (kg m⁻²) of
- 282 placentae for the decade were also estimated considering the area of the island at
- 283 mean low water (697102 m^2).

284 Carcass consumption by scavengers

The consumption of carrion by gulls was calculated for the 2012 pupping season. To do so, the estimated percentage of scavenged mass, calculated according to the given consumption state (from A to E), was removed from each carcass mass. The scavenged portion of the total biomass available was then calculated as: starting mass – consumption state mass = scavenged mass.

290 Statistical analysis

We calculated means, standard errors, and relative standard errors (RSE = standard error / mean *100) expressed as a percentage, as measures of inter-annual variability of the different characteristics of the seal carrion resource (mean pupping date, biomass, energy and density of both placentae and carcasses).

We used two-way analysis of variance (ANOVA) to test for the effect of pup status (dead/alive), developmental stage (1-5), and their interaction on carcass mass.

298 We used Ripley's function (Ripley 1977) to analyse inter-annual variation in 299 the spatial pattern and distribution of carcasses. This method is used in 300 epidemiology, but also in forensic science and plant ecology (Lancaster and Downes 301 2004), and provides a visual comparison of the spatial distribution of points 302 belonging to different patterns, as well as the identification of clustering. Considering 303 a circle of radius r and the number of points inside the circle, the Ripley's K function 304 represents the expected number of points within a distance r from an arbitrary point, 305 divided by the intensity of points occurring in the study area. The linearised 306 \hat{L} function is the corresponding transformation of the Ripley's K function and here it is 307 used to facilitate the visualisation (Santos and Schiavetti 2014). As the point patterns 308 of this study varied from location to location, the inhomogeneous *Linhom (r)* function 309 was used as proposed by Baddeley (2008). The three functions for 2008, 2012 and 310 2013 were estimated using the Ripley's isotropic correction for a polygonal window,

311 as an edge correction is needed to reduce bias (Ripley 1988, Baddeley 1998). They 312 were then tested for Complete Spatial Randomness (CSR) by running Monte Carlo 313 permutations (n = 99) for statistical tests (Lancaster and Downes 2004) providing a 314 level of significance of clustering. The latter functions were finally compared by 315 computing a bootstrap 95% confidence band for each function and plotted together 316 in order to evaluate whether the pattern of the functions was repeated in years 317 showing overlaps. We interpreted the observed spatial patterning of carcasses 318 against the null hypothesis of a random distribution of carcasses. This null model 319 was indicated by a horizontal line of null x values between the x and y axes 320 corresponding respectively to the radius r (in metres) and the \hat{L} function values. The 321 function *Linhom (r)* is located above the horizontal line when the spatial pattern of 322 points is aggregated and below it when it is uniform. Spatial analyses were run using 323 R 3.1.1 and the following packages: Spatstat, Maptools, Rgdal, Rgeos, Ggplot2 and 324 Splancs (Baddeley and Turner 2005, Wickham 2009, Bivand and Lewin-Koh 2014, 325 Bivand and Rundel 2014, Bivand et al. 2014).

2.4 Comparison with salmon runs

327 To complement our empirical analyses, we investigated if carried provided by 328 the Isle of May seal colony was a predictable food supply relative to other published 329 studies. We focused on salmon runs during the spawning season as a predictable 330 resource for a wide range of consumers, in particular predatory bears which predict 331 salmon availability (Lugue and Stokes 1976; Fortin et al. 2007). The papers reviewed 332 were Reimchen (2000), Gende et al. (2001), Dickerson et al. (2002), Quinn and 333 Adams (1996), and Hocking and Reimchen (2009) which included data on annual 334 variability of salmon runs.

335

336 3 Results

337 **3.1 Timing of mean pupping dates**

The annual mean pupping date observed during 2000-2010 and 2012 was 30 October (SE = 0.58; n = 12; RSE = 1.97%) (Fig. 2a).

340 3.2 Quantity of seal carrion

Aerial survey data showed that the mean pup production was 1988 (SE = 47.72; n = 12; RSE = 2.40%) pups per year. Therefore, the same mean number of

343 placentae was released yearly. The mean mass of a placenta was 1.6 kg (SD =

- 0.24 (Table 2) and the resulting total biomass for the period 2000-2010 and 2012 was estimated to be 3124.3 kg·yr⁻¹ (SE = 74.99; n = 12; RSE = 2.40%). The mean number of dead pups estimated by aerial surveys was 262 (SE = 16.32; n = 12; RSE = 6.23%), representing a mean annual mortality of 13.3 % (SE = 0.91) (Fig. 2b).
- 348 Ground visual censuses performed in 2008 and 2012 recorded similar 349 numbers of dead seals (adult and pups), with 229 and 233 respectively, whereas in 350 2013 a total of 165 carcasses were observed. The mean number of seal carcasses 351 counted for the three years was 209 (SE = 22.03; n = 3; RSE = 10.54%). On 352 average, more than half the carcasses (55.63%; SE = 4.15) were second stage pups 353 (Fig. 3), aged between 1.7 and 7.9 days. Starvelings formed 8.66% (SE = 3.94) of 354 the total number of carcasses. Dead adults were found every year (2.12% of the total 355 number of carcasses; SE = 0.70), and 12 out of 14 individuals were female.
- 356 There was a significant difference in mass between dead and live pups 357 (ANOVA, $F_{1,115}$ = 34.84, P < 0.001), and between pup stages (ANOVA, $F_{2,114}$ = 46.49, P < 0.001), but there was no significant interaction (ANOVA, $F_{2,114}$ = 0.11, P = 358 359 0.9). Therefore, the average difference in mass of 8.02 kg measured between alive 360 and dead pups for the three developmental stages was used to calculate the 361 biomass contributed by dead pups (Table 3). The mean annual biomass of dead 362 seals (adults and pups) was estimated to be 3768.2 kg (SE = 713.73; n = 3; RSE = 363 18.94%). The second developmental stage contributed the greatest proportion and represented 42.1% (SE = 5.16) while the mass of adult carcasses represented 364 365 14.6% (SE = 5.04) of the total biomass released.

366 3.3 Quality of seal carrion

- Mean gross energy density of placentae was estimated to be 21.8 $MJ \cdot kg^{-1}$ (SD = 1.15). From this the estimated average annual energy delivered as placentae on the Isle of May was 68.1 x 10³ MJ (SE = 1.64; n = 12; RSE = 2.41%).
- The biomass of dead seals was primarily composed of water (58.6%; SE = 1.46). Fat (18.9%, SE = 2.15) and protein (19.9%; SE = 0.62) inputs combined reached 1481.7 kg·yr⁻¹ (SE = 329.43; n = 3; RSE = 22.23%) and were estimated to release 42.4 x 10^3 MJ·yr⁻¹ (SE = 10.42; n = 3; RSE = 24.58%) of energy to the ecosystem. Fat was the largest contributor to the total energy released (67.2%; SE =
- **3**75 **3.12**).

376 3.4 Spatial distribution of seal carrion

- The mean annual placental density on the island was 0.003 placentae m⁻², (SE = 0.00007; n = 12; RSE = 2.33%) while the mean annual placental biomass per area unit was 0.004 kg m⁻² (SE = 0.0001; n = 12; RSE = 2.50%).
- 380 Carcasses found on the Isle of May were distributed according to the main 381 breeding sites, mainly in the north and south of the island. The three years showed 382 similar patterns with most dead pups located above the mean high water line. 383 However, 13.1% (SE = 3.85) of carcasses occurred in the area between mean high 384 water and mean low water (166902 m²). The mean kernel smoothed intensity was estimated to be 0.0003 carcasses m^{-2} (SE = 0.00003; n = 3; RSE = 10.00%) on the 385 whole island for the three years, whereas the mean density of carcasses in the 386 387 polygons was 0.005 carcasses m^{-2} (SE = 0.001). The highest intensity each year was observed in the area between Rona and North Ness, particularly in 2012 when 388 389 the maximum intensity was 0.03 carcasses m⁻² (Fig. 4). The union of polygons 390 characterised by an intensity equal or higher than 0.001 covered 11% of the total 391 area of the island at mean low water. Two thirds of this area was located in the north. 392 The intersection of polygons (areas shared across years) corresponded to 3% of the 393 total area of the island and 28% of the union of polygons (Fig. 5). The biomass per unit area was 0.006 kg m⁻² (SE = 0.001; n = 3; RSE = 16.67%) for the whole island 394 395 and 0.09 kg m⁻² (SE = 0.03, n = 3; RSE = 33.33%) for the area covered by the 396 polygons.

Initially the spatial pattern of carcasses was highly aggregated, but as the value of the inhomogeneous *Linhom (r)* function increased, it became uniform. The overlap of the three curves demonstrated that areas with carcasses were repeated and had a similar distance across years, both in the north and in the south of the island. The 2008 curve, however, showed a faster decrease towards uniformity (Supplementary material Appendix 1 Fig. A1).

403 **3.5 Carcass consumption by scavengers**

The mean cumulative mass loss (% of starting mass) for consecutive consumption states (A-E) observed on experimental carcasses deployed in 2013 was calculated (Table 4). In 2012, the total biomass scavenged was 1032.0 kg representing 21.4% of the total mass available. The energy intake by scavengers was 12.8 x 10³ MJ.

409 **3.6 Comparison with salmon runs**

410 Our results showed that the annual variability in the timing, quantity 411 (expressed as number and mass of carcasses) and quality (expressed as energy) of 412 dead seals and placentae ranged between 1.97% and 24.50% of relative standard 413 errors (RSE). The spatial intensity of biomass for dead seals had the highest value 414 among all (33.33%; Fig. 6). The analysis of published data indicates that variability in 415 the timing of salmon spawning, quantity and biomass per unit of salmon across 416 years ranges between 1.67 and 25% (RSE). Both seal carrion and salmon runs 417 showed the lowest RSE in their timing relative to quantity and density (Fig. 6; 418 Supplementary material Appendix 1 Table A1).

419

420 4 Discussion

421 *Predictability of the carrion resource*

422 Although predictability is a widely used concept in ecology, there has not been 423 a standard definition of resource predictability. In this study, we used relative 424 standard errors (RSE) to standardise different kinds of data (timing, abundance, 425 biomass, energy, density of two separate carrion sources) to compare different 426 studies. We showed that seal carrion had a similar range of RSE as spawning 427 salmon runs. Salmon runs clearly are predictable because brown bears (Ursus 428 arctos, in undisturbed conditions) move towards rivers an average of two weeks 429 before the arrival of salmon (Fortin et al. 2007) to prey on the spawning fish to 430 accumulate fat for hibernation (Hilderbrand et al. 1999). We suggest this range of 431 RSE values could be used as a benchmark for comparing predictability in future 432 studies dealing with other resources and ecosystems.

433 The timing, quantity, quality and spatial distribution of seal carrion were 434 selected as key attributes of the resource that can be expected to have a major 435 influence on the ecological response of its consumers. Quantity and nutrient content 436 of the food were chosen as they correlate with survival, health, reproductive success 437 of animals, but also population and ecosystem dynamics (White 2008; Marcarelli et 438 al. 2011). Moreover, when a resource is predictable in time and space, animals can 439 learn how to predict its future occurrence (e.g. Reimchen 2000; Wittemver et al. 440 2007; Schindler et al. 2013). The animal learning process involves the temporal 441 awareness (when food occurs), which can be promoted by environmental cues

442 (Frederiksen et al. 2004), but also spatial perception (where food occurs) (Sherry et 443 al. 1992). Individual bears, for instance, return to the same river to fish for salmon 444 year after year (Luque and Stokes 1976). Timing and location were therefore the 445 other crucial features used to describe the predictability of a resource. Furthermore, 446 predictability of salmon runs in riparian systems leads to synchronisation between 447 feeding strategy and food input in black bears (Ursus americanus), wolves (Canis 448 lupus) and riverine and coastal mink (Neovison vison) (Ben-David et al. 1997; 449 Reimchen 2000; Darimont et al. 2003). Recently, it has been demonstrated that 450 brown bears (U. arctos) and glaucous-winged gulls (Larus glaucescens) shift their 451 spatial distribution from early to late sockeye salmon (Oncorhynchus nerka) 452 spawning areas according to the precise timing of spawning of different populations 453 (Schindler et al. 2013).

454 The relative influence of the timing, spatial distribution, quantity, and quality of 455 a food resource on the behaviour and learning of its consumers, seems largely 456 unknown. Our comparison between seal carrion and salmon runs showed that timing 457 had lower RSEs than quantity and density. Does this mean that low variability in 458 timing is the most important factor determining resource predictability for scavenging 459 gulls and bears? Or simply, are foragers less responsive to interannual variation in 460 food quantity rather than timing? In Sigler et al. 2004, pre-spawning aggregation of 461 eulachon (Thaleichthys pacificus), a small ocean fish, had seasonal impacts on the 462 abundance of Steller sea lions (Eumetopias jubatus) attending the area for feeding 463 despite the peak in quantity of this resource greatly varies between years (RSE = 464 65%). Also in this case, the timing was more consistent (RSE = 33.33%; values 465 derived from Sigler et al. 2004). There are very few studies giving details on the 466 inter-annual variation of pulsed resources, and none considering all four components 467 of predictability considered here. Thus, it is difficult to understand how animals 468 assess the individual and combined variations of timing, spatial distribution, quantity, 469 and quality before responding to predictable pulses.

Considering absolute values given in studies on other apparently predictable resources, it appeared that sea turtle eggs provide an amount of energy to plants, decomposers and detritivores similar to seal carrion at pupping (91.1 x 10³ MJ on 21 km stretch of beach; Bouchard and Bjorndal 2000). Seabird chick mortality on nesting islands in the Gulf of California, instead, provides up to 10-fold higher mass of carrion per area unit than seals on the Isle of May (Polis and Hurd 1996a, Sánchez-Piñero and Polis 2000). Moreover, a much higher biomass of marine
plankton deposits on the shore of upwelling areas (ca. 4 kg m⁻²; Davenport 1995).

479 Carrion consumption

480 At the end of the breeding season 21.4% of the dead seal biomass available 481 on the Isle of May was scavenged resulting in an estimated energy intake of 12.8 x 482 10³ MJ. Because of low air temperatures, decomposition rate in winter at high 483 latitude is slow (Beasley et al. 2012), and scavenging is likely the main determinant 484 of mass loss by carcasses. Considering that the daily energy required by a great 485 black-backed gull is 2.6 MJ (Camphuysen et al. 1993; Walter and Becker 1997), the 486 total amount of energy released as dead seals and placentae (110.5 x 10^3 MJ) would 487 be equivalent to 42500 daily rations for these birds, which would be enough to feed 488 more than a thousand gulls per day during the duration of the seal pupping season. 489 This is likely an underestimate as the scavenging activity may continue once the seal 490 pupping season has concluded. Incomplete depletion of the carcasses may have 491 occurred because of their high abundance (Reimchen 2000; Wilmers et al. 2003), 492 and/or low abundance of scavengers and availability of alternative food sources 493 (e.g., fishery discards). In fact, it may take six weeks for a single seal pup carcass to 494 be entirely scavenged by the local gull population (Quaggiotto et al. 2016). Placentae 495 are likely to be immediately and fully consumed by gulls because they are easier to 496 handle and 1.6 times more nutritious than seal carcasses. Energy density of 497 placenta has rarely been measured. In this study grey seal placenta contained three 498 times more energy than harp seal *Pagophilus groenlandicus* placenta (4.73 MJ kg⁻¹; 499 Lavigne and Stewart 1979), but similar amounts to that of west Atlantic grey seals 500 (23 MJ kg⁻¹; Yunker et al. 2005). 501 We cannot ascertain here whether gulls predict carrion resource pulses. However, 502 gulls are known to respond to other predictable food sources, such as fishery

discards (Bartumeus et al. 2010; Cama et al. 2012). Gulls at the open sea distribute
 according to fishery vessel density during fish discarding time, with apparent optimal

- 505 foraging strategies (Cama et al. 2012). Seal carrion on the Isle of May might be
- 506 crucially important for scavenging gulls as a predictable food resource, in particular,
- 507 in preparation for winter. This is similar to salmon for many predatory and
- scavenging animals during autumn (Cederholm et al. 1999). Changes in seal carrion
- 509 consumption by gulls may occur in the near future as a result of the fishery discards

510 ban. Their reliance on carcasses, in fact, could increase after discards availability

511 drops. It may also lead to the specialisation of the great black-backed gull in

512 exploiting carrion, as this species is seen regularly scavenging at the grey seal

colony on Sable Island (Canada) (Ronconi et al. 2014) and also on North Rona (N.

514 Scotland) (Twiss et al. 2003).

515 We found that a proportion of dead seals (13%) occurred between mean high 516 and low water. This means that carrion is exposed not only to gulls, but also to 517 marine scavengers (Quaggiotto et al. 2016). However, these carcasses may 518 represent only a proportion of carrion available to the marine system, as other dead 519 seals are likely to have been already transported by tides and waves before the 520 surveys were undertaken. To our knowledge, the effects of predictable carrion from 521 seabird and pinniped (and therefore from the wider ocean) to small areas of the 522 coastal marine environment have been little explored so far (Watts et al. 2011).

523

524 Broader ecosystem effects

525 Repeated and predictable events lead to long term impacts on ecosystems, 526 and can promote lasting changes in the soil, microbial, and plant communities. 527 Stable isotope δ^{15} N signatures of salmon found in plants and invertebrates (Hocking 528 and Reimchen 2009), for instance, persist for decades after the salmon population 529 disappeared (Koyama et al. 2005). Similar effects are also likely to occur at our study 530 site. Moreover, a higher nitrogen enrichment is also expected in plants and soil than 531 at other animal colonies because of the high position that pinnipeds have in the 532 trophic chain (Erskine et al 1998). Since the distribution of seal carcasses was 533 similar across years, hotspots of soil and vegetation enrichment by marine-derived 534 nutrients might be identified from the maps produced here. The relationship between 535 seal carrion distribution and nutrient hotspots at different spatial scales is an 536 interesting avenue for future research.

537

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553 preparation. All authors read and approved the final manuscript.

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- Figure 1. Map of the Isle of May. Maps provided by EDINA Digimap Service,
- 775 <u>http://digimap.edina.ac.uk/roam/os</u>.



Figure 2. a) Mean pupping date for the decade 2000-2010 and 2012 from aerial
survey data (1 = 1st Oct); b) Estimated maximum number of dead pups and total pup
production for the years 2000 – 2010 and 2012. Number of dead pups re-estimated
from aerial survey data is represented by columns and scaled to the y-axis on the left
side. Pup productivity is represented by line with CV (%, ranged between 0.02 and
0.05) and scaled to the y-axis on the right side.



Number of carcasses

- Figure 3. Number of dead pups per year belonging to different sex and
- developmental stages from the ground visual census performed in 2008, 2012 and
- 787 2013: M = male, F = female and N/A = not identified sex; 1 = first, 2 = second, 3 =
- third, 4 = fourth, 5 = fifth developmental stage, starveling = pups who died of
- starvation, n/a = not identified developmental stage and adult.



Figure 4. Kernel smoothed intensity of carcasses and polygons of density \geq 0.001.

- Left: maps of kernel smoothed intensity (n carcasses m^{-2} ; range= 0 0.03) for
- carcasses counted during the ground visual censuses performed in 2008, 2012 and
- 2013. The smoothing bandwidth was selected by likelihood cross-validation. Pixel
- size: 20 x 20 m. Right: maps showing polygons (in green) characterised by intensity
- equal or higher than 0.001 carcasses m^{-2} in 2008, 2012 and 2013.
- 797

Union and intersection of polygons (intensity \geq 0.001)



798

- Figure 5. Union (sum) and intersection (overlap) of polygons of intensity \ge 0.001.
- 800 Union of polygons characterised by intensity equal or higher than 0.001 carcasses
- m^{-2} for any of the three years (2008, 2012 and 2013) (blue), and polygons of areas
- 802 used in all three years (yellow).

803



807 Figure 6. Comparison between RSE of resource availability resulting from the Isle of 808 May seal carrion and spawning salmon. Inter-annual variability, measured as RSE, 809 of carcasses (black) and placentae (red) occurring on the Isle of May during seal 810 pupping seasons 2008, 2012 and 2013 and spawning salmon (orange) according to 811 time (mean date for seal pupping and salmon spawning), quantity (abundance as n 812 and total mass available as kg), quality (energy released as MJ) and space (density as n m⁻² and kg m⁻²). Reviewed papers: (a) Quinn and Adams 1996, (b) Reimchen 813 814 (2000), (c) Gende et al. 2001, (d) Dickerson et al. 2002 and (e) Hocking and 815 Reimchen (2009).

- Table 1. Energy content for developmental stages of grey seal pups, adult female,
- 818 and adult and juvenile male individuals according to energy density calculated per
- 819 body component (%).

Stage	Body component	Component content (%) (Lang et al. 2011; Lidgard et al. 2005)	Energy density (MJ kg ⁻¹)	Energy content (MJ kg ⁻¹)
	water	65.1	0	
1 and 2	fat	9.4	3.8	7.9
	protein	22.6	4.1	
	water	47.3	0	
3	fat	35.5	14.0	16.8
	protein	15.1	2.8	
	water	45.7	0	
4 and 5	fat	38.1	15.0	17.6
	protein	14.4	2.6	
	water	55.2	0	
Adult female	fat	23.9	9.4	12.7
	protein	18.4	3.3	
	water	52.8	0	
Adult or juvenile male	fat	27.3	10.7	13.8
-	protein	17.4	3.1	

Table 2. Grey seal placentae (n = 6) collected showing date, pup sex, total, amnion

822 and placenta mass (kg).

Dato	Pup	Total Wet	Amnion	Placenta	Wet	Dry Matter	Energy
	Sex	Mass	Mass	Mass	Sample		Density
of collection		(kg)	(kg)	(kg)	Mass (kg)	(70)	(MJ kg⁻¹)
29-Oct	F	1.42	0.76	0.66	0.03	15.66	22.165
30-Oct	n/a	1.91	0.46	1.45	0.03	16.96	22.474
31-Oct	n/a	1.46	0.34	1.12	0.03	15.44	22.099
31-Oct	F	1.19	0.64	0.55	0.05	14.88	20.297
01-Nov	М	1.81	0.33	1.48	0.04	17.23	20.328
05-Nov	n/a	1.63	0.36	1.27	0.03	17.20	23.490

823

- Table 3. Re-estimated mass of dead grey seal pups belonging to different
- 826 developmental stages. Each developmental stage is associated with the re-
- 827 estimated body mass (kg) for male and female pups calculated according to the
- 828 equations of Kovacs and Lavigne (1986) and adjusted by the average difference in
- 829 mass measured between alive and dead pups. When sex of pup was unavailable,
- the body mass was calculated using the mean body mass of both sexes.

termined
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Table 4. States of consumption of scavenged carcasses (n = 11) with associated

State of consumption	Description	Mean % mass loss (SE)	n sample
Α	intact	0.4 % (0.49)	15
В	lack of both eyes and occurrence of one opening on the body	10.2 % (2.23)	30
С	additional openings	31.3 % (4.89)	8
D	body appears flat and lacking internal organs; loss of the head; missing bones	41.2 % (2.43)	7
E	remains (only bones and skin)	65.2 % (3.00)	4

834 description and cumulative percentage mass loss.

835

837 Supplementary material - Appendix 1.

Appendix 1 Figure A1. Test for Complete Spatial Randomness on *Linhom* summaryfunctions.

- 841 Comparison among *Linhom* summary functions illustrating spatial point patterns of
- carcasses found in 2008 (light grey, dashed line), 2012 (dark grey, midline ellipsis)
- and 2013 (medium grey, continuous line) during the ground visual census. Bands of
- 844 95% confidence interval were obtained by bootstrap simulations (n = 99). The three
- ⁸⁴⁵ functions were tested for Complete Spatial Randomness (CSR, dashed red line) by
- running Monte Carlo permutations (n = 99) for statistical tests, bands borders
- represent the p-value 0.05.



Two peaks in the aggregation pattern were observed in each year: one around 40-70 m and the other around 150-180 m, showing that the area of carcasses were repeated and had a similar distance across years, both in the north and in the south

- of the island. Around 215-230 m, the pattern became uniform and the function
- curves fell under the line illustrating complete spatial randomness (CSR). As the
- 854 maximum r displayed does not exceed 400 m, aggregations further than this
- distance are not shown. There was no statistical significance for the test of CSR, as
- the observed patterns were outside of the simulation envelopes representing the p-
- value 0.05. This indicated that well defined areas of carrion availability occurred.
- 858 Moreover the bootstrap 95% confidence bands of the functions overlapped
- 859 demonstrating that the spatial pattern of carcasses is consistent across years.

- 862 Appendix 1 Table A1. Comparison between the Isle of May seal carrion availability
- and apparently predictable pulses in resource availability according to time (a),
- 864 quantity (b), quality (c) and space (d).

Time (a)							
Reference	Resource	Mean	SE	RSE	Effect		
Present study	Seal pupping	30 th Oct	0.58	1.97	Scavenging great black-backed gulls		
Quinn and Adams 1996	Salmon spawning	2 nd July	0.58	1.83	n/a		
		Quant	ity (b)				
Reference	Resource	Mean	SE	RSE	Effect		
Present	Dead seals (ground visual census)	n = 209.0 3777.7 kg	22.03 720.87	10.54% 19.08%	Scavenging great black-		
study	Placentae (aerial survey)	n = 1988.1 3124.3 kg	47.72 74.99	2.40% 2.40%	backed guils		
Reimchen 2000	Salmon spawning run	n = 4,000	1011.60	25.00%	Black bears predation		
Gende et al. 2001	Salmon spawning run (Hansen Creek)	n = 13791.3	3281.44	23.80%	Brown and black bears predation		
Dickerson et al. 2002	Salmon spawning run	MaxN = 587.3	119.78	20.40%	Brown bears predation		
		Quali	ty (c)				
Reference	Resource	Mean	SE	RSE	Effect		
	Dead seals (ground	42.4 10 ³ MJ	10.42	0.42 4.58%			
Present	visual census)	42.4 10 103	10.42		Scavenging great black-		
study	Placentae		1.64	2.41%	backed gulls		
	(aerial survey)	68.1 10 ³ MJ					
Space (d)							
Reference	Resource	Mean	SE	RSE	Effect		
	Dead seals (ground	0.005 carc m ^{-2 A}	0.001	20.00%			
Present	visual census)	0.09 kg m ^{-2 A}	0.03	33.33%	Scavenging great black-		
study	Placentae	0.04 plac m ^{-2 B}	0.001	2.50%	backed gulls		
	(aerial survey)	0.07 kg m ^{-2 B}	0.002	2.86%			
siday	Placentae (aerial survey)	0.04 piac m ^{-2 b}	0.001	2.50% 2.86%	Dacked guils		

Gende et al.	Salmon spawning	1.72 colm m ⁻²	0.40	23.68%	Brown and black bears
2001	run	1.73 Sain m ² °	0.40		predation
Hocking and					
Peimchen	Salmon spawning	16.3 kg m ⁻¹	5.00	1 67%	Marine enrichment in
Reinichen	run	10.3 kg 11	5.00	1.07 /0	riparian food webs
2009					·

- 866 Summary table showing results obtained by the present study and other published
- sources. The table is divided horizontally according to the factors explaining
- 868 predictability (time, quantity, quality and space) and by columns according to the
- 869 type of predictable resource, values considered and effect observed in the
- 870 ecosystem. Values in kg are of wet mass.
- A calculated considering polygons of intensity higher than 0.001 (2008 = 55976 m²,

872 **2012 = 32746 m²**, **2013 = 50803 m²**);

- ⁸⁷³ ^B calculated considering polygons of intensity higher than 0.001 (mean of the three
- 874 years, mean = 46508 m²);
- ^c calculated using data for Hansen Creek (8000 m²).