- 1 October 22nd 2015
- 2 Dr. Antonio Uzal
- 3 School of Animal Rural and Environmental Sciences
- 4 Nottingham Trent University
- 5 Southwell NG25 0QF, UK
- 6 Phone +44 (0)115 848 5338
- 7 <u>antonio.uzal@ntu.ac.uk</u>
- 8 RH: Laforge et al. Effects of Density on Foal Survival

# 9 Scale-Dependent Effects of Density and Habitat on Foal Survival

- 10 MICHEL P. LAFORGE,<sup>1</sup> Department of Biology, University of Saskatchewan, 112 Science
- 11 Place, Saskatoon, SK S7N 5E2, Canada
- 12 ANTONIO UZAL,<sup>2,3</sup> Department of Biology, University of Saskatchewan, 112 Science Place,
- 13 Saskatoon, SK S7N 5E2, Canada
- 14 SARAH A. MEDILL, Department of Biology, University of Saskatchewan, 112 Science Place,
- 15 Saskatoon, SK S7N 5E2, Canada
- 16 PHILIP D. MCLOUGHLIN, Department of Biology, University of Saskatchewan, 112 Science
- 17 Place, Saskatoon, SK S7N 5E2, Canada
- 18 **ABSTRACT** Identifying the most appropriate scale to study factors influencing life history is
- 19 important to evolutionary ecology and wildlife management. For example, the scale at which
- 20 density is assessed and explains variation in survival can affect how biologists observe and
- 21 interpret population dynamics, which can influence plans for managing populations. Feral horses
- 22 (Equus ferus caballus) contrast with most ungulates by exhibiting a mating system characterized

<sup>&</sup>lt;sup>1</sup> Present affiliation and address: Department of Animal and Poultry Science, College of Agriculture and Bioresources, University of Saskatchewan, Saskatoon SK, S7N 5A8 Canada

<sup>&</sup>lt;sup>2</sup> Corresponding author's email address: <u>antonio.uzal@ntu.ac.uk</u>

<sup>&</sup>lt;sup>3</sup> Present affiliation and address: School of Animal Rural and Environmental Sciences, Nottingham Trent University, Southwell NG25 0QF, United Kingdom

by female-defense polygyny with persistent, non-territorial breeding groups (bands) and female-23 biased initial (natal) and subsequent (breeding) dispersal. We predicted that for horses, offspring 24 movements coupled with female-biased breeding dispersal would increase the scale at which 25 density best related to juvenile survival compared to species with greater female philopatry. From 26 27 2008 to 2013, we censused the population of feral horses on Sable Island, Canada. We annually computed individual-specific local densities for 442 foals (horses/km<sup>2</sup> in radii of 2.000 m, 4.000 28 m, and 8,000 m fixed to a band's centroid of movements) and whole-island (total) population 29 density, group (band) size, and local access to surface freshwater, which affected movement 30 patterns and selection of vegetation by females. The population of feral horses increased from 31 380 in 2008 to 559 in 2013. Overwinter survival of foals averaged 82.8%. Island-wide density 32 was the most important predictor of foal mortality and was negatively associated with survival, 33 with a lesser negative effect from local density. Increased access to surface freshwater (ponds) 34 35 was an important predictor of foal survival but only at certain scales. Our study emphasizes the relevance of a multi-scale approach when analyzing the response of fitness components to 36 changes in habitat and population processes, which may be influenced by the particular social 37 organization of the species. 38

KEY WORDS density dependence, *Equus ferus caballus*, feral horse, foal, habitat, population
dynamics, Sable Island, scale, survival.

In ecology, scale refers to "...several concepts, including the physical extent of the processes (the 41 42 'range') and the spatial and temporal resolution of the data ('grain')" (Fortin and Dale 2005:9). Scale affects interpretation of ecological processes, including resource selection (Anderson et al. 43 2005, Boyce et al. 2003, Leblond et al. 2011, Laforge et al. 2015b), sexual segregation (Bowyer 44 et al. 1996), and interspecific (Whittaker and Lindzey, 2004) and intraspecific competition 45 (Mayor and Schaefer 2005). It is important to use the most appropriate scale, or at least develop 46 models that bridge across scales, to understand an ecological system (Chave 2013). 47 Juvenile survival decreases with increasing intraspecific competition (population density) 48 in many species of wildlife, including Soay sheep (Ovis aries; Milner et al. 1999), bighorn sheep 49 (Ovis canadensis; Portier et al. 1998), and mallard ducks (Anas platyrhynchos; Gunnarsson et al. 50 2006). In examining how an organism might experience effects of intraspecific competition 51 through exposure to conspecific density, Coulson et al. (1997) reported that it was not at the 52 53 largest spatial scale of population density (total population size) that the greatest amount of variation in juvenile survival was explained for red deer (*Cervus elaphus*). Rather, the most 54 statistically explanatory scale isolated a population substructure of intermediate local scales of 55 individual experience related to spatial heterogeneity in high quality food resources (grasslands). 56 The link between use of grasslands and a female's lifetime reproductive success was 57 subsequently reported to interact with its local experience of density (McLoughlin et al. 2006), 58 and adaptive use of habitat depended on how far the local (as opposed to total) population was 59 from reaching carrying capacity (Fortin et al. 2008). A similar situation was reported for juvenile 60 survival of Atlantic salmon (Salmon solar), where fine-scale local density was most important in 61 explaining juvenile mortality (Einum and Nislow 2005). 62

63 There are many factors that may dictate how and why a life-history trait like density64 dependent juvenile survival may vary with scale. Mating systems and social organization can be

65	important to fine-scale, spatial effects on inbreeding (Chesser 1991) and underpin relationships
66	leading to intraspecific competition (Clutton-Brock 1989) affecting distribution, and hence
67	heterogeneity in population and evolutionary processes (e.g., by affecting reproductive success or
68	sex ratio and dispersal; Dobson 1982, Leturque and Rousset 2004). We might therefore expect
69	social system to play a role in density-dependent phenomena and factor into scale-related
70	questions of density dependence. Little is known, however, about how something like an
71	animal's social system might affect scale dependence in ecological interactions. One reason for
72	this is the paucity of multi-scale, individual-based analyses of population dynamics for
73	continuously distributed organisms (i.e., where each individual possesses a unique perspective of
74	density, as opposed being grouped into a discrete subpopulation).
75	Feral horses (Equus ferus caballus) contrast with most other ungulates by exhibiting a
76	mating system that more resembles that of primates (Cameron et al. 2009), being characterized
77	by year-round, female-defense polygyny that results in non-territorial breeding groups (bands)
78	consisting of a harem of females and 1 or 2 adult males (stallions), and other groups of non-
79	reproducing males (bachelors) and adolescents (Berger 1986, Linklater et al. 2000). In horses,
80	both sexes disperse and will move among bands (Berger 1986, Clutton-Brock 1989), although
81	natal (initial) and secondary (breeding) dispersal from bands is generally female-biased
82	(Marjamäki et al. 2013). Dispersal rate in feral horses on Sable Island, Nova Scotia (Fig. 1), was
83	previously reported to result in compensatory and depensatory density feedbacks (Herrando-
84	Pérez et al. 2012), contingent on the state of individuals and the spatial scale at which density
85	acted on individuals (Marjamäki et al. 2013, van Beest et al. 2014). Density affects the ecology of
86	horses on Sable Island, including behaviors such as density-dependent habitat selection (van
87	Beest et al. 2014, Rozen-Rechels et al. 2015) and dispersal (Marjamäki et al. 2013), and rates of
88	survival and reproduction (with density negatively affecting survival for all age and sex

categories; Richard et al. 2014). Horses on Sable Island provide an opportunity to compare scaledependent population dynamics against what has been reported for ungulates with different social
systems.

Originally introduced sometime in the mid-1700s, Sable Island's feral horses have always 92 93 been free-ranging with minimal interference by humans (Christie 1995). The horses of Sable Island present a markedly distinct genetic structure from other horse breeds but are most closely 94 related to the Nordic breeds of horses and ponies (Plante et al. 2007, Prystupa 2012). Though 95 introduced, the horses are treated by Parks Canada Agency as a naturalized species constituting 96 an important part of the ecosystem. Over the course of our study (2008–2013), total population 97 size increased from 380 individuals to 559, an increase of 47%. Foals dispersed from their natal 98 band 33.4% of the time, with approximately a third of foals (10.2% of all foals) dispersing on 99 their own (without their mother). Natal dispersal distance of female foals over winter averaged 100 101  $3.572 \text{ m} \pm 1.546 \text{ m}$  (SD; Contasti 2011). Female dispersal to different bands was common (30.5%) of mothers). 102

Following Coulson et al. (1997), we examined density feedbacks on juvenile overwinter 103 survival at multiple spatial scales for horses on Sable Island, using an individual-based approach. 104 Horses on Sable Island exist in a natural though simplified system without predation, human 105 interference, or interspecific competition (the horses are the island's only terrestrial mammal). As 106 part of a long-term, individual-based monitoring program for this population (Contasti et al. 107 2012, Marjamäki et al. 2013, Richard et al. 2014, van Beest et al. 2014, Rozen-Rechels et al. 108 2015), we tracked the overwinter fates of all foals alive on the island at the end of each summer 109 (n = 442). We quantified each animal's unique experience of the environment including attributes 110 of social group (band size) and a foal's (and its mother's) access to freshwater and exposure to 111 population density at 4 spatial scales of increasing size (2,000-m, 4,000-m, and 8,000-m buffers 112

around median band centroid locations and total population density). We chose these scales to
reflect different percentiles of mean annual horse movement relating to the intensity of use of
their home ranges.

We generated 2 competing hypotheses for how density may affect foal survival. Our first 116 117 hypothesis was that the scale at which foal mortality would be detected in relation to density would mirror that for red deer fawns in Coulson et al. (1997), where local density in a defined 118 area around the focal individual would best predict foal mortality due to local competition for 119 120 resources. By contrast, social system and trends in philopatry may play an important role in determining the most appropriate scale at which to examine density dependence in population 121 processes, including juvenile survival. Horses on Sable Island show female-biased natal and 122 breeding dispersal (social dispersal), and females with accompanying offspring can and do leave 123 bands in response to increases in band size and conspecific density (Marjamäki et al. 2013). 124 125 Dispersal events largely occur during winter (Marjamäki et al. 2013, Debeffe et al. 2015). Departure of foals with females during dispersal increases foal overwinter survival (Debeffe et al. 126 2015), suggesting a role for dispersal to affect variation in foal survival. Therefore, our alternate 127 hypothesis was that dispersal of females and foals would result in a situation where individuals 128 are better able to exploit resources evenly across the island, leading to population density at 129 larger scales being a better predictor of overwinter survival than measurements taken at smaller 130 scales. 131

#### 132 STUDY AREA

Sable Island National Park Reserve, located 275 km southeast of Halifax, Nova Scotia, Canada ( $43^{\circ} 55' \text{ N}, 60^{\circ} 00' \text{ W}$ ), is a crescent-shaped sand bar home to a population of free-ranging feral horses (Fig. 1). It is approximately 49 km long and 1.25 km in width at its widest point, and is comprised of sandy beaches and vegetated and bare sand dunes  $\geq 30$  m high. The island's climate

was temperate oceanic with cool summers and wet winters. Catling et al. (1984), Freedman et al. 137 138 (2011), and Tissier et al. (2013) detailed the flora present. Sable Island was treeless, and the vegetation community was dominated by American beach grass, or marram (Ammophila 139 *breviligulata*), which occurred throughout most of the vegetated parts of the island. Shrub-140 141 dominated heath (Empetrum nigrum, Juniperus communis, Myrica pensylvanica, Rosa virginiana, Vaccinium angustifolum) was the climax vegetation association found on Sable 142 Island. The west end of the island contained patches of sandwort (Honckenya peploides) and 143 beach pea (Lathyrus japonicus var. maritimus). Several permanent freshwater ponds covering 144 approximately 20 ha were used by horses and confined to western and central areas of the island 145 (Contasti 2011, Contasti et al. 2012). Ephemeral melt- and rain-water ponds occurred on the east 146 half of the island but generally disappeared in summer, and horses on east Sable Island had to 147 excavate drinking holes to access freshwater (Contasti 2011). This environmental gradient 148 149 underlaid density-dependent habitat selection processes with major consequences for population and social dynamics across the length of the island (Marjamäki et al. 2013, Richard et al. 2014, 150 Manning et al. 2015). Aside from a small human presence (researchers and tourists), feral horses 151 were the only terrestrial mammals on the island. 152

## 153 **METHODS**

## 154 Data Collection and GIS Analysis

155 We obtained direct observations of individual horses via systematic ground censuses on Sable

- 156 Island (weekly observations from Jul–Sep), including 21,392 individual observations
- encompassing the entire population (N = 380, 437, 503, 448, 534, and 559 individuals,
- respectively, between 2008 and 2013, including 442 foals alive in Sep 2008 through 2013; counts
- updated from Richard et al. [2014]). Each daily sampling effort included 1 of 7 sections of the
- island, stratified to allow complete coverage of a section in 1 day and roughly complete coverage

of the island in 1 week. We approached horses on foot (which were largely indifferent to our 161 162 presence) and recorded the location of an individual using a hand-held global positioning system (GPS) with location error to within 5 m. We determined the horse's identity (to determine 163 survival status and band membership from year to year) from facial features and other 164 165 distinguishing marks (verified using digital photographs at every sampling event), sex, reproductive status, and group membership. Our photography-based approach to cataloguing 166 each individual is described in Contasti (2011). If a horse was not observed during an entire 167 season, we assumed that it had died. We evaluated whether our censuses were adequate by 168 comparing, for 2010 data, our summer counts of non-foals with that obtained from high-169 resolution aerial photography in January 2010 (prior to births). The latter was flown by aircraft at 170 171 an altitude of approximately 450 m as part of a census of the population of grey seals (Halichoerus grypus) on Sable Island, which also proved excellent for counting horses (Bowen et 172 173 al. 2011). This confirmed that our 2010 census accounted for >99% of the horses present as expected. A previous study (Welsh 1975) reported that horses have small home ranges on Sable 174 Island (2.8 ± 1.2 km<sup>2</sup> [ $\bar{x}$  ± SD], n = 32 bands), which generally range from one end of the island 175 to the other and are compressed along its length. 176

All collection and sampling methods (strictly behavioral observation) were approved 177 178 by the University of Saskatchewan's Animal Research and Ethics Board, under University of Saskatchewan Animal Care Protocol 20090032 and guidance of the Canada Council on Animal 179 Care. Access to Sable Island for this research project was granted by written permission of M. 180 181 Voigt, Canada Coast Guard for years 2008–2012; however, in the year of 2012, as management of the island transitioned from Coast Guard to Parks Canada Agency, we also obtained and 182 followed conditions of a Research License (14668) provided by Parks Canada Agency 2012-183 2014. 184

We used ArcGIS 10.1 (Environmental Systems Research Institute, Redlands, CA) to 185 186 compute conspecific densities and proportion of freshwater in 3 buffers of increasing radius (2,000 m, 4,000 m, and 8,000 m) centered on each band's median centroid location. Buffer radii 187 generally corresponded to the 80th, 95th, and 99th percentile of annual movements of all horses 188 189 and were selected to reflect a continuum of scales. We selected the 99th percentile (8,000-m buffer) to reflect the annual movement rate of nearly all horses in the population. We selected the 190 95th percentile (4,000-m buffer) to eliminate outliers of extraordinary dispersal, and the 2,000-m 191 192 buffer (80th percentile) to approximate the core area used by individuals. Within each buffer, we calculated the density of conspecifics as a function of vegetated area (km<sup>2</sup>; hereafter local 193 density) because horses used beaches primarily for transitory movements or resting. We also 194 calculated the proportion of freshwater in each buffer and total population density over the whole 195 island as a function of horses/km<sup>2</sup> of vegetated area (hereafter total density). We included 196 vegetation availability implicitly in the density term as horses/km<sup>2</sup> of vegetated area; therefore, 197 we did not include any other explicit vegetation terms in our models; therefore we did not include 198 any other explicit vegetation terms in our models. We obtained habitat and vegetation data from 199 200 high-resolution aerial photography and a Light Detection and Ranging (LIDAR) map (2009) classified and ground-truthed by the Applied Geomatics Research Group (AGRG) at Nova Scotia 201 Community College, Middleton, Nova Scotia, Canada. 202

#### 203 Statistical Analyses

Foal overwinter survival was a binary response variable, with 0 representing animals that died between 1 August of the birth year (t) and 15 July of the following year (t + 1), and 1 describing those that survived. We used generalized linear mixed effects models (GLMM) with a binomial error structure in R Version 3.2.1 (R Core Team 2015) to model survival as a function of local density and habitat variables at the 3 local scales measured and total island density. Our maximal

model included as covariates local density and proportion of freshwater (computed at 1 of the 3 209 210 identified scales), band size, and total density. We used Z-scores to compare effect sizes between 211 different parameters and to facilitate convergence of models. We tested for collinearity between predictors at all scales and ensured no pair of variables had a Pearson's correlation coefficient of 212 213 r > |0.7| (Dormann et al. 2013) and that no variable had a variance inflation factor >5 (Menard 1995). Unlike population density, surface freshwater availability did not vary among years 214 because we used 1 map (dated 2009); hence, we could not generate a variable for the whole-215 216 island scale. Marjamäki et al. (2013) noted significant collinearity between local density (at the 8,000-m scale) and distance of overwinter movements, and so we also did not include distance of 217 dispersal (if it occurred) as a covariate in our models. We tested our final models for trends in 218 average residuals versus expected values using binned residuals (Gelman and Hill 2007) with the 219 R package arm (Gelman and Su 2015) because plotting raw residuals versus fitted values are 220 221 generally not useful after logistic regression (we did not detect any serious anomalies). We tested 3 model structures for random effects: band identification (ID), maternal ID, and both (not 222 nested). We were not able to test maternal ID nested within band ID because of dispersal of 223 224 mothers to different bands.

Following Burnham and Anderson (2002), we developed a set of candidate models (Table 225 1) that we evaluated using small sample size adjusted Akaike's Information Criterion (AIC $_c$ ). We 226 employed a hierarchical method of model selection, first determining the optimal random effect 227 structure and then performing model selection on the fixed effects. We tested the 3 random effect 228 structures using the maximal (fully parametrized) model with local density and proportion 229 freshwater modeled at each of our 3 scales using a GLMM using the R package lme4 (Bates et al. 230 2015) and selected the model with the lowest AIC<sub>c</sub> (Zuur et al. 2009), which resulted in our using 231 models with maternal ID only. Once we had determined the most informative random term 232

structure, we ran each model in our candidate model set at each of our 3 scales and ranked each 233 234 model by AIC<sub>c</sub>. We used model-averaging techniques using the R package AICcmodavg (Mazerolle 2015) to generate model-averaged parameter estimates for all models that had an 235  $AIC_{c} < 10$  from the top model at each scale. To evaluate predictive capacity of our final GLMM. 236 237 we adopted the approach of Boyce et al. (2002), which applies to this situation. We evaluated this model using k-fold cross-validation (test-to-training ratio of 5 data subsets), whereby we tested 238 239 predictive capacity of partitioned models against withheld training data using the mean Spearman's rank correlation ( $\bar{r}_s$ ) between training and test data, grouped within 10 bins (Boyce et 240 al. 2002). In a separate analysis, we re-ran all models replacing total island density with year, to 241 determine whether other inter-annual effects besides density may have had an impact on foal 242 survival. 243

## 244 **RESULTS**

Of 442 foals entering their first winter from 2008 to 2013, 82.8% survived to the next census 245 period. A foal's exposure to conspecific density in the year it was born (t) had a larger mean and 246 247 standard deviation at small scales than at large ones (Table 2). Using maternal ID only as a random effect provided the best fit for our models as determined by AIC<sub>c</sub>. Using a model 248 structure with band ID or band ID and maternal ID raised the AIC<sub>c</sub> of our fully parametrized 249 250 model by between 1.6 and 3.0 (see Table S1, available online at www.onlinelibrary.wiley.com). Our results suggested that no single model was clearly superior in describing foal overwinter 251 survival; the top 6 models all had a  $\triangle AIC_c$  from the top model < 2.0 (Table 3). All top models 252 253 included local density and total density. Proportion of freshwater was important in many of the 254 top models, was retained in our best model at the 8,000-m and the 2,000-m scale (Table 3), and enhanced probability of survival; however, the term was not informative in our model-averaged 255 estimates at any scale (Table 4). Both local density and total density negatively affected survival 256

in our models at each of our 3 local scales (Table 4); however, the effect was greater for total
density than for local density at all 3 scales. Band size was uninformative at all scales (Table 4).
The *k*-fold cross-validation values for the best model at the 2,000-m, 4,000-m, and 8,000-m
scales were 0.62, 0.75, and 0.68, respectively.

Models generated using year instead of total island density were better overall predictors of foal overwinter mortality (Table 3). Year as a factor, however, reveals little about the underlying ecological processes in how foal survival changes as a function of year, whereas modeling density allowed us to make meaningful ecological conclusions (we could not include both in our main analysis because of collinearity).

#### 266 **DISCUSSION**

It has become increasingly recognized that animal life-history traits are influenced differentially 267 across scales (Wiens 1989, Levin 1992, Wheatley and Johnson 2009). Conspecific densities at 268 269 larger scales were generally more informative than smaller scales in terms of predicting foal overwinter survival for horses on Sable Island, with total density being most explanatory and 270 negatively associated with survival. Both total density and local densities appeared in our top 271 272 models, suggesting that density at several scales can be important when assessing foal overwinter survival; however, the effect was nearly twice as strong for total island density as for local 273 density. Our study supports previous work that has emphasized the relevance of scale in 274 275 ecological studies, in particular the response of fitness components to changes in habitat quality 276 or population dynamics (Coulson et al. 1997, Johnson et al. 2001, Maurer and Taper 2002, Leblond et al. 2011, Laforge et al. 2015b). 277

Our results contrast with numerous other studies that have investigated scalar processes in life-history traits. Studies of roe deer (*Capreolus capreolus*) reported that fine-scale variation in food resources at the home-range scale have the greatest effect on fawn biomass (Pettorelli et al.

2003) and survival (Pettorelli et al. 2005). Einum and Nislow (2005) reported juvenile survival in
salmon was density-dependent at small scales and overwinter survival in red deer fawns was
reported by Coulson et al. (1997) to be affected by conspecific density at intermediate scales. By
contrast, Vreeland et al. (2004) did not find any effect of home-range scale habitat variation in
fawn survival and suggested that factors at the landscape level may have been important in their
system.

Habitat quality, availability of cover habitat, and predation pressure have been identified 287 as important influences of large herbivore life-history traits for many species (Coulson et al. 288 1997, Milner-Gulland et al. 2000, Nilsen et al. 2004). For horses on Sable Island, habitat quality 289 is relatively homogeneous, and cover habitat (e.g., trees and shrubs) and predation pressure are 290 291 non-existent. Cost of dispersal for horses in our study was therefore likely much less than in many other studies. In their study of salmon, Einum and Nislow (2005) suggest that the 292 293 metabolic costs and risk of predation pressure make dispersal costly in this species, resulting in less dispersal and greater density-dependent mortality at local scales. Similarly, Stuart-Smith et 294 al. (1997) reported that caribou (*Rangifer tarandus*) calf mortality was higher in fragmented 295 296 landscapes where individuals had smaller home ranges, suggesting similar barriers to dispersal were responsible for fine-scale juvenile mortality. 297

Our results suggest that differences in social system and juvenile dispersal ability may play a role in the scale at which density affects juvenile mortality. For example, in red deer, female dispersal is rare (Clutton-Brock and Albon 1989, Albon et al. 1992), which may explain why intermediate scales were clearly best at describing overwinter survival in juvenile red deer (Coulson et al. 1997). This contrasts to the situation on Sable Island, where natal dispersal of female foals overwinter can be several kilometers (Contasti 2011). Female horses are generally accompanied by their offspring when leaving social groups in response to density (Marjamäki et

al. 2013) and we suspect that this process may increase variability in local density effects on theoverwinter survival of accompanying foals.

Availability of freshwater appeared to be a lesser (compared to local density) but still 307 important influence of foal survival across most scales of observation. Heterogeneity in 308 309 freshwater as a resource influences spatial distribution and movement patterns for horses on Sable Island (Rozen-Rechels et al. 2015), other populations of feral horses (Rubenstein 1981, 310 Berger 1986), and other large herbivores such as springbok (Antidorcas marsupialis; Ritter and 311 312 Bednekoff 1995) and African elephants (Loxodonta africana; Chamaillé-Jammes et al. 2007). Freshwater availability was a positive predictor of foal survival, which we suspect is mainly 313 manifest through effects on body condition and resources for lactation in mothers. Contasti et al. 314 315 (2012) noted that body condition in female horses significantly declined from west to east on Sable Island (2008–2010) in accordance with the west to east reduction in availability of surface 316 317 freshwater on the island. Heterogeneity in fixed, high quality resources, like ponds including freshwater and their associated vegetation, may act to explain survival similar to the case in red 318 deer (Coulson et al. 1997). Density-related associations on high-nutrition grasslands (Agrostis-319 320 *Festuca*) were attributed to small-scale influences on red deer calf survival; high-resolution scales were good descriptors of population substructure during summer, when good grazing was 321 distributed heterogeneously. Small-scale effects of density aggregations, which were likely linked 322 to sources of permanent freshwater, did translate directly into density effects on survival in our 323 study, however, not to the extent that larger-scale density effects predicted mortality (Table 4). 324 Many studies reported inter-annual, density-independent processes (or processes interacting

Many studies reported inter-annual, density-independent processes (or processes interacting with density) acting on population dynamics to affect juvenile survival. Spring temperatures have been correlated with juvenile survival in bighorn sheep lambs (Portier et al. 1998) and red deer (Albon et al. 1987). Soay sheep juvenile overwinter survival appears to be driven by a

combination of both density and weather effects (Grenfell et al. 1998, Milner et al. 1999). Our
results suggest a similar situation in horses on Sable Island; models using year instead of total
density improved our models, suggesting density-dependent and density-independent effects are
at play in juvenile survival. It is likely that greater total island densities are associated with
overwinter resource depletion, which is likely exacerbated by extreme weather effects leading to
increased mortality (Manning et al. 2015).

## 335 MANAGEMENT IMPLICATIONS

336 Managers of wide-ranging, long-lived species (e.g., large mammals) may be at particular risk of making prior assumptions about the scale at which population dynamics should be monitored or 337 studied. In light of our findings, a more reliable methodology to analyse the response of fitness 338 components to changes in habitat and population processes (in particular early survival) may 339 involve a multi-scale approach in which an understanding of the social system plays a role. In 340 341 species that exhibit strong dispersal responses such as horses, management efforts should be focussed on broad scales. Using a framework of multiple grains in animal habitat selection 342 studies (Laforge et al. 2015a, b), informed by knowledge of the species' social system, will 343 344 provide valuable insight into selection and dispersal patterns. Following such a protocol will assist in the development of more effective and directed conservation and management plans, 345 which is especially critical for the viability of isolated species and populations. 346

#### 347 ACKNOWLEDGMENTS

We thank A. L. Contasti, J. Poissant, S. E. Simpson, K. R. Lysak, J. A. Weisgerber, and E. J.
Tissier for assisting in fieldwork. L. Debeffe and 2 anonymous reviewers critiqued an earlier
version of this paper. Mapping support was provided with assistance of D. Colville, A. M. Muise,
and the Sable Island Preservation Trust. In-kind and logistical support was provided by Parks
Canada Agency and Fisheries and Oceans Canada including Canada Coast Guard and the

- 353 Bedford Institute of Oceanography, Environment Canada, Maritime Air Charters Limited, and
- 354 Sable Island Station (Meteorological Service of Canada).

## 355 LITERATURE CITED

- Albon, S. D., T. H. Clutton-Brock, and F. E. Guinness. 1987. Early development and population
- 357 dynamics in red deer. II. Density-independent effects and cohort variation. Journal of
  358 Animal Ecology 56:69–81.
- Albon, S. D., H. J. Staines, F. E. Guinness, and T. H. Clutton-Brock. 1992. Density-dependent
  changes in the spacing behaviour of female kin in red deer. Journal of Animal Ecology
  61:131–137.
- Anderson, D. P., M. G. Turner, J. D. Forester, J. Zhu, M. S. Boyce, H. Beyer, and L. Stowell.
- 363 2005. Scale-dependent summer resource selection by reintroduced elk in Wisconsin, USA.
  364 Journal of Wildlife Management 69:298–310.
- Bates D, M. Maechler, B. Bolker and S. Walker. 2015. lme4: linear mixed-effects models using
- Eigen and S4. R package version 1.1-8, <a href="http://CRAN.R-project.org/package=lme4">http://CRAN.R-project.org/package=lme4</a>>.
- 367 Accessed 10 05 2015.
- Berger J. 1986. Wild horses of the Great Basin: social competition and population size.
- 369 University of Chicago Press, Chicago, Illinois, USA.
- Bowen, W. D., C. den Heyer, J. I. McMillan, and M. Hammill. 2011. Pup production at Scotian
- 371 Shelf grey seal (*Halichoerus grypus*) colonies in 2010. Canadian Science Advisory
- 372 Secretariat Research Document 2011/066. Fisheries and Oceans Canada, Ottawa, Canada.
- 373 <a><br/>www.dfo-mpo.gc.ca/csas-sccs>. Accessed 12 May 2014.</a>
- 374 Bowyer, R. T., J. G. Kie, and V. Van Ballenberghe. 1996. Sexual segregation in black-tailed
- deer: effects of scale. Journal of Wildlife Management 60:10–17.

- Boyce, M., J. Mao, E. Merrill, D. Fortin, M. Turner, J. Fryxell, and P. Turchin. 2003. Scale and
  heterogeneity in habitat selection by elk in Yellowstone National Park. EcoScience 10:421–
  431.
- Boyce, M., P. R. Vernier, S. E. Nielsen, and F. K. Schmiegelow. 2002. Evaluating resource
  selection functions. Ecological Modelling 157:281–300.
- Burnham, K. P., and D. R. Anderson. 2002. Model selection and multimodel inference: a
- 382 practical information-theoretic approach. Second edition. Springer, New York, USA.
- Cameron, E. Z., T. H. Setsaas, and W. L. Linklater. 2009. Social bonds between unrelated
- 384 females increase reproductive success in feral horses. Proceedings of the National Academy
- 385 of Sciences of USA 106:13850–13853.
- Catling, P. M., B. Freedman, and Z. Lucas. 1984. The vegetation and phytogeography of Sable
  Island, Nova Scotia. Proceedings of the Nova Scotia Institute of Science 24:181–248.
- 388 Chamaillé-Jammes, S., M. Valeix, and H. Fritz. 2007. Managing heterogeneity in elephant
- 389 distribution: interactions between elephant population density and surface-water
- availability. Journal of Applied Ecology 44:625–633.
- Chave, J. 2013. The problem of pattern and scale in ecology: what have we learned in 20 years?
  Ecology Letters 16:4–16.
- 393 Chesser, R. K. 1991. Gene diversity and female philopatry. Genetics 127:437–447.
- Christie, B. 1995. The horses of Sable Island. Second edition. Pottersfield Press, Lawrencetown
  Beach, Nova Scotia, Canada.
- Clutton-Brock, T. H. 1989. Mammalian mating systems. Proceedings of the Royal Society of
  London B 236:339–372.
- 398 Clutton-Brock, T. H., and S. D. Albon. 1989. Red deer in the highlands. Blackwell Scientific
- 399 Publications, Oxford, United Kingdom.

- 400 Contasti, A. L. 2011. Structure in vital rates, internal source-sink dynamics, and their influence
- 401 on current population expansion for the feral horses (*Equus ferus caballus*) of Sable Island,
  402 Nova Scotia. Thesis, University of Saskatchewan, Saskatoon, Canada.
- 403 Contasti, A. L., E. J. Tissier, J. F. Johnstone, and P. D. McLoughlin. 2012. Explaining spatial
- 404 heterogeneity in population dynamics and genetics from spatial variation in resources for a
- 405 large herbivore. PLOS ONE 7:e47858.
- 406 Coulson, T., S. Albon, F. Guinness, J. Pemberton, and T. Clutton-Brock. 1997. Population
- 407 substructure, local density, and calf winter survival in red deer (*Cervus elaphus*). Ecology
  408 78:852–863.
- Debeffe, L., E. Richard, S. A. Medill, J. A. Weisgerber, and P. D. McLoughlin. 2015. Costs of
  social dispersal in a polygynous mammal. Behavioral Ecology:in press.
- 411 Dobson, F. S. 1982. Competition for mates and predominant juvenile male dispersal in mammals.
  412 Animal Behaviour 30:1183–1192.
- 413 Dormann, C. F., J. Elith, S. Bacher, C. Buchmann, G. Carl, G. Carré, J. R. García Marquéz, B.
- 414 Gruber, B. Lafourcade, P. J. Leitão, T. Münkemüller, C. McClean, P. E. Osborne, B.
- 415 Reineking, B. Schröder, A. K. Skidmore, D. Zurell, and S. Lautenbach. 2013. Collinearity:
- 416 a review of methods to deal with it and a simulation study evaluating their performance.
- 417 Ecography 36:27–46.
- 418 Einum, S., and K. H. Nislow. 2005. Local-scale density-dependent survival of mobile organisms
- 419 in continuous habitats: an experimental test using Atlantic salmon. Oecologia 143:203–210.
- 420 Fortin M.-J., and M. R. T. Dale 2005. Spatial analysis: a guide for ecologists. Cambridge
- 421 University Press, Cambridge, United Kingdom.

- Fortin, D., D. Morris, and P. D. McLoughlin. 2008. Adaptive habitat selection and the evolution
  of specialists in heterogeneous environments. Israel Journal of Ecology and Evolution
  54:311–328.
- 425 Freedman, W., P. M. Catling, and Z. Lucas. 2011. Effects of feral horses on vegetation of Sable
- 426 Island, Nova Scotia. Canadian Field-Naturalist 125:200–212.
- 427 Gelman, A., and J. Hill. 2007. Data analysis using regression and multilevel/hierarchical models.
  428 Cambridge University Press, Cambridge, United Kingdom.
- 429 Gelman, A. and Y.-S. Su. 2015. arm: data analysis using regression and multilevel/hierarchical
- 430 models. R package version 1.8-6. <<u>http://CRAN.R-project.org/package=arm</u>>. Accessed 10
  431 05 2015.
- 432 Grenfell, B. T., K. Wilson, B. F. Finkenstädt, T. N. Coulson, S. Murray, S. D. Albon, J. M.
- Pemberton, T. H. Clutton-Brock, and M. J. Crawley. 1998. Noise and determinism in
  synchronised sheep dynamics. Nature 394:674–677.
- 435 Gunnarsson, G., J. Elmberg, K. Sjöberg, H. Pöysä, and P. Nummi. 2006. Experimental evidence
- 436 for density-dependent survival in mallard (*Anas platyrhynchos*) ducklings. Oecologia
  437 149:203–213.
- Herrando-Pérez, S., S. Delean, B. W. Brook, and C. J. A. Bradshaw. 2012. Density dependence:
  an ecological Tower of Babel. Oecologia 170:585–603.
- 440 Johnson, C. J., K. L. Parker, and D. C. Heard. 2001. Foraging across a variable landscape:
- behavioural decisions made by woodland caribou at multiple spatial scales. Oeclogia
  127:590–602.
- 443 Laforge, M.P., R.K. Brook, F.M. van Beest, E.M. Bayne, and P.D. McLoughlin. 2015a. Grain-
- 444 dependent Functional Responses in Habitat Selection. Landscape Ecol., in press.

445	Laforge, M. P., E. Vander Wal, R. K. Brook, E. M. Bayne and P. D. McLoughlin. 2015b.
446	Process-focussed, multi-grain resource selection functions. Ecological Modelling 305:10-
447	21.
448	Leblond, M., J. Frair, D. Fortin, C. Dussault, JP. Ouellet, and R. Courtois. 2011. Assessing the
449	influence of resource covariates at multiple spatial scales: an application to forest-dwelling
450	caribou faced with intensive human activity. Landscape Ecology 26:1433–1446.
451	Leturque, H., and F. Rousset. 2004. Intersexual competition as an explanation for sex-ratio and
452	dispersal biases in polygynous species. Evolution 58:2398–2408.
453	Levin, S. A., 1992. The problem of pattern and scale in ecology: the Robert H. MacArthur award
454	lecture. Ecology 73:1943–1967.
455	Linklater, W. L., E. Z. Cameron, K. J. Stafford, and C. J. Veltman. 2000. Social and spatial
456	structure and range use by Kaimanawa wild horses (Equus caballus: Equidae). New
457	Zealand Journal of Ecology 24:139–152.
458	Manning, J. A., S. A. Medill, and P. D. McLoughlin. 2015. Climate fluctuations interact with
459	local demography and resources to predict spatially dynamic adult sex ratios in a
460	megaherbivore. Oikos 124:1132–1141.
461	Marjamäki, P. H., A. L. Contasti, T. N. Coulson, and P. D. McLoughlin. 2013. Local density and
462	group size interacts with age and sex to determine direction and rate of social dispersal in a
463	polygynous mammal. Ecology and Evolution 3:3073–3082.
464	Maurer, B. A, and M. L. Taper. 2002. Connecting geographical distributions with population
465	processes. Ecology Letters 5:223–231.
466	Mayor, S. J., and J. A. Schaefer. 2005. The many faces of population density. Oecologia
467	145:276–281.

468	Mazerolle	, M. J. 2015	. AICcmodavg:	model selection an	nd multimodel	inference based on
-----	-----------	--------------	---------------	--------------------	---------------	--------------------

- 469 (Q)AIC(c). R package version 2.0-3. <a href="http://CRAN.R-project.org/package=AICcmodavg">http://CRAN.R-project.org/package=AICcmodavg</a>>.
  470 Accessed 10 05 2015.
- 471 McLoughlin, P. D., M. S. Boyce, T. Coulson, and T. Clutton-Brock. 2006. Lifetime reproductive
- 472 success and density-dependent, multi-variable resource selection. Proceedings of the Royal
- 473 Society of London B 273:1449–1454.
- 474 Menard, S. 1995. Applied logistic regression analysis: Sage University series on quantitative
  475 applications in the social sciences. Sage, Thousand Oaks, California, USA.
- 476 Milner, J. M., D. A. Elston, and S. D. Albon. 1999. Estimating the contributions of population
- density and climatic fluctuations to interannual variation in survival of Soay sheep. Journal
  of Animal Ecology 68:1235–1247.
- 479 Milner-Gulland, E. J., T. N. Coulson, and T. H. Clutton-Brock. 2000. On harvesting a structured
  480 ungulate population. Oikos 88:592–602.
- 481 Nilsen, E. B., J. D. C. Linnell, and R. Andersen. 2004. Individual access to preferred habitat
- 482 affects fitness components in female roe deer *Capreolus capreolus*. Journal of Animal
  483 Ecology 73:44–50.
- 484 Pettorelli, N., S. Dray, J.-M. Gaillard, D. Chessel, P. Duncan, A. Illius, N. Guillon, F. Klein, and
- G. Van Laere. 2003. Spatial variation in springtime food resources influences the winter
  body mass of roe deer fawns. Oecologia 137:363–369.
- 487 Pettorelli, N., J.-M. Gaillard, N. G. Yoccoz, P. Duncan, D. Maillard, D. Delorme, G. Van Laere,
- 488 and C. Toïgo. 2005. The response of fawn survival to changes in habitat quality varies
- 489 according to cohort quality and spatial scale. Journal of Animal Ecology 74:972–981.

490	Plante, Y., J. L. Vega-Pla, Z. Lucas, D. Colling, B. De March, and F. Buchanan. 2007. Genetic
491	diversity in a feral horse population from Sable Island, Canada. Journal of Heredity
492	98:594–602.
493	Portier, C., M. Festa-Bianchet, JM. Gaillard, J. T. Jorgenson, and N. G. Yoccoz. 1998. Effects
494	of density and weather on survival of bighorn sheep lambs (Ovis canadensis). Journal of
495	Zoology 245:271–278.
496	Prystupa, J. M., R. Juras, E. G. Cothran, F. C. Buchanan, and Y. Plante. 2012. Genetic diversity
497	and admixture among Canadian, Mountain and Moorland and Nordic pony populations.
498	Animal 6:19–30.
499	R Core Team. 2015. R: a language and environment for statistical computing. R Foundation for
500	Statistical Computing, Vienna, Austria.
501	Richard, E., S. E. Simpson, S. A. Medill, and P. D. McLoughlin. 2014. Interacting effects of age,
502	density, and weather on survival and current reproduction for a large mammal. Ecology and
503	Evolution 4:3851–3860.
504	Ritter, R. C., and P. A. Bednekoff. 1995. Dry season water, female movements and male
505	territoriality in springbok: preliminary evidence of waterhole-directed sexual selection.
506	African Journal of Ecology 33:395–404.
507	Rozen-Rechels, D., F. M. van Beest, E. Richard, A. Uzal, S. A. Medill, and P. D. McLoughlin.
508	2015. Density-dependent, central-place foraging in a grazing herbivore: competition and
509	tradeoffs in time allocation near water. Oikos 124:1142–1150.
510	Rubenstein, D. I. 1981. Behavioural ecology of island feral horses. Equine Veterinary Journal
511	13:27–34.

512	Stuart-Smith, A. K., C. J. A. Bradshaw, S. Boutin, D. M. Hebert, and A. B. Rippin. 1997.
513	Woodland caribou relative to landscape patterns in northeastern Alberta. Journal of
514	Wildlife Management 61:622–633.
515	Tissier, E., P. D. McLoughlin, J. Sheard, and J. F. Johnstone. 2013. Distribution of vegetation
516	along environmental gradients on Sable Island, Nova Scotia. Écoscience 20:361-372.
517	van Beest, F. M., A. Uzal, E. Vander Wal, M. P. Laforge, A. L. Contasti, D. Colville, and P. D.
518	McLoughlin. 2014. Increasing density leads to generalization in both coarse grained habitat
519	selection and fine-grained resource selection in a large mammal. Journal of Animal
520	Ecology 83:147–156.
521	Vreeland, J. K., D. R. Diefenbach, and B. D. Wallingford. 2004. Survival rates, mortality causes,
522	and habitats of Pennsylvania white-tailed deer fawns. Wildlife Society Bulletin 32:542-
523	553.
524	Welsh, D. A. 1975. Population, behavioural and grazing ecology of the horses of Sable Island.
525	Dissertation, Dalhousie University, Halifax, Canada.
526	Wheatley, M., and C. Johnson. 2009. Factors limiting our understanding of ecological scale.
527	Ecological Complexity 6:150–159.
528	Whittaker, D. G., and F. G. Lindzey. 2004. Habitat use patterns of sympatric deer species on
529	Rocky Mountain Arsenal, Colorado. Wildlife Society Bulletin 32:1114–1123.
530	Wiens, J. A. 1989. Spatial scaling in ecology. Functional Ecology 3:385–397.
531	Zuur, A., E. N. Ieno, N. Walker, A. A. Saveliev, and G. M. Smith. 2009. Mixed effects models
532	and extensions in ecology with R. Springer, New York, USA.
533	Associate Editor: Fanie Pelletier.

## 534 Figure Caption

- 535 Figure 1. Sable Island, Nova Scotia, Canada. Sable Island National Park Reserve (43° 55' N; 60°
- 536 00' W), located approximately 275 km east-southeast of Halifax, Nova Scotia, Canada. The
- island is a crescent shaped sand bar 49 km long and 1.25 km at its widest (vegetation in grey).

538

- Table 1: List of a priori candidate models for linear regression analysis of foal survival, Sable
- 540 Island, Nova Scotia, Canada, 2008–2013. Each model was evaluated with local density and
- 541 proportion water computed at each of 3 scales: 2,000 m, 4,000 m, and 8,000 m.

Local density + total density + proportion freshwater + band size Local density + total density + band size Local density + total density + proportion freshwater Local density + total density Local density Total density Local density + proportion freshwater Null

Table 2: Distribution of density (horses/km<sup>2</sup> of vegetated area) in summer of birth across 4 spatial scales for foals born into bands of

feral horses on Sable Island, Nova Scotia, Canada, 2008–2013. Presented is the mean  $(\bar{x})$ , standard deviation (SD), and coefficient of

544 variation  $(c_v)$  for increasing scales of individual experience, each metric centered on the median centroid of a band's movements in the

545 year (*t*) in which a foal was born. Annual means at the whole-island scale (total) are total number of horses on the island that year

546 divided by vegetated area on the island (without SD or  $c_v$ ).



			Scale (m)	)	
Year		2,000	4,000	8,000	Total
2008 ( <i>n</i> = 69)	x	28.2	14.7	7.7	12.7
	SD	17.0	9.0	5.3	
	$C_{V}$	0.60	0.61	0.69	
2009 ( <i>n</i> = 62)	$\bar{x}$	42.4	20.3	10.2	14.6
	SD	31.2	12.9	6.9	
	$C_V$	0.74	0.63	0.68	
2010 ( <i>n</i> = 83)	$\bar{x}$	37.1	21.4	12.1	16.8
	SD	16.4	13.1	9.2	

23	Laforge	et	al.
----	---------	----	-----

	$C_{V}$	0.44	0.61	0.76	
2011 ( <i>n</i> = 56)	$\bar{x}$	41.7	24.9	15.9	15
	SD	31.6	21.9	16.5	
	$C_V$	0.76	0.88	1.03	
2012 ( <i>n</i> = 93)	$\bar{x}$	37.1	20.6	11	17.8
	SD	13.8	10.9	6.7	
	$C_V$	0.37	0.53	0.61	
2013 ( <i>n</i> = 79)	$\bar{x}$	45.5	25.7	14.1	18.7
	SD	19.7	13.1	7.3	
	$C_V$	0.43	0.51	0.52	
All years $(n = 442)$	$\bar{x}$	38.6	33.9	29.8	16.2
	SD	22.2	16.5	12.3	2.1
	$C_{V}$	0.57	0.49	0.41	0.13

Table 3: Scale, degrees of freedom (df), adjusted Akaike's Information Criterion (AIC<sub>c</sub>), log

549 likelihood (LL) and  $R^2$  for the top models describing foal overwinter survival on Sable Island,

550 Nova Scotia, Canada, 2008–2013. Scale refers to the size of the buffer around which we

quantified local density and proportion of freshwater. For comparison, we also present in the 2

last columns AIC<sub>c</sub> and  $R^2$  values for models with year (factorial) in the model as opposed to total

553 island density.

Model	Scale (m)	df	AIC <sub>c</sub>	LL	$R^2$	Year AIC <sub>c</sub>
Local density + total density (year) +	8,000	5	391.08	-190.47	0.112	387.60
proportion freshwater						
Local density + total density (year)	8,000	4	391.23	-191.57	0.100	387.24
Local density + total density (year) +	2,000	5	391.23	-190.55	0.112	387.93
proportion freshwater						
Local density + total density (year)	2,000	4	391.82	-191.87	0.095	388.54
Local density + total density (year) +	8,000	6	392.93	-190.37	0.113	389.52
proportion freshwater + band size						
Local density + total density (year)	4,000	4	392.93	-192.42	0.093	389.20
Local density + total density (year) + band	8,000	5	393.15	-191.51	0.101	389.19
size						
Local density + total density (year) +	2,000	6	393.21	-190.51	0.113	389.99
proportion freshwater + band size						
Local density + total density (year) + band	2,000	5	393.82	-191.84	0.096	390.61
size						
Null		2	406.75	-201.36	0.000	406.75

Table 4. Model-averaged coefficients and 95% confidence intervals of models with difference in adjusted Akaike's Information Criterion ( $\Delta AIC_c$ ) <10 from the top model at each scale of observation for foal overwinter survival (*t* to *t* + 1) for foals of feral horses on Sable Island, Nova Scotia, Canada, 2008–2013. Density was measured as horses/km<sup>2</sup> of vegetated area. Scale refers to the size of the buffer around which we quantified local density and proportion of freshwater.

Scale (m)	Variable	В	95% CI <sup>a</sup>		
			Lower	Upper	
2,000	Intercept*	1.92	1.49	2.36	
	Local density*	-0.32	-0.60	-0.03	
	Island density*	-0.55	-0.87	-0.23	
	Proportion freshwater	0.25	-0.06	0.57	
	Band size	-0.04	-0.32	0.24	
4,000	Intercept*	1.90	1.47	2.33	
	Local density	-0.28	-0.57	0.02	
	Island density*	-0.55	-0.86	-0.23	
	Proportion freshwater	0.14	-0.16	0.44	
	Band size	-0.05	-0.32	0.23	
8,000	Intercept*	1.89	1.47	2.31	
	Local density*	-0.33	-0.61	-0.05	
	Island density*	-0.52	-0.84	-0.20	
	Proportion freshwater	0.22	-0.08	0.52	
	Band size	-0.06	-0.33	0.22	

<sup>&</sup>lt;sup>\*</sup> Informative parameters (CIs not overlapping 0).