

# Density-dependent habitat selection alters drivers of population distribution in northern Yellowstone elk

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## Abstract

Although it is well established that density dependence drives changes in organismal abundance over time, relatively little is known about how density dependence affects variation in abundance over space. We tested the hypothesis that spatial trade-offs between food and safety can change the drivers of population distribution, caused by opposing patterns of density-dependent habitat selection (DDHS) that are predicted by the multidimensional ideal free distribution. We addressed this using winter aerial survey data of northern Yellowstone elk (*Cervus canadensis*) spanning four decades. Supporting our hypothesis, we found positive DDHS for food (herbaceous biomass) and negative DDHS for safety (openness and roughness), such that the primary driver of habitat selection switched from food to safety as elk density decreased from 9.3 to 2.0 elk/km<sup>2</sup>. Our results demonstrate how population density can drive landscape-level shifts in population distribution, confounding habitat selection inference and prediction and potentially affecting community-level interactions.

## KEY WORDS

cougar, density dependence, food-safety tradeoff, habitat selection, ideal free distribution, predation risk, predator-prey interactions, RSF, spatial distribution, wolf

## INTRODUCTION

Density dependence is a pervasive ecological process, and incorporating it into models of population abundance is critical for understanding population dynamics and informing management (Abadi et al., 2012; Guthery & Shaw, 2013). Whereas much is known about how density dependence affects variation in abundance over time, relatively little is known about how it affects abundance over space. This is a particular problem for the study of habitat selection, where density-dependent habitat selection (DDHS) is a foundational assumption (Rosenzweig, 1981) that is rarely tested and often ignored (Avgar et al., 2020). This gap is significant because unmeasured density-dependent variation in habitat selection may limit the accuracy of empirical models

for inferring drivers of fitness (e.g. food vs. safety) and predicting spatial distribution and abundance (Boyce & McDonald, 1999; Matthiopoulos et al., 2015, 2019).

The expectation of DDHS arises from optimal foraging theory (MacArthur & Pianka, 1966) via the ideal free distribution (IFD; Fretwell & Lucas, 1969). The IFD postulates that a population's density influences the fitness benefits that individuals receive from a habitat (Morris, 1987), and that individuals use habitats in a way that equalises fitness across occupied habitats (Fretwell & Lucas, 1969; Křivan et al., 2008). According to IFD, as population density increases, individuals occupy progressively lower-quality habitats ('spill-over'), resulting in 'negative DDHS'—the strength of selection for high-quality habitat decreases with density (Morris, 2003; Rosenzweig, 1991). Alternatively, fitness

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in certain habitats can increase with population density (Stephens & Sutherland, 1999), leading to an increase in selection strength for these habitats—‘positive DDHS’ (Morris, 2002). Positive DDHS manifests as habitat switching, where individuals leave habitats associated with high fitness at a low density, shifting into habitats associated with high fitness at a high density (Greene & Stamps, 2001).

Fitness is never determined by a single environmental driver; thus, habitat selection reflects a balancing act along multiple dimensions and scales, each with its own context-dependent relative contribution to overall fitness (multidimensional IFD sensu Avgar et al., 2020). Population density is one of these contexts, and whether fitness in a habitat decreases or increases with density varies across different habitat dimensions. For example, fitness typically decreases with density due to reduced per-capita food acquisition (Le Bourlot et al., 2014), whereas it typically increases with density due to increased safety (i.e. reduced per-capita predation risk; reviewed by Lehtonen & Jaatinen, 2016). If food and safety are spatially independent, we expect negative DDHS along both dimensions: at low density, individuals select habitats with more food and safety, but as density increases, individuals spill into habitats with less food or safety. Conversely, if food and safety are negatively correlated in space, individuals select habitats with more food *or* safety. Thus, we expect negative DDHS along the safety dimension and positive DDHS along the food dimension; as density increases, safety in numbers increases and simultaneously, increasing intraspecific competition makes food more limiting to fitness, hence a stronger driver of habitat selection. (Avgar et al., 2020).

Despite the theoretical and practical importance of DDHS, empirical understanding of DDHS and its effects on inference and prediction from habitat selection models is underdeveloped (Avgar et al., 2020). This is especially true in free-living systems (McLoughlin et al., 2010), which involve complex trade-offs (e.g. food for safety) that are often missing in experimental systems (Lima & Dill, 1990), habitat selection at multiple scales (Johnson, 1980) and multiple interacting species. To fill this gap, we tested the hypothesis that trade-offs between food and safety generate positive DDHS for food resources and negative DDHS for safe habitats, such that the drivers of habitat selection switch from food to safety as density decreases. We did so by constructing and applying a population-level habitat selection model to winter aerial-survey data of northern Yellowstone elk collected over 16 years spanning 4 decades. Our findings demonstrate how ignoring DDHS can confound both ecological inference and prediction. We provide novel evidence of multidimensional IFD and of DDHS as an important driver of habitat selection and spatial distribution in a free-living system.

## METHODS

### Study area

Our study occurred in the winter range of the northern Yellowstone elk population (Houston, 1982; Lemke & Mack, 1998). We expanded the area previously used to define the winter range (e.g. Tallian et al., 2017) to include adjacent areas where elk were also occasionally counted (Figure S1). We believe the modified polygon better captures the full extent used by this population. Our study area encompassed 1900 km<sup>2</sup>, compared with the 1520 km<sup>2</sup> previous authors have cited. Approximately two-thirds of the winter range falls within the boundaries of Yellowstone National Park (YNP), with the remaining one-third to the north in the state of Montana. Elevations range between 1500 and 3000 m, and the area experiences long, cold winters (Houston, 1982). Northern Yellowstone elk migrate from high-elevation summer ranges, often in the interior of YNP, to the lower elevation winter range, where they are found from December to April (Houston, 1982; White et al., 2010).

Wolves (*Canis lupus*) and cougars (*Puma concolor*) are the two main predators of northern Yellowstone elk during winter (Kohl et al., 2019). Elk comprised 96% of the wolf diet in winter from 1995 to 2009 (Metz et al., 2012) and 75% of the cougar diet from 1998 to 2005 (Ruth et al., 2019). Alternative prey for wolves and cougars in the system include bison (*Bison bison*), deer (*Odocoileus hemionus*, *O. virginianus*), moose (*Alces alces*), pronghorn (*Antilocapra americana*) and bighorn sheep (*Ovis canadensis*) (Metz et al., 2012; Ruth et al., 2019).

### Data collection

A timeseries of winter counts of northern Yellowstone elk extends to the 1920s, and the population has fluctuated with various management practices and climatic conditions throughout it (MacNulty et al., 2020). Since 1988, these data have included georeferenced locations of elk groups in many years, allowing us to estimate habitat selection. Elk were counted via aerial, fixed-wing surveys designed to provide a full population census. Surveys took place mainly between 08:00 and 12:00 over 1 and 4 days between late December and March of each winter (Appendix S1). In some years, the count did not occur or georeferenced group data were only available within YNP. We used a state-space model (see Appendix S1 in Tallian et al., 2017) to interpolate total abundance in years with incomplete data so that we could predict using our fitted habitat selection model. We used the partial georeferenced data for validation (see *Model evaluation*). Counts reached an all-time high in 1994 near 20,000 elk and a low of less than 4000 in 2013 (MacNulty et al., 2020). Thus, this timeseries of georeferenced counts provided information about elk

distribution across a wide range of population densities, ideal for measuring DDHS.

During the study period (1988–2020), densities of wolves and cougars also varied in the study area. Wolves were reintroduced to YNP during 1995–1997 (Bangs & Fritts, 1996), and their population increased to a maximum in 2003 and declined thereafter (Smith et al., 2020). Cougar densities generally increased across the study period (Marcus et al., 2022; Ruth et al., 2019). Consequently, the risk of predation by each of these predators has varied substantially across the timeseries of elk distribution analysed here.

## Model structure

We modelled elk counts ( $n_{i,t}$ ) within a pixel ( $i$ ) in a year ( $t$ ) using a Bayesian generalised linear mixed model (GLMM). We used 1 km<sup>2</sup> pixels to average over elk daily movements (Appendix S1). We did not have data to directly account for imperfect detection, but we found our results were insensitive to it (Appendix S2). We used a negative-binomial likelihood to accommodate overdispersion in our count data, a distribution commonly used to model animal group sizes (Ma et al., 2011). Factors other than habitat, for example, sex-specific social interactions, are at least partially responsible for group size distributions (Gerard et al., 2002), which results in overdispersion that should be modelled to avoid overfitting. Elk group sizes in our dataset range from 1 to over 1000 individuals, as seen with elk in other areas (Brennan et al., 2015; Proffitt et al., 2012).

We modelled the expected count ( $\lambda_{i,t} = E[n_{i,t}]$ ) in each pixel and year as a function of 22 covariates (indexed by  $k$ ; Table S1), a time-varying offset ( $\alpha_t$ ), a temporal random effect ( $\eta_{i,t}$ ) and a spatial random effect ( $s_i$ , equation 1), all of which we describe in detail in Appendix S3. We used the natural logarithm (hereafter, log) as the link function.

$$\log(\lambda_{i,t}) = \alpha_t + \eta_{i,t} + s_i + \sum_{k=1}^K \beta_k x_{i,t,k} \quad (1)$$

We estimated the size parameter of the negative binomial distribution as a single free parameter ( $r$ ) such that:  $n_{i,t} \sim \text{NegBin}\left(\frac{r}{r+\lambda_{i,t}}, r\right)$ .

Because we set the area of each pixel to 1.0 km<sup>2</sup>, we can interpret  $\lambda_{i,t}$  as expected density in elk/km<sup>2</sup>. Note that ‘expected density’ refers to the expected value of our response variable. Contrast this with ‘(log) average range-wide density’, which is a predictor variable, hereafter  $\log(\text{Dens})$ .

## Habitat variables

We treated habitat variables as fixed effects representing food, safety, or other conditions. The slope of each

variable measures habitat selection (see Appendix S4). We rasterised all count data and variables on a 1-km grid, retaining only pixels falling within the study area ( $N = 1978$ ). We describe processing for all variables in Appendix S5. Variables describing conditions—snow-water equivalent (SWE), elevation and cosine and sine of aspect (northing and easting respectively; Table S1)—were included to control for important known drivers of elk density but were not the target of inference.

Because northern Yellowstone elk are primarily grazers (Houston, 1982), we measured food using total herbaceous biomass from the Rangeland Analysis Platform (RAP) annual biomass (v2.0) layer (Jones et al., 2021; Robinson et al., 2019). This layer combines information about total growth from 16-day NDVI data with plant functional type estimates to calculate growth of grasses and forbs (the annual layer sums all 16-day layers). For each winter, we used the layer corresponding to the preceding growing season to measure potential forage availability for elk that winter. We log-transformed this covariate to reduce the influence of very high values from agriculture outside YNP. We refer to this as the food variable.

We measured predation risk using the risky places approach such that habitat covariates indexed predation risk (Moll et al., 2017). Previous research has shown that risk to elk from wolves and cougars varies with tree canopy openness and terrain roughness. The wolf habitat domain is characterised by high openness and low roughness, whereas the cougar habitat domain is characterised by lower openness and high roughness (Kohl et al., 2019). On a fine temporal (5 h) and spatial scale (30m), Kohl et al. (2019) showed that individual elk manage risk from wolves and cougars by moving into each predator's habitat domain at the time when that predator is least active. At the coarser scale of our analysis, we expected elk to select 1-km<sup>2</sup> pixels with a mixture of intermediate levels of openness and roughness that facilitate efficient switching between wolf and cougar habitat domains across the diel cycle. Thus, we expected elk density to be greatest at intermediate values of these variables, which indicate high heterogeneity in the pixel (Appendix S1; Figure S4). To test this, we included linear and quadratic terms for openness and roughness to parameterise a parabola that quantified how elk select for safety at the 1-km<sup>2</sup> scale. Hereafter, we refer to these as the safety variables. To check the raw data for a negative correlation between food and safety, we created a composite safety variable by taking the product of the openness and roughness rasters. We used Spearman's correlation to check this assumption. All other food-safety comparisons were model based (see Quantifying DDHS).

To support our assertion that the safety variables reflect how elk perceive risk, we included interactions between wolf and cougar densities and each of the safety variables. If the safety variables were good metrics of safe and risky places, we expected predator densities

to alter the strength of selection for the safety variable or shift the parabola vertex (most preferred openness/roughness). We predicted increasing wolf density would decrease selection strength or push the vertex away from open and smooth (i.e. away from the wolf and into the cougar habitat domain). Conversely, we predicted increasing cougar density would increase selection strength or push the vertex towards open and smooth habitat. Whereas aerial surveys occurred mostly during daylight hours before noon when wolves were more active than cougars (Kohl et al., 2019), we did not expect an effect of survey timing due to our coarse spatial scale (Appendix S1).

To measure DDHS, we included an interaction between the food/safety variables and  $\log(\text{Dens})$ . These interactions allowed for flexibility in the patterns of DDHS, but due to the linear and quadratic terms involved in the formulation, interpretation of the effects is most easily accomplished graphically (see *Quantifying DDHS*). Note that we assume that DDHS occurs as a function of the current density, not a time-lagged density. Whereas time lags are important for density dependence to operate on population growth rates (Turchin, 1990), we assume that the mechanisms for DDHS (competition for resources, safety in numbers) operate more instantaneously than their fitness consequences.

## Model fitting

We created all quadratic and interaction terms and separately scaled and centered them before fitting. We did not transform the log-density offset or the sine and cosine of aspect, but we z-transformed all other variables to facilitate model fitting. We performed all data preparation and analyses in R (v. 4.1.1), and we fitted the GLMM via MCMC using R package NIMBLE v. 0.11.1 (de Valpine et al., 2017, 2021). We used Laplace priors with mean 0 on all regression parameters, referred to as the ‘Bayesian lasso’, to prevent overfitting (Hooten & Hobbs, 2015). We ran the model for 100,000 iterations across three chains, discarded the first 20,000 as burn-in (including adaptation) and thinned by 20 to obtain 4000 posterior samples/chain for inference. All analysis code and data are available on GitHub and published through Zenodo (doi:10.5281/zenodo.6687904).

## Model evaluation

We used the Gelman–Rubin statistic to evaluate MCMC convergence (Gelman & Rubin, 1992). We then used out-of-sample data to validate our model. Counts outside of YNP in the Montana portion of the study area were unavailable in 1994, 2002 or 2004, so we withheld the YNP data for these years from model fitting and used them to validate our model. Additionally, we withheld data from

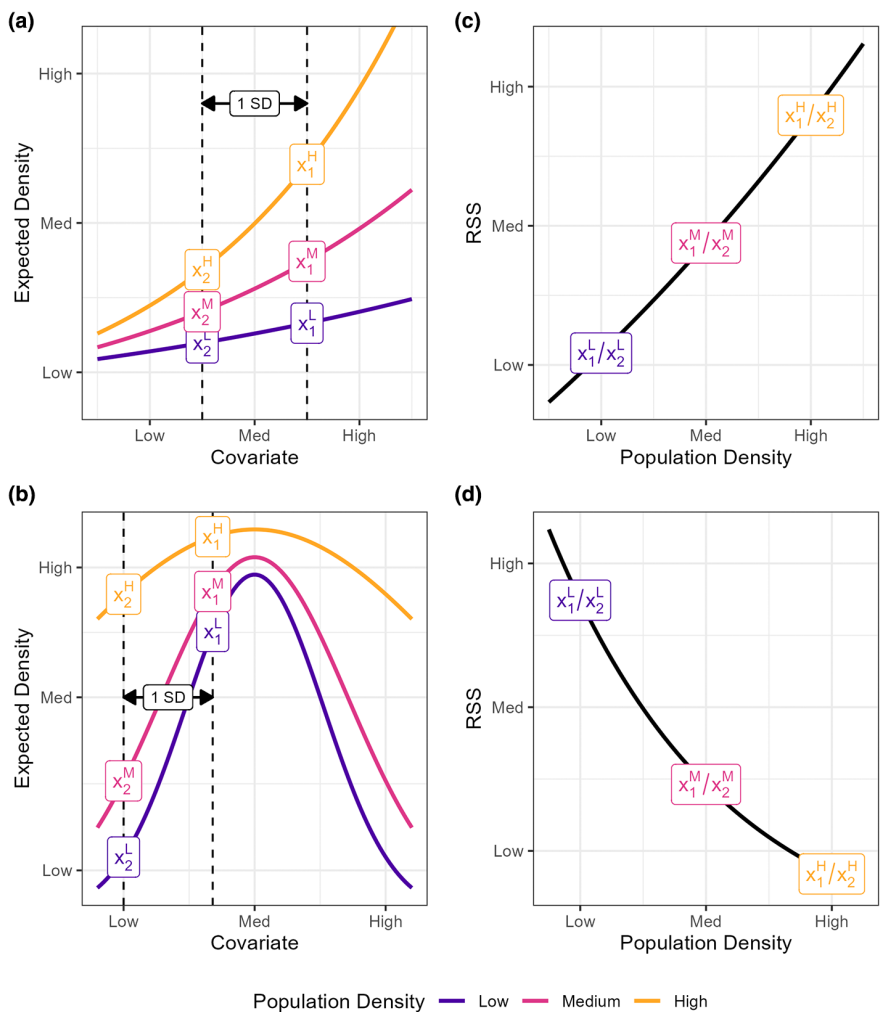
2020, which were available for the entire study area, as an additional year of validation data.

To perform validation, we used our fitted model to predict expected elk densities for each year, including all fixed and random effects. We compared expected elk densities under the model to observed densities from the count by calculating ordinary residuals ( $n_{i,t} - \lambda_{i,t}$ ), Pearson residuals ( $\frac{|n_{i,t} - \lambda_{i,t}|}{\sqrt{\lambda_{i,t}}}$ ) and Spearman's correlation coefficient between predicted and observed densities. We compared ordinary residuals and Spearman's correlation between the training and testing years. We used Pearson residuals to check for residual spatial autocorrelation by fitting non-parametric spline correlograms to these residuals (Bjørnstad & Falck, 2001).

To estimate model goodness-of-fit, we calculated a likelihood-based pseudo- $R^2$  for each posterior sample from the MCMC, yielding a distribution of pseudo- $R^2$ . We calculated our pseudo- $R^2$  using the method of Nagelkerke (1991), which compares the likelihood of the data under the fitted model to a null model. For our null model, we fitted a model where expected elk density was a function of only the time-varying offset ( $\alpha_t$ ), and the only other parameter estimated by the model was the size parameter of the negative binomial distribution,  $r$ .

## Quantifying DDHS

We quantified DDHS by measuring relative selection strength (RSS), the ratio of expected densities in two habitats (Avgar et al., 2017; Fieberg et al., 2021).  $\text{RSS}(x_1, x_2)$  is how many times more elk we expect at the habitat in the numerator ( $x_1$ ) compared to the habitat in the denominator ( $x_2$ ). The effect sizes of the habitat variable–density interactions indicate the strength of the DDHS, but because (1) this is on the scale of the link function, and (2) multiple variables are involved in the safety interaction, it is easier to visualise DDHS via RSS. Credible intervals around our RSS predictions account for the uncertainty in parameter estimates and their covariance, which clarifies inference on the quantity of interest (DDHS). Figure 1 provides a hypothetical example of how RSS indicates DDHS. Our model estimates expected density using linear (food; Figure 1a) or quadratic (safety; Figure 1b) formulations. A plot of RSS as a function of density reveals the pattern of DDHS: positive slopes show positive DDHS (Figure 1c), negative slopes show negative DDHS (Figure 1d) and a horizontal line shows no DDHS. We calculated RSS for the food and safety variables by comparing habitats that differ by one standard deviation (SD) in the focal variable (with all other variables held at their mean), which makes the magnitude of RSS comparable between variables. For safety, we altered openness and roughness by 0.5 SD each (1 SD total). The habitat we chose for the numerator ( $x_1$ ) always had the higher expected elk density. For the food variable,



**FIGURE 1** Conceptual figure depicting density-dependent habitat selection (DDHS). Left column (a, b) shows how expected density could change with a habitat covariate and average population density (colour). Right column (c, d) recasts the patterns in (a) and (b) in terms of relative selection strength (RSS), the ratio of expected densities in different habitats. We calculated RSS( $x_1, x_2$ ) as the ratio of expected densities for a 1-SD change in the covariate (dashed vertical lines). For our purposes, the numerator ( $x_1$ ) is always larger than the denominator ( $x_2$ ). ‘H’, ‘M’ and ‘L’ in superscript refer to high, medium and low population density. Calculating RSS across a range of average population densities (e.g.  $x_1^H/x_2^H, x_1^M/x_2^M, x_1^L/x_2^L$ ) yields the RSS curve, which more clearly demonstrates DDHS. In (a), expected density is modelled with just a linear term for the covariate, and expected density increases monotonically with an increase in the covariate (positive habitat selection). In this example, RSS (slope of each line) increases with population density; this is positive DDHS (c). Alternatively, if RSS decreased with population density, this would be negative DDHS (not shown). In (b), expected density is modelled with linear and quadratic terms such that expected density peaks at an intermediate value. A narrow parabola at low density indicates stronger selection, whereas a wider parabola at high density indicates weaker selection. This example demonstrates negative DDHS but note that positive DDHS is also possible. We calculated RSS as the ratio of expected densities when the covariate is near the vertex of the parabola ( $x_1$ ) that is, when the habitat covariate is lower ( $x_2$ ). Calculating RSS across a range of population densities yields the RSS curve (d), which in this case demonstrates negative DDHS. In summary, whether a habitat covariate is modelled with solely a linear term or also includes a quadratic term, the slope of the RSS curve plotted against population density shows the pattern of DDHS.

$x_1 = 661 \text{ kg/ha}$  and  $x_2 = 376 \text{ kg/ha}$  (Figure 1a). For the safety variables, we chose values that did not span the vertex (Figure 1b). Under our hypothesis of a trade-off between food and safety, we predicted positive DDHS for food and negative DDHS for safety.

To understand the relative drivers of habitat selection, we compared the magnitude of RSS between our habitat variables. Together with our predictions of positive DDHS for food and negative DDHS for safety, we predicted that habitat selection was driven by safety at low densities and food at high densities; that is, we predicted

that the two RSS curves with opposite slopes would cross.

To understand the impact of density on predicted elk distribution, we created a map of the study area showing the change in RSS from high ( $9.3 \text{ elk/km}^2$ ) to low ( $2.0 \text{ elk/km}^2$ ) density which approximated the observed decrease over time. We fixed all spatial covariates and predator densities to their value in 2008, then we calculated RSS where  $x_1$  was each observed pixel and  $x_2$  was a habitat with mean values for all covariates. We took the log of RSS so that relative to mean conditions, positive values

represent selection, zero represents no preference and negative values represent avoidance. We repeated this at low and high elk density. For each pixel, we subtracted the high-density log-RSS from the low-density log-RSS, which we term  $\Delta$  log-RSS. Positive values indicate that selection for the pixel increased as elk density decreased, whereas negative values indicate that selection decreased with elk density.

## RESULTS

### Model overview & assessment of key assumptions

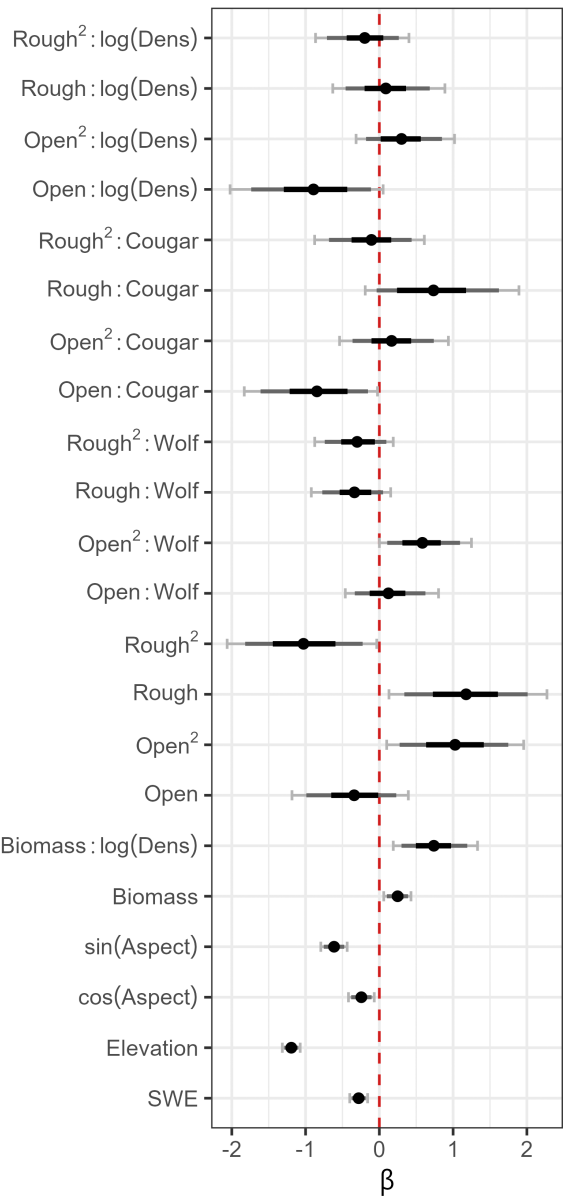
Our model indicated that elk selected for southwest-facing, low-elevation slopes with minimal snow cover. This is indicated by negative coefficients for  $\cos(\text{Asp})$  [aspect northing],  $\sin(\text{Asp})$  [aspect easting], Elevation and SWE (Figure 2, Figure S6).

Expected elk density increased with herbaceous biomass (Figure S7a), consistent with our assertion that herbaceous biomass reflects elk forage availability. Contrary to our prediction that elk would most prefer an intermediate openness, expected elk density was greatest at 100% openness, and the relationship between openness and elk density was largely monotonic for the observed range of openness (Figure S7b). Expected elk density was greatest for a roughness of 23.5 m (intermediate, as expected) with all other covariates held at their mean (Figure S7c). Spearman's correlation between biomass and the product of openness and roughness was  $-0.18$ , supporting our assumption that food and safety are negatively correlated.

Elk altered selection for safety variables with increasing predator densities in a manner that indicated openness and roughness were valid indices of spatial variation in predation risk. As wolf density increased, RSS for openness decreased (Figure 3a), whereas it increased as cougar density increased (Figure 3b), consistent with our predictions. Wolf density increased RSS for roughness, but the effect of cougars was negligible (Figure S8). Wolf density shifted the vertex of the parabola in the expected direction (from wolf to cougar habitat domains): at low wolf density (0 wolves/100 km<sup>2</sup>), the vertex was 22.3 m (90% credible interval [CI]: 20.1–25.0), and at high wolf density (10 wolves/100 km<sup>2</sup>), the vertex was 26.1 m (90% CI: 22.1–32.1; Figure 3c). Cougar density did not shift the vertex (Figure 3d).

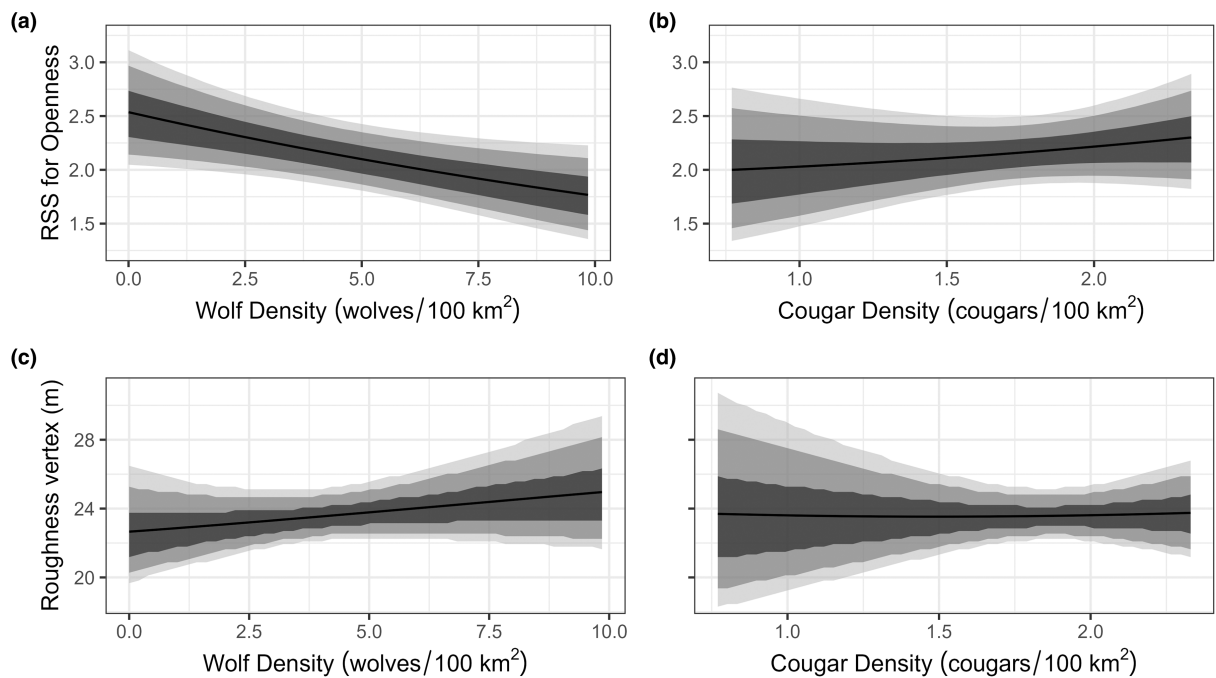
### Density-dependent habitat selection

Elk exhibited positive DDHS with respect to food, demonstrated by the positive Biomass:log(Dens) coefficient (Figure 2) and the positive slope of the RSS curve (Figure 4). Elk exhibited negative DDHS for openness,

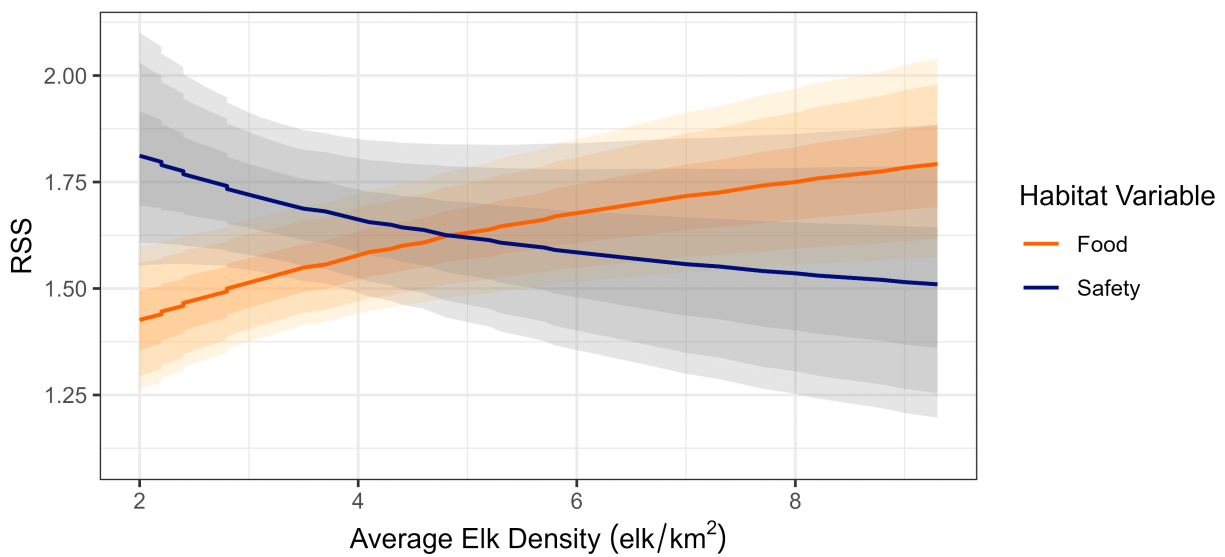


**FIGURE 2** Fitted model coefficients. Points are posterior means and bars are credible intervals (black bars: 50% credible intervals; dark grey bars: 80% credible intervals; light grey bars with end caps: 90% credible intervals). Red dashed line indicates 0 (no effect).

shown by the negative Open:log(Dens) coefficient (Figure 2) and the negative slope of the RSS curve (Figure 4). Since the Open<sup>2</sup>:log(Dens) coefficient was estimated near 0 (Figure 2), the linear coefficient drove the pattern (Figure 4). These patterns are consistent with a trade-off between food and safety, expected under the multidimensional IFD (Avgar et al., 2020), supporting our main hypothesis. By contrast, elk exhibited no DDHS with respect to roughness. The 50% CIs for the Rough:log(Dens) and the Rough<sup>2</sup>:log(Dens) terms overlapped 0 (Figure 2). Although the mean trend in RSS for roughness with increasing elk density was slightly positive, high uncertainty indicated this effect was negligible (Figure S9).



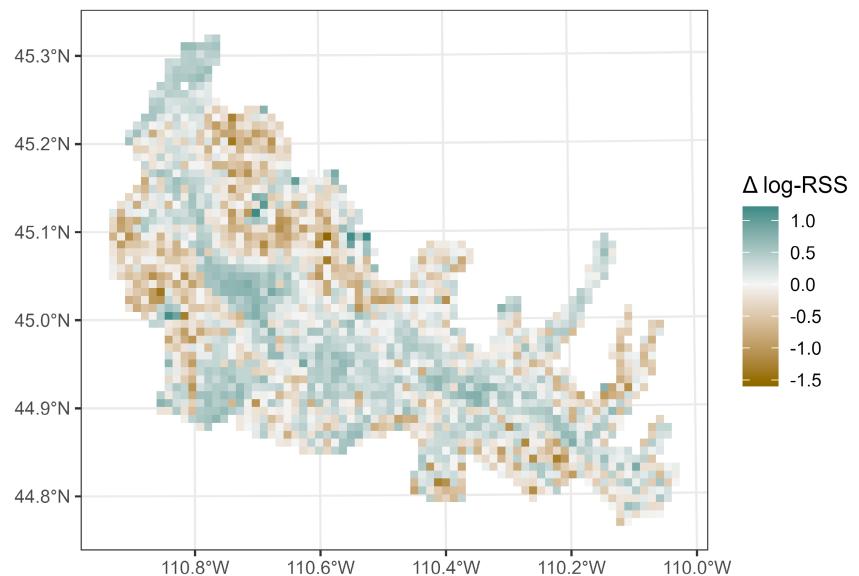
**FIGURE 3** Changing elk selection for risk variables with predator density. Openness and roughness describe important dimensions of the wolf and cougar habitat domains, that is, those habitats where they are most likely to kill elk. The wolf habitat domain is open and smooth, whereas the cougar habitat domain is rough and more forested. Relative selection strength (RSS) for openness (top row) is the ratio of expected density when openness is 100% versus when openness is 82%, a 1 SD change in openness. (a) RSS for openness decreases with wolf density (less selection for the wolf habitat domain), whereas (b) it increases with cougar density (less selection for the cougar habitat domain). The roughness vertex (bottom row) is the value of roughness that is most strongly selected by elk, that is, the vertex of the parabola describing selection. (c) The roughness vertex shifts to rougher habitat as wolf density increases (away from the wolf habitat domain); however, (d) elk did not shift the roughness vertex in response to cougar density. In all panels, predictions were made using samples from the entire posterior distribution. Solid black lines are mean effects and shaded grey envelopes are 50%, 80% and 90% credible intervals.



**FIGURE 4** Density-dependent habitat selection for food and safety. At low density (x-axis), relative selection strength (RSS) for a one standard deviation (1 SD) change in safety (0.5 SD of openness and 0.5 SD of roughness) is greater than the RSS for a 1 SD change in food (herbaceous biomass). At high density, this relationship flips, with a greater RSS for food than for safety. Changing average range-wide elk density alters the main driver of elk habitat selection from safety at low density to food at high density. Solid lines are posterior mean estimates and shaded envelopes are 50%, 80% and 90% credible intervals.

The stronger driver of elk habitat selection was safety at low elk density and food at high elk density (Figure 4), matching our prediction. At low elk density

(2.0 elk/km<sup>2</sup>), the RSS for a one-SD change in food was 1.43 (90% CI = 1.26–1.60) and the RSS for a one-SD change in safety was 1.81 (90% CI = 1.55–2.10). At high



**FIGURE 5** Change in log-RSS from high elk density to low elk density. Relative selection strength (RSS) is the ratio of expected density in each pixel of the landscape to the expected density in a habitat with all habitat variables held at their mean. The natural logarithm of RSS (log-RSS) is a measure of habitat selection, with positive values indicating preference versus the mean conditions, zero indicating no preference versus the mean conditions, and negative values indicating avoidance versus the mean conditions. Here, we plot  $\Delta \log\text{-RSS}$ , the difference between log-RSS when average range-wide elk density is high ( $9.3 \text{ elk}/\text{km}^2$ ) versus when average range-wide elk density is low ( $2.0 \text{ elk}/\text{km}^2$ ). Positive values (blue-green pixels) indicate that selection for the pixel increased as elk density decreased (the observed pattern over time), whereas negative values (brown pixels) indicate that selection for the pixel decreased as elk density decreased. All habitat variables and predator densities are held at their 2008 levels for demonstration.

elk density ( $9.3 \text{ elk}/\text{km}^2$ ), the RSS for a one-SD change in food was 1.79 (90% CI = 1.57–2.04) and the RSS for a one-SD change in safety was 1.51 (90% CI = 1.20–1.88).

Posterior mean  $\Delta \log\text{-RSS}$  varied from  $-1.6$  to  $3.0$  with a mean of  $0.0$  across all pixels, indicating that changing selection resulted in habitat switching across space—the selection strength for some habitats decreased while the selection for other habitats increased, resulting in a redistribution of the population. Figure 5 illustrates the spatial pattern in  $\Delta \log\text{-RSS}$ .

## Model evaluation

The Gelman–Rubin statistics indicated MCMC convergence for all top-level parameters (Appendix S6). Ordinary residuals and Spearman's correlations were similar between in-sample and out-of-sample predictions (Figure S10). Residuals had mean near 0, indicating good model accuracy. Mean pseudo- $R^2$  for our model was 0.065 (90% CI = 0.064–0.066), indicating low precision (to be expected under the negative binomial distribution). We found little to no residual spatial autocorrelation in all years (Figure S11).

## DISCUSSION

Density dependence is a fundamental concept in ecology, and its importance for a species' abundance over time is

well understood (Berryman & Turchin, 2001; Brook & Bradshaw, 2006; Dennis & Taper, 1994). Although theory establishes that density dependence should also act on abundance in space (Morris, 2003; Rosenzweig, 1991), comparatively little empirical work has demonstrated the role of density dependence in shaping population distribution. In this study, we provide rare empirical evidence that density alters drivers of habitat selection and population distribution in a free-living system, consistent with theoretical expectations under the multidimensional IFD (Avgar et al., 2020). This is a broadly important conceptual advance because it links population density to the community-level interactions (i.e. consumer–resource, predator–prey) that determine an organism's distribution in space (Rosenzweig & Abramsky, 1997). Observations of this system at a constant density would lead to erroneous inference on the drivers of habitat selection, biased predictions of population distribution and misunderstanding of community interactions under unobserved densities.

Interactions between predator densities and safety variables indicate that the latter were valid indices of predation risk. RSS for openness decreased with wolf density (Figure 3a) and increased with cougar density (Figure 3b). The effect of predator densities on RSS for roughness was weak; rather, increasing wolf density pushed the most preferred roughness into rougher habitat (Figure 3c). We found no effect of cougar densities on selection for roughness (Figure 3d), and the predator density effects were greater for wolves than for cougars.

Nevertheless, elk avoided densely forested, rough habitats across all elk and predator densities (Figure S7b,c), indicating that risk from cougars was an important driver of elk distribution. Our finding that elk selected for maximum openness (Figure S7b) and that DDHS for safety was primarily driven by openness (Figure S9b) further suggests that elk distribution was more strongly influenced by cougars than wolves. These results independently corroborate the findings of Kohl et al. (2019) that multiple predators (wolves and cougars) influence elk space use.

We found support for our hypothesis that the food-safety trade-off in our system leads to a switch from safety driving distribution at low density to food driving distribution at high density (Figure 4). We found evidence for positive DDHS for food and negative DDHS for safety, which we would expect when there is a trade-off between food and predation risk. Indeed, biomass and the product of openness and roughness were negatively correlated (Spearman's  $r = -0.18$ ). While this food-safety trade-off is expected to be ubiquitous in nature (Brown & Kotler, 2004), it is difficult to measure, and our approach provides evidence of it. Were there no trade-off, elk at low density could occupy habitats that provided both food and safety, and we would expect spillover from those habitats as density increases (i.e. negative DDHS). The negative correlation between food and safety means that elk cannot satisfy both of these requirements in the same place, thus giving rise to the trade-off and habitat switching as density increases (i.e. positive DDHS; Greene & Stamps, 2001; Avgar et al., 2020). The switch in the relative importance of food and safety leads to some habitats on the landscape becoming less selected while others become more selected; the result is a redistribution of the population (Figure 5). This switch is consistent with theoretical and empirical work on behavioural ecology in predator-prey systems. Organisms should change their risk assessment based on conspecific density (Peacor, 2003), and antipredator behaviours such as vigilance and habitat selection should adjust accordingly (Mooring et al., 2004). This is also consistent with the predation-sensitive food hypothesis, whereby both food and predation limit prey populations (Sinclair & Arcese, 1995); as competition increases and food becomes limiting, prey will increase selection for food and decrease selection for safety. Our work demonstrates how these behaviours manifest in space.

Current empirical understanding of DDHS in free-living vertebrate systems is mostly limited to small mammals (e.g. Morris, 1989; Morris et al., 2000; Rosenzweig & Abramsky, 1985) or domesticated mammals on islands (e.g. Mobaek et al., 2009; van Beest et al. 2014b). Previous studies of DDHS in free-living systems, though rare, have also focused on elk (Merrill et al., 2020; van Beest et al. 2014a) or congeneric red deer (*Cervus elaphus*; McLoughlin et al., 2006; Pérez-Barbería et al., 2013); see

Supplemental Discussion for a comparison with our work (Appendix S7). Often, space-use data are collected on short time scales during which population densities are similar (McLoughlin et al., 2010), and the resulting habitat selection models are snapshots of true, DDHS patterns (Boyce et al., 2016; Boyce & McDonald, 1999; Northrup et al., 2022). Our results show that inference or predictions from these snapshots may be unreliable under changing densities.

Space use of organisms can be measured from two perspectives: a population-level (Eulerian) perspective which measures changes in population density at various places over time or an individual-level (Lagrangian) perspective which tracks the locations of individuals over time (Turchin, 1998). Analyses from the Eulerian perspective lend themselves more readily to projection across space at large scales, whereas analyses from the Lagrangian perspective can uncover more detailed mechanism. Individual behaviour scales up to population-level distributions (Mueller & Fagan, 2008), so results from the two perspectives can, in principle, be reconciled. In practice, few studies have compared the Eulerian and Lagrangian approaches to studying population-level space use (but see Phillips et al., 2019; Bassing et al., 2022). In our case, we find both similarities and discrepancies between our work (Eulerian) and previous studies (Lagrangian) of elk habitat selection in northern Yellowstone (Appendix S7). Without the Lagrangian perspective, we could not identify whether DDHS occurred because (1) individuals changed their habitat selection traits (behavioural plasticity), or (2) individuals that exhibited certain habitat selection traits had differential survival and/or reproduction (demographic sorting), although previous research has suggested it is a demographic sorting effect (White et al., 2012). Which perspective provides the best understanding of DDHS is still an open question. Most empirical work on DDHS has made use of isodar theory (Morris, 1988, 2011), which is constructed from the Eulerian perspective and naturally integrates fitness. An important knowledge gap concerns how to connect DDHS to fitness from the Lagrangian perspective, which can reveal more detailed mechanisms of how population regulation plays out in space.

We believe the capacity of DDHS to qualitatively alter community-level interactions is underappreciated and understudied (see Rosenzweig & Abramsky, 1997), especially in predator-prey systems where positive DDHS may lead to landscape-scale habitat switching. For example, the relative importance of consumptive versus non-consumptive effects (Peacor et al., 2013; Sheriff et al., 2020) may critically depend on DDHS (habitat selection is an important antipredator trait; Trussell et al., 2006). If prey only respond to risk by altering their habitat-selection traits at low density (when competition is less costly), the importance of non-consumptive effects may be overstated. Similarly, DDHS may impact competition between prey species. Habitat selection is an

important mechanism for reducing competition between species, which may determine the level of competition (Rosenzweig, 1981), especially when prey species face differential risk. For example, spatial overlap between bison and elk in northern Yellowstone is potentially greater than expected at high elk density if neither species responds strongly to predators.

In conclusion, consistent with theoretical expectations, density dependence alters habitat selection and distribution in free-living systems. Incorporating DDHS into models of distribution is crucial whenever density is variable; ignoring its effects may lead to severely compromised inferential and predictive performance. Our findings underscore that the effects of the food-safety trade-off on prey distribution are dynamic, and that inference and prediction in these systems depends on prey density.

## AUTHOR CONTRIBUTIONS

BJS, DRM and TA conceived and developed the idea. DRS and DWS collected data and supervised the project. BJS performed analyses. All authors contributed to the interpretation of the results. BJS wrote the first draft of the manuscript and all authors contributed substantial edits.

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## DATA AVAILABILITY STATEMENT

All data (and code) are available on GitHub and published through Zenodo (doi:[10.5281/zenodo.6687904](https://doi.org/10.5281/zenodo.6687904)).

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## SUPPORTING INFORMATION

Additional supporting information can be found online in the Supporting Information section at the end of this article.

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## Supporting Information

### Density-dependent habitat selection alters drivers of population distribution in northern Yellowstone elk

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## 1 Appendix S1 – Aerial Survey Timing and Scale

2 We analyzed aerial survey data – designed to census the population – collected from 1988 to 2020.  
3 Elk were counted by 3 – 4 fixed-wing aircraft flying non-overlapping areas of the winter range (Fig. S1).  
4 Surveys took place over one to four days between late December and March of each year (Fig. S2A).  
5 Winter surveys were assigned to the year of that January; e.g., the survey for the winter of 1991 occurred  
6 in January 1991 and the survey for 1992 occurred in December 1991. The count did not occur in 1996 or  
7 1997. The count was done via helicopter in 2019 and was not included due to differing protocol. Counts  
8 for 1989, 1991, 2006, and 2014 are considered unreliable as a census of the population and so were not  
9 included. For three years (1994, 2002, and 2004), georeferenced group data are only available inside of  
10 Yellowstone National Park (data from the State of Montana were not georeferenced). We excluded these  
11 data from model fitting and used them for model validation (see main text). A full census was available  
12 for 2020, but we chose to withhold it as an additional validation year with complete georeferencing.

13 Since survey timing varied from year to year, we checked that survey timing did not introduce a  
14 bias in our model. We plotted the model’s ordinary residuals as a function of survey date, and visually  
15 checked for any pattern by fitting a very flexible GAM smoothing line (Fig. S3). The fitted line showed  
16 no obvious pattern. We tested this further with an ANOVA, where we treated each of the 15 unique mean  
17 survey dates as categorical variables and the Pearson residuals as the response. We computed p-values for  
18 pairwise differences using Tukey’s Honest Significant Difference method to check for differences  
19 between Pearson residuals for any pair of dates. All p-values were large ( $>0.23$ ; Table S2), which indicated  
20 no meaningful differences between residuals for any pair of dates.

21 Surveys recorded groups between 07:00 and 15:00, but 92% of observations were between 08:00  
22 and 12:00 (Fig. S2B). Kohl *et al.* (2019) found that wolf movements peaked at 10:00 (in the middle of

23 these survey times), whereas cougar activity peaked at 03:00 and 04:00 for male and female cougars,  
24 respectively (outside all survey times). Thus, surveys occurred at a time of day when wolves were the  
25 more active predator. To account for this and other potential temporal effects, we chose a coarse spatial  
26 scale to average over daily movements. Our 1 km x 1 km raster cells were chosen to be coarse enough to  
27 accomplish this. We verified that elk daily winter movements rarely exceeded this area by comparing the  
28 area of a raster cell ( $1 \text{ km}^2$ ) to the area used daily by GPS-collared elk during winter (data not included in  
29 this manuscript). We found that on 93% of winter elk-days, a 100% MCP around all of an elk's locations  
30 had an area  $< 1 \text{ km}^2$  (B. Smith, pers. comm.). This was based on 44,444 elk-days from 169 unique  
31 individuals tracked between 2001 – 2022.

32 Furthermore, this coarse spatial grain is large enough to capture (potentially multiple) patches of  
33 trees within a matrix of openness. For example, a raster cell with 50% openness typically contains a mix  
34 of open and forested patches – not a homogenous patch of intermediate forest cover. This patchy habitat  
35 allows elk to shuttle between patches across the diel cycle, whereas a raster cell with 0% or 100% openness  
36 hinders this diel shuttling behavior (Fig. S4). Fitting a parabola to openness therefore measures how much  
37 heterogeneity in openness is selected for by elk.

38 We have designed our modeling approach with respect to spatial and temporal scale, while  
39 considering variation in survey dates and times, and, thus, we believe our model is a valid representation  
40 of the winter distribution of northern Yellowstone elk that is insensitive to survey date and time of day.

41

42 Appendix S2 – Sensitivity to Imperfect Detection

43 Previous research evaluating aerial counts of elk, including work in northern Yellowstone, has  
44 found that elk group size and vegetation cover (i.e., tree canopy openness) affect detection of elk groups  
45 (Samuel *et al.* 1987; Singer & Garton 1994). Because openness is an important covariate in our habitat  
46 selection model, we were concerned that imperfect detection as a function of openness could bias our  
47 results. We did not have data to explicitly estimate imperfect detection for each of the survey years we  
48 analyzed (e.g., mark-resight data, double observers, or multiple surveys). Instead, to evaluate any potential  
49 bias, we conducted a sensitivity analysis wherein we excluded pixels outside a certain openness threshold  
50 from the analysis and refitted the model. We compared coefficients and predicted DDHS patterns between  
51 the original model and the reduced dataset models.

52 We used three openness cutoffs: <30%, <50%, and >99.9%. We discarded all pixels that had  
53 openness less than/greater than the cutoff in *any* year and refitted the model. The first two cutoffs (<30%  
54 and <50%) discarded the pixels we considered least reliable, i.e., due to their low openness, we expected  
55 detectability to be low. The third cutoff (>99.9%) discarded the pixels we considered most reliable; we  
56 did this to determine if the remaining pixels were reliable enough to replicate the pattern on their own.  
57 We ran each MCMC procedure with 2 chains and 50,000 iterations per chain, discarded the first 22,000  
58 iterations as burn-in and adaptation, and then thinned by 7 to 4,000 iterations per chain. All code for this  
59 analysis can be found with the main analysis on GitHub and archived through Zenodo  
60 (doi:10.5281/zenodo.6687905).

61 The resulting dataset for the <30% cutoff was 95% of the original data, the resulting dataset for  
62 the 50% cutoff was 70% of the original data, and the resulting dataset for the >99.9% cutoff was 73% of  
63 the original data. We found all of our resulting patterns were qualitatively similar between the models: we

64 found positive DDHS for biomass, negative DDHS for safety, and a change in their relative importance  
65 (Fig. S5). Note that for the <99.9% cutoff, the means do not cross in the plotted domain, but the credible  
66 intervals overlap, and the pattern remains similar (Fig. S5D). Thus, we concluded imperfect detection did  
67 not meaningfully impact our inference.

68 Appendix S3 – Model Structure and Random Effects

69 As detailed in the main text, we modeled the expected count ( $\lambda_{i,t}$ ) in each pixel in each year as a  
70 function of 22 covariates (indexed by  $k$ ; Table S1), a time-varying offset ( $\alpha_t$ ), a temporal random effect  
71 ( $\eta_{i,t}$ ), and a spatial random effect ( $s_i$ , eqn. 1). We described the fixed covariates in the main text, and we  
72 described the offset and random effects in detail here.

73 *Offset*

74 To account for changing total abundance across years, we included a time-varying offset,  $\alpha_t$ . The  
75 offset was the log of average range-wide density of elk across the study area in each year, i.e., the natural  
76 logarithm of the total count ( $N_t$ ) divided by the number of pixels in the study area ( $\Omega$ ),  $\log\left(\frac{N_t}{\Omega}\right)$ . If none  
77 of the covariates had any effect (i.e., for all  $k$ ,  $\beta_k = 0$ ), then the expected density in each pixel would  
78 simplify to  $\lambda_{i,t} = \exp[\alpha_t] = \exp\left[\log\left(\frac{N_t}{\Omega}\right)\right] = N_t/\Omega$ , which is simply the average range-wide density for  
79 that year. In that way, the offset acts like a time-varying intercept. Note that when we refer to “average  
80 range-wide density” or “log average range-wide density” throughout, we are referring to a predictor  
81 variable, such as this offset described here or the interaction terms described below. Contrast this with  
82 “expected density,” which is the expectation of our response variable,  $\lambda_{i,t}$ .

83 *Temporal Random Effect*

84 We used the temporal random effect ( $\eta_{i,t}$ ) to account for temporal fluctuations in the effect of a  
85 raster pixel being outside YNP compared to inside YNP. The two main motivations for this were (1) an  
86 observed shift in the elk distribution from primarily inside YNP in the early years of our study to primarily  
87 outside the northern boundary of YNP in later years (Tallian et al. 2017; MacNulty et al. 2020), and (2) a  
88 late hunt managed by the state of Montana that occurred in January of each year prior to 2010 but ceased

89 afterward (MacNulty et al. 2020). We estimated the effect in each year ( $t$ ) as coming from a normal  
90 distribution with mean  $\mu_\eta$  and variance  $\sigma_\eta^2$ . We multiplied the effect by a binary covariate indicating  
91 whether each pixel was outside of YNP, with outside = 1 and inside = 0.

92 The temporal random effect ( $\eta_{i,t}$ ) had a mean of -1.74 (90% credible interval = -2.05 – -1.42), and  
93 a variance of 0.56 (90% credible interval = 0.24 – 1.09). The posterior mean effect for each year ranged -  
94 2.77 – -0.39, and the trend was positive over time (Fig. S12). In this context, negative values indicate  
95 lower elk densities outside YNP than would be expected based on the fixed effects in the model and the  
96 positive trend indicates increasing densities outside YNP over time.

97 *Spatial Random Effect*

98 We used the spatial random effect ( $s_i \in \mathbf{s}$ ) to account for residual spatial autocorrelation in elk  
99 density, i.e., consistent spatial pattern in elk density not captured by the other terms in the model. We  
100 modeled it as a Gaussian process (a.k.a., kriging), i.e., the vector  $\mathbf{s}$  is a realization of a multivariate normal  
101 distribution with mean 0 and variance-covariance matrix,  $\Sigma$  ( $\mathbf{s} \sim N(\mathbf{0}, \Sigma)$ ). Each element of the variance-  
102 covariance matrix was calculated as a decaying exponential function of the distance between the two  
103 pixels,  $\|x_i - x_j\|$ , and two free parameters,  $\sigma$  and  $\rho$ , estimated by the model:  $\Sigma_{i,j} = \sigma^2 \times \exp\left(-\frac{\|x_i - x_j\|}{\rho}\right)$ .  
104 The parameter  $\sigma$  controlled the variance within a pixel (i.e., the distance between a pixel and itself is 0,  
105 thus  $\Sigma_{i,i} = \sigma^2$ ). The parameter  $\rho$  controlled the rate of decay with distance, with smaller  $\rho$  indicating faster  
106 decay. To improve model fitting, we scaled distances between pixels to range from 0 – 1 by dividing by  
107 the maximum distance.

108 The estimated spatial random effect, which was constant across time, was near 0 inside YNP and  
109 positive outside YNP (Fig. S13). The posterior mean of each  $s_i$  ranged from -0.03 to 1.61, but the mean

110 of each mean  $s_i$  inside YNP was 0.01, whereas outside YNP it was 0.22. The decay parameter of the  
111 spatial covariance function ( $\rho$ ) had posterior mean = 14.08 (90% credible interval = 6.67 – 19.49) and the  
112 standard deviation parameter ( $\sigma$ ) had posterior mean = 0.63 (90% credible interval = 0.43 – 0.79). Thus,  
113 the covariance between two pixels separated by 1 km was 0.41 (90% credible interval = 0.19 – 0.63) and  
114 the covariance between two pixels separated by 100 km was 0.38 (90% credible interval = 0.16 – 0.59),  
115 indicating overall low spatial autocorrelation (Fig. S14). Together, the two random effects resulted in  
116 some spatially autocorrelated variation in elk densities outside YNP, with densities inside YNP almost  
117 entirely captured by the fixed effects (Fig. S15).

118 Appendix S4 – Equivalence to Habitat Selection Functions

119 Our approach to modeling habitat selection at the population level was parallel to more familiar  
120 approaches taken with individual telemetry locations, habitat-selection functions (HSFs, a.k.a., RSFs). In  
121 most HSFs, the observed locations (coded as 1) and background locations (coded as 0) are analyzed with  
122 logistic regression, the parameters of which are exactly equivalent to the parameters of an inhomogeneous  
123 Poisson point process (IPP) model (reviewed in Fieberg *et al.* 2021). The IPP model describes the *density*  
124 of points as a function of habitat variables; however, in telemetry-based HSFs, the resulting predictions  
125 are only *proportional* to density (Avgar *et al.* 2017). An equivalent model can be fit by discretizing space  
126 and using Poisson regression (Manly *et al.* 2002; Hooten *et al.* 2017). As we describe below, we fit our  
127 HSF using the latter approach – we modeled elk counts as a function of habitat covariates in discrete raster  
128 pixels (note that we used the negative binomial distribution as an overdispersed Poisson distribution). The  
129 resulting coefficients (the  $\beta$ s described in the main text for the covariates listed in Table S1) are interpreted  
130 in exactly the same way as if they had come from a telemetry HSF – but in our population-level study, we  
131 interpret the model predictions as spatiotemporally explicit elk densities.

132

133     Appendix S5 – Covariate Processing

134       We began with a master template raster, with a projected coordinate reference system (EPSG:  
135       26912; UTM zone 12, NAD83) – the extent determined by the northern Yellowstone elk winter range  
136       polygon (Fig. S1) – and 1 km x 1 km resolution. We rasterized all count data and covariates (see below)  
137       on this template, using bilinear interpolation when resampling the covariate rasters. We projected rasters  
138       from their native resolution to our master template raster using the function `projectRaster()` from  
139       the `raster` package (Hijmans 2022). After all covariate processing, we retained only pixels falling  
140       within the northern winter range boundary for analysis, a total of 1,978 1-km<sup>2</sup> pixels.

141     *Conditions*

142       We included conditions to account for known drivers of elk density that were not the target of  
143       inference. These were snow-water equivalent (SWE), elevation, cosine of aspect (aspect northing), and  
144       sine of aspect (aspect easting).

145       We obtained SWE data from Daymet v 4.0 (Thornton *et al.* 2020).

146       We obtained a digital elevation model (DEM) from the National Map’s 3DEP program (U.S.  
147       Geological Survey 2020) The DEM was projected to a 30 x 30 m resolution in a UTM projection (zone  
148       12N, NAD83). We extracted elevation from the DEM directly and processed the DEM to calculate aspect  
149       (and roughness, see *Safety*) rasters in R using the `terrain()` function from the `raster` package  
150       (Hijmans 2022). We calculated aspect northing (cosine of aspect) and aspect easting (sine of aspect) from  
151       the aspect raster. Following all calculations at a 30-m resolution, we reprojected the raster to our master  
152       template raster.

153     *Food*

154       We measured food using total herbaceous biomass from the annual biomass (v2.0) layer from the  
155       Rangeland Analysis Platform (RAP; Jones *et al.* 2021). For each winter, we used the layer corresponding  
156       to the previous growing season to measure potential elk forage available from the previous growing season  
157       (e.g., for winter 1988, we used total annual biomass from 1987). We downloaded total biomass layers  
158       from RAP for 1987 – 2019, which were natively in geographic coordinates (longitude and latitude;  
159       WGS84) at approximately 30 m resolution (0.0002694946 degrees) and organized in two layers: annual  
160       herbaceous biomass and perennial herbaceous biomass. We summed these two layers to obtain total  
161       herbaceous biomass. We reprojected a stack of total herbaceous biomass rasters for each year to our master  
162       template. Finally, we log-transformed this covariate to reduce the influence of very high values from some  
163       areas of irrigated agriculture north of YNP.

164     *Safety*

165       We measured predation risk using the risky places approach, and previous research has shown that  
166       risk to elk from both wolves and cougars can be described by using tree canopy openness and terrain  
167       roughness (see main text for details).

168       We measured openness using the cover product from RAP. For each winter, we used the layer  
169       corresponding to that year (e.g., for winter 1988, we used cover for 1988). We downloaded the cover  
170       rasters from RAP for 1988 – 2020, and we extracted the trees layer, which measures the tree coverage in  
171       a pixel in %. As with herbaceous biomass, the cover layers were natively in geographic coordinates  
172       (longitude and latitude; WGS84) at approximately 30 m resolution (0.0002694946 degrees). We converted  
173       from tree cover to openness by subtracting the values from 100%, and then we divided by 100 to leave

174 the data in the range [0, 1], where 1 is completely open (no tree cover) and 0 is 100% tree cover. We  
175 reprojected a stack of openness rasters for each year to our master template.

176 We measured roughness using the DEM described above (see *Conditions*). We used the  
177 `terrain()` function with `opt = "roughness"` to calculate roughness from the DEM. For a focal  
178 pixel, roughness was measured as the difference between the maximum and the minimum value of the  
179 pixel and its 8 surrounding pixels, thus the units were m (inherited from the DEM). After calculating  
180 roughness at the 30-m resolution, we reprojected the raster to our master template raster.

181 *Predator densities*

182 To support our assertion that the safety variables reflect how elk perceive risk, we included  
183 interactions between wolf and cougar densities and each of the safety variables. Wolf counts were  
184 available from National Park Service monitoring, and we used wolf population size for the northern winter  
185 range, specifically. We converted wolf counts to densities by dividing by the area of the northern winter  
186 range polygon that falls within the boundaries of Yellowstone National Park (wolves/km<sup>2</sup>; Fig. S1) and  
187 then multiplied by 100 to get wolves/100 km<sup>2</sup>.

188 Cougar densities in northern Yellowstone have been estimated from field data during 3 phases  
189 across the period of time we analyzed. Phase 1 estimated the density each year from 1987 – 1993, Phase  
190 2 from 1998 – 2004, and Phase 3 from 2014 – 2017. Densities increased from Phase 1 through Phase 2,  
191 leveling off by Phase 3. We used linear interpolation across the entire dataset to fill in the gaps between  
192 Phase 1 – 2 and between Phase 2 – 3. From the end of Phase 3 until the end of our elk timeseries (2018 –  
193 2020), we used the average cougar density in Phase 3, rather than assuming continued growth. Fig. S16  
194 shows estimated and interpolated cougar densities.

195

196     Appendix S6 – Model Evaluation

197           We evaluated convergence of our MCMC algorithm by using the Gelman-Rubin statistic (Gelman  
198       & Rubin 1992), which we calculated using the function `gelman.diag()` from the R package `coda`  
199       (Plummer *et al.* 2006). Potential scale reduction factors (PSRFs) near 1 indicate MCMC convergence.  
200       PSRF point estimates for all top-level parameters were < 1.02 and the upper bounds of all confidence  
201       limits were < 1.06 (Table S3), indicating good convergence.

202           To estimate model goodness-of-fit, we calculated a likelihood-based pseudo- $R^2$  for each posterior  
203       sample from the MCMC, yielding a distribution of pseudo- $R^2$ . We calculated pseudo- $R^2$  using the method  
204       of Nagelkerke (1991), which compares the likelihood of the data under the fitted model to a null model.  
205       For our null model, we fitted a model where expected elk density was a function of only the time-varying  
206       offset ( $\alpha_t$ ), and the only other parameter estimated by the model was the size parameter of the negative  
207       binomial distribution,  $r$ .

208           Ordinary residuals and Spearman's correlations were similar between in-sample and out-of-sample  
209       predictions (Fig. S10). Residuals had mean near 0, indicating good model accuracy (Fig. S10A), and  
210       Spearman's correlations were similar between in-sample data (mean  $r = 0.28$ ) and out-of-sample data  
211       (mean  $r = 0.27$ ; Fig. S10B). Mean pseudo- $R^2$  for our model was 0.065 (90% CI = 0.064 – 0.066), indicating  
212       relatively low precision. We found little to no residual spatial autocorrelation in all years (Fig. S11).

213           Our model's low pseudo- $R^2$  is not surprising. Factors other than habitat, e.g., social interactions among  
214       elk, are at least partially responsible for group size distributions (Gerard *et al.* 2002; Brennan *et al.* 2015).  
215       Elk group sizes in our dataset range from 1 to over 1,000 individuals, and this is seen with elk in other  
216       study areas (Proffitt *et al.* 2012; Brennan *et al.* 2015). Thus, to capture the large variance in group size  
217       distributions, we chose to model elk counts using the negative binomial rather than the Poisson

218 distribution. The free dispersion parameter of the negative binomial can soak up the variation in counts  
219 rather than attributing that variation to covariates. This is an important feature of our model: modeling elk  
220 counts with the Poisson distribution would be misleading as it would require a stronger effect of habitat  
221 to capture the variation in the observed data (i.e., placing more weight on the covariates and potentially  
222 overfitting). One result of this is that predicting observed number of elk in a particular pixel shows low  
223 precision, i.e., a wide range of counts is plausible for a given expected value. While our model has low  
224 precision, it does have good accuracy, shown by out-of-sample validation (Fig. S10). Thus, our model  
225 estimates the expected number of elk in a pixel accurately and captures the slopes of our covariates from  
226 which we draw inference. Translating this expectation to an actual count was not the goal of our model.

227

228 Appendix S7 – Supplemental Discussion

229 *Comparison to previous studies*

230 **Elk and Red Deer**

231 Previous studies of DDHS in free-living systems, though rare, have also focused on elk (van Beest  
232 *et al.* 2014; Merrill *et al.* 2020) or congeneric red deer (*Cervus elaphus*; McLoughlin *et al.* 2006; Pérez-  
233 Barbería *et al.* 2013). In the absence of predation risk, red deer exhibited negative DDHS and spillover,  
234 as expected (McLoughlin *et al.* 2006; Pérez-Barbería *et al.* 2013).

235 Most similar to our work, Merrill *et al.* (2020) looked at DDHS in elk in Ya Ha Tinda under risk  
236 of wolf predation. This system exhibits the negative correlation between food and safety that we would  
237 expect would generate results similar to ours; however, the results do not show this. Ya Ha Tinda elk  
238 exhibited negative DDHS for forage biomass and positive DDHS for risky places – exactly the opposite  
239 of what we found. Additionally, an interaction between forage biomass and wolf risk became more  
240 strongly negative with elk density – i.e., elk selected the best forage even less under wolf predation risk,  
241 and this response to predators became stronger when elk were at high density (see Fig. 2 in Merrill *et al.*  
242 2020). One possible explanation is that elk can make use of the “human-shield” effect to effectively reduce  
243 their risk from wolves in this system. If this effect sufficiently reduces the predation risk, then we would  
244 no longer expect positive but instead negative DDHS for food, since elk can use the “shield” to select for  
245 the best forage, even at low density. In that case, the correlation between the food and safety variables  
246 would result in both being subject to negative DDHS, just as Merrill *et al.* (2020) reported. Nevertheless,  
247 this is another piece of evidence that DDHS occurs in large herbivores.

248 **Northern Yellowstone**

249 Several previous studies have focused on individual winter habitat selection by telemetered  
250 northern Yellowstone elk. Ours is the first to take a population-level (Eulerian) perspective; nonetheless,  
251 there is some broad agreement in our conclusions. Mao et al. (2005) compared habitat selection in two  
252 distinct time periods: 1985–1990 (pre-wolf reintroduction) and 2000–2002 (post-wolf reintroduction).  
253 While they assumed no DDHS, the population declined >25% ( $9.3 \text{ elk/km}^2$  in 1988;  $6.7 \text{ elk/km}^2$  in 2001)  
254 between these periods. They found elk selected more open habitats in winter post-wolf reintroduction – at  
255 lower density – agreeing with our finding of negative DDHS for openness (Fig. S9B). Kohl et al. (2018,  
256 2019) used step-selection functions (SSFs) to show that elk selected habitat to avoid wolves (2018) or  
257 wolves and cougars (2019), but only when those predators were most active. Elk use of risky places at  
258 times of day when risk was low may suggest that during the timing of our aerial surveys, elk would indeed  
259 prioritize food over safety. These studies used data from 2001 – 2004, during which time elk densities  
260 were at moderate levels but decreasing; they did not account for possible DDHS. Elk density in 2001 was  
261  $6.7 \text{ elk/km}^2$  and dropped to  $4.3 \text{ elk/km}^2$  by 2004, densities at which our model predicts food and safety  
262 are approximately equally important (Fig. 4).

263 Other studies of northern Yellowstone elk have qualitative discrepancies with our results. Fortin  
264 et al. (2005) concluded that elk used resources less under risk from wolves. Their study occurred at an  
265 intermediate density ( $6.7 \text{ elk/km}^2$  in 2001), when our model suggests food should be a stronger driver than  
266 safety (Fig. 4). This discrepancy could be due to scale and study design; they focused at a fine  
267 spatiotemporal scale using SSF. Cusack et al. (2020) looked at spatial overlap with risk variables. They  
268 analyzed elk data from 2012 – 2015, when elk densities were consistently near  $2.1 \text{ elk/km}^2$  – the lowest  
269 in our timeseries. Our model suggests that safety should be most important during this time, yet Cusack  
270 et al. (2020) found very little evidence that elk avoid wolves in time or space. This discrepancy is possibly  
271 explained by methodological differences, with their analysis mostly relying on either winter-long spatial

272 overlap or very fine scale encounters by individuals and our analysis relying on an annual snapshot of the  
273 population distribution.

## Appendix S8 – Supplemental Tables

*Table S1. Model covariates*

**Table S1.** Covariates included to model expected density ( $\lambda_{i,t}$ ). Colons (:) in the “Name” column indicate interactions. “Rationale” describes the reason we included a parameter in the model. The first four variables ( $k = 1 - 4$ ) measure fixed conditions that are known to affect elk density but were not the main target of inference. Variables including biomass ( $k = 5 - 6$ ) are the food variables and variables including openness or roughness ( $k = 7 - 22$ ) are the safety variables. DDHS = density-dependent habitat selection.

Index ( $k$ )	Name	Description	Rationale	
1	SWE	Snow-water equivalent	Control for autecological conditions.	
2	Elev	Elevation		
3	cos(Asp)	Cosine of aspect, N-S component		
4	sin(Asp)	Sine of aspect, E-W component		
5	Biomass	Above ground biomass of grasses and forbs from previous growing season (log-transformed)	Measure DDHS for food.	
6	Biomass:Dens	Interaction between biomass and log(density)		
7	Open	Openness (1 - % tree)	Measure DDHS for safety.	
8	Rough	Roughness		
9	Open <sup>2</sup>	Open-squared		
10	Rough <sup>2</sup>	Rough-squared		
11	Open:Dens	Interaction between safety and log(density)		
12	Rough:Dens			
13	Open <sup>2</sup> :Dens			
14	Rough <sup>2</sup> :Dens			
15	Open:Wolf	Interaction between safety and wolf density	Measure change in perceived safety with predator density; corroborates safety variables as biologically meaningful measures of predation risk.	
16	Rough:Wolf			
17	Open <sup>2</sup> :Wolf			
18	Rough <sup>2</sup> :Wolf			
19	Open:Cougar	Interaction between safety and cougar density		
20	Rough:Cougar			
21	Open <sup>2</sup> :Cougar			
22	Rough <sup>2</sup> :Cougar			

*Table S2. ANOVA for residuals by month*

**Table S2.** ANOVA for model residuals by month. We examined whether the date of the survey affected model residuals. “Day-of-year Comparison” indicates the two compared dates, and “Residual Difference” indicates the difference in mean residuals between the two dates. The p-value was adjusted using Tukey’s Honest Significant Difference test. All p-values are  $> 0.23$ , showing that residuals do not differ between any pairs of dates. These same data are visualized in Fig. S3.

Day-of-year Comparison	Residual Difference	p-value (adjusted)
2-1	-1.44	0.909
5-1	-1.69	0.743
15-1	-1.98	0.239
18-1	-0.77	1.000
20-1	-1.46	0.900
30-1	-2.26	0.252
45-1	-0.90	0.999
49-1	-0.44	1.000
57-1	-2.19	0.304
67-1	-1.87	0.589
350-1	-1.03	0.995
355-1	-0.71	1.000
358-1	-0.06	1.000
364-1	-0.94	0.998
5-2	-0.26	1.000
15-2	-0.54	1.000
18-2	0.67	1.000
20-2	-0.02	1.000
30-2	-0.82	1.000
45-2	0.54	1.000
49-2	1.00	0.997
57-2	-0.75	1.000
67-2	-0.43	1.000
350-2	0.41	1.000
355-2	0.73	1.000
358-2	1.38	0.933
364-2	0.50	1.000
15-5	-0.28	1.000

18-5	0.92	0.998
20-5	0.24	1.000
30-5	-0.57	1.000
45-5	0.80	1.000
49-5	1.25	0.970
57-5	-0.50	1.000
67-5	-0.17	1.000
350-5	0.67	1.000
355-5	0.99	0.997
358-5	1.64	0.788
364-5	0.75	1.000
18-15	1.21	0.928
20-15	0.52	1.000
30-15	-0.29	1.000
45-15	1.08	0.971
49-15	1.53	0.677
57-15	-0.21	1.000
67-15	0.11	1.000
350-15	0.95	0.991
355-15	1.27	0.897
358-15	1.92	0.285
364-15	1.04	0.980
20-18	-0.69	1.000
30-18	-1.49	0.881
45-18	-0.13	1.000
49-18	0.33	1.000
57-18	-1.42	0.917
67-18	-1.10	0.991
350-18	-0.26	1.000
355-18	0.06	1.000

358-18	0.71	1.000
364-18	-0.17	1.000
30-20	-0.81	1.000
45-20	0.56	1.000
49-20	1.01	0.996
57-20	-0.73	1.000
67-20	-0.41	1.000
350-20	0.43	1.000
355-20	0.75	1.000
358-20	1.40	0.926
364-20	0.52	1.000
45-30	1.37	0.938
49-30	1.82	0.632
57-30	0.07	1.000
67-30	0.40	1.000
350-30	1.24	0.973
355-30	1.55	0.845
358-30	2.21	0.292
364-30	1.32	0.952
49-45	0.46	1.000
57-45	-1.29	0.960
67-45	-0.97	0.997
350-45	-0.13	1.000
355-45	0.19	1.000

358-45	0.84	0.999
364-45	-0.04	1.000
57-49	-1.75	0.697
67-49	-1.42	0.915
350-49	-0.59	1.000
355-49	-0.27	1.000
358-49	0.39	1.000
364-49	-0.50	1.000
67-57	0.32	1.000
350-57	1.16	0.984
355-57	1.48	0.887
358-57	2.13	0.349
364-57	1.25	0.970
350-67	0.84	0.999
355-67	1.16	0.985
358-67	1.81	0.642
364-67	0.93	0.998
355-350	0.32	1.000
358-350	0.97	0.997
364-350	0.09	1.000
358-355	0.65	1.000
364-355	-0.23	1.000
364-358	-0.88	0.999

*Table S3. Gelman-Rubin Statistics*

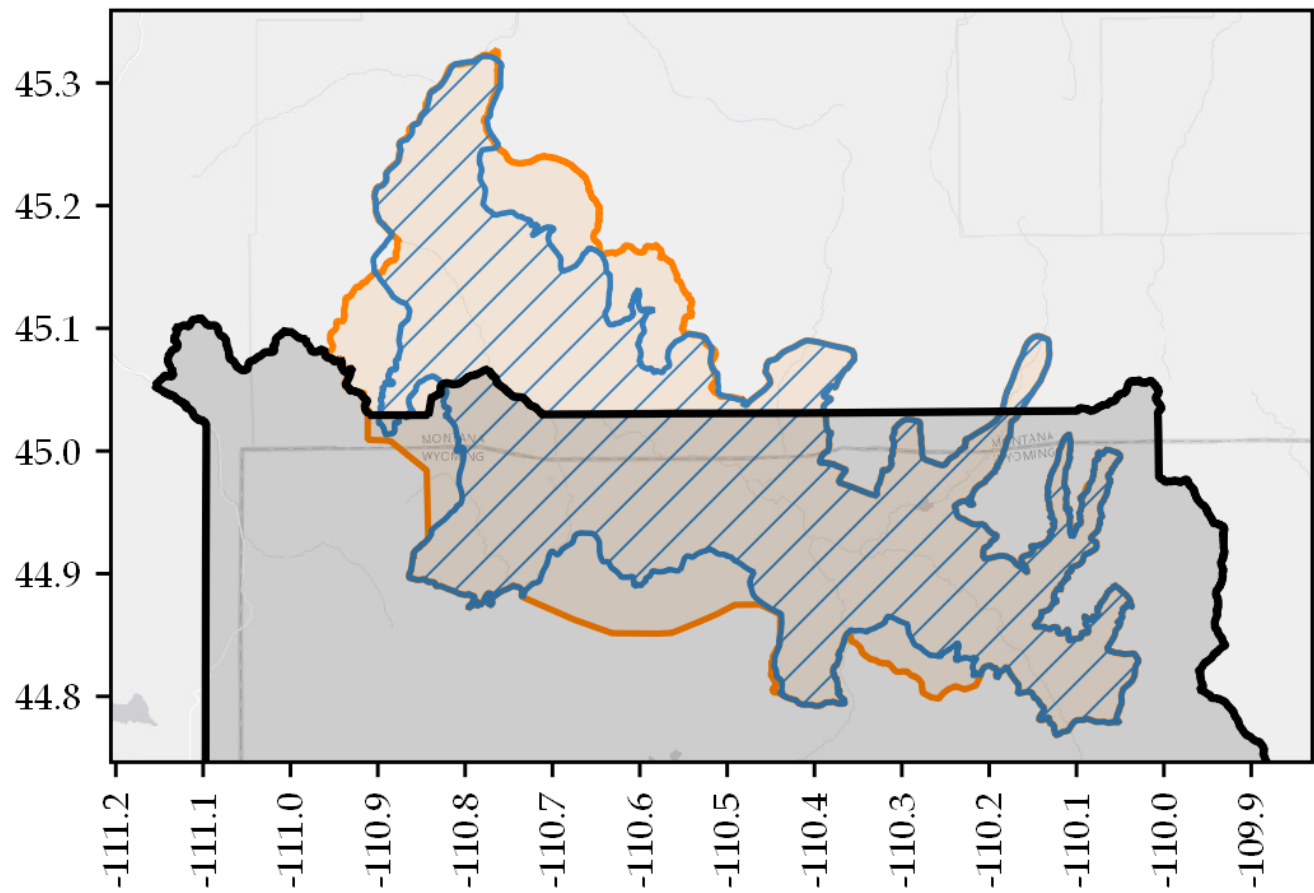
**Table S3.** Gelman-Rubin statistics for MCMC convergence diagnosis. Values for potential scale reduction factors (PSRFs) close to 1 indicate model convergence. All PSRFs for top-level parameters were estimated to be < 1.02, indicating good convergence. Parameter names match the NIMBLE code, which is available on GitHub and archived on Zenodo (doi:10.5281/zenodo.6687905). The column “Point est.” gives the estimated PSRF, and the column “Upper C.I.” gives the upper bound on the PSRF.

Parameter	Point est.	Upper C.I.
beta[1]	1.00	1.00
beta[2]	1.01	1.02
beta[3]	1.00	1.01
beta[4]	1.00	1.02
beta[5]	1.00	1.00
beta[6]	1.00	1.00
beta[7]	1.00	1.00
beta[8]	1.00	1.00
beta[9]	1.00	1.00
beta[10]	1.00	1.00
beta[11]	1.00	1.00
beta[12]	1.00	1.00
beta[13]	1.00	1.00
beta[14]	1.00	1.00
beta[15]	1.00	1.00
beta[16]	1.00	1.00
beta[17]	1.00	1.00
beta[18]	1.00	1.00
beta[19]	1.00	1.01
beta[20]	1.00	1.01
beta[21]	1.00	1.00

beta[22]	1.00	1.00
eta_out[1]	1.00	1.00
eta_out[2]	1.00	1.00
eta_out[3]	1.00	1.00
eta_out[4]	1.00	1.00
eta_out[5]	1.00	1.01
eta_out[6]	1.01	1.02
eta_out[7]	1.00	1.01
eta_out[8]	1.00	1.00
eta_out[9]	1.00	1.00
eta_out[10]	1.00	1.00
eta_out[11]	1.00	1.00
eta_out[12]	1.00	1.00
eta_out[13]	1.00	1.01
eta_out[14]	1.00	1.00
eta_out[15]	1.00	1.01
eta_out[16]	1.00	1.00
mu_out	1.00	1.01
nb_r	1.00	1.00
rho	1.00	1.01
sigma	1.02	1.06
sigma_out	1.00	1.00

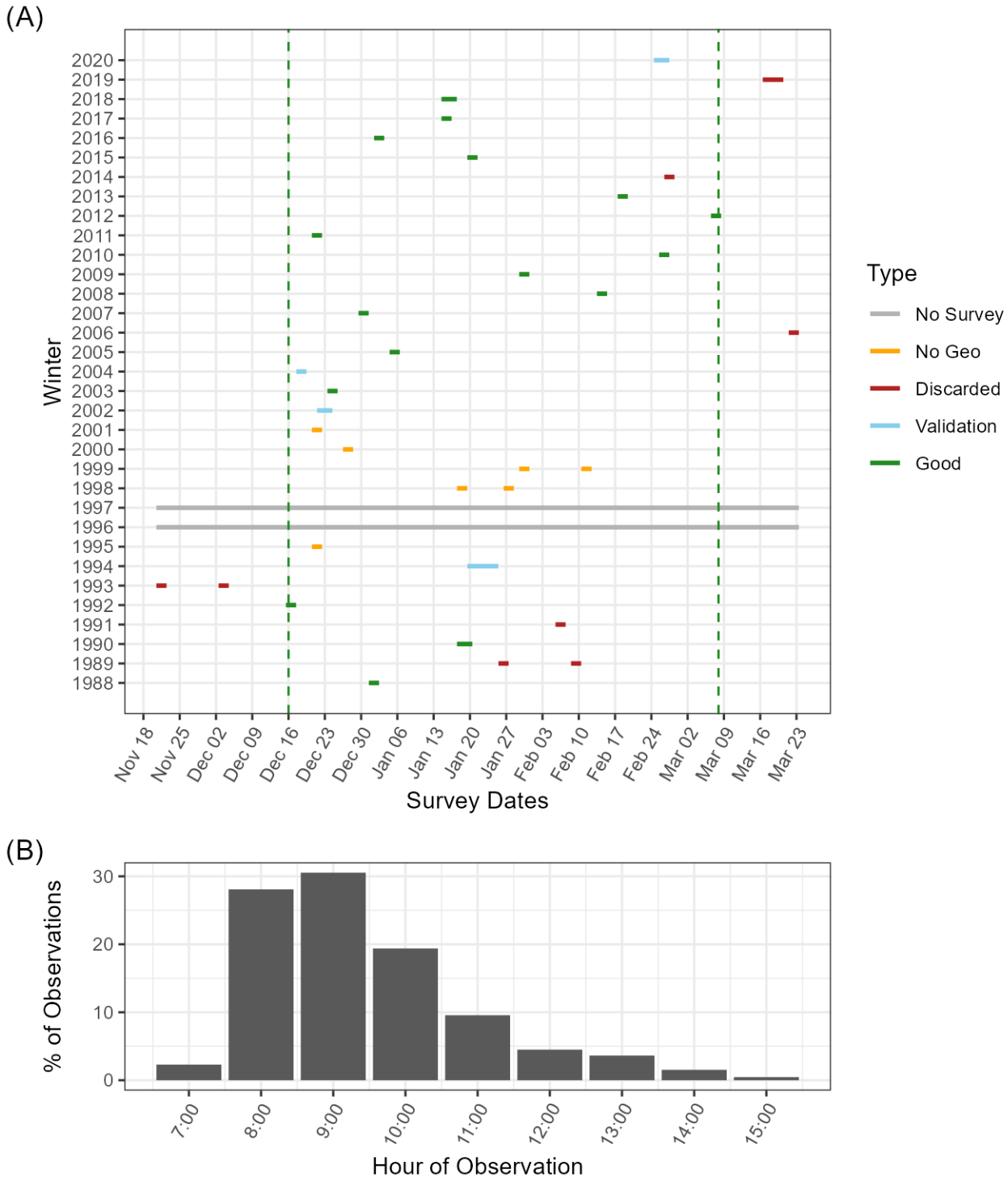
## Appendix S9 – Supplemental Figures

*Figure S1. Study area map*



**Figure S1.** Study area showing historic northern Yellowstone elk winter range polygon (blue; 1,520 km<sup>2</sup>), and our adjusted winter range polygon (orange; 1,900 km<sup>2</sup>), and the boundary of Yellowstone National Park (black).

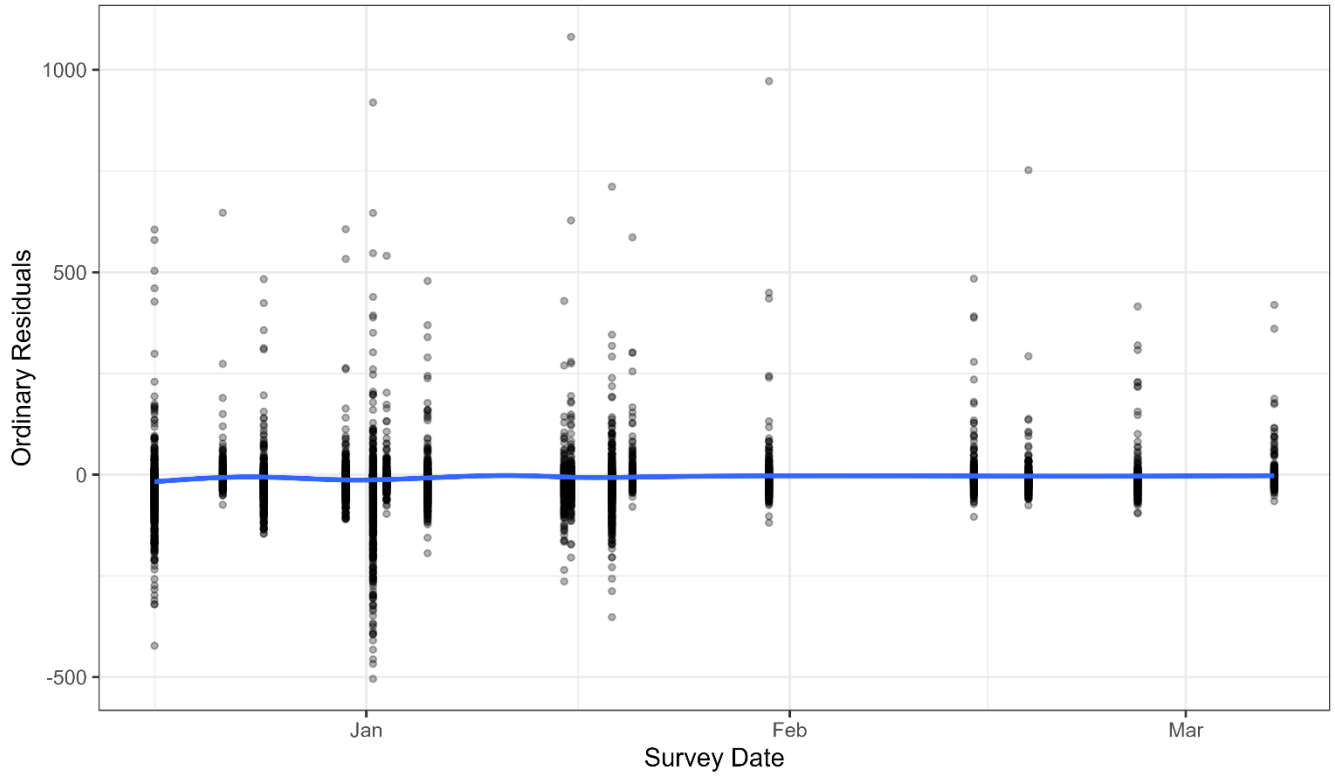
**Figure S2.** Survey dates and times



**Figure S2.** We fit our model using data collected from 1988 to 2020. Locations of elk groups ( $\geq 1$  elk) and their sizes were recorded during aerial surveys from fixed-wing aircraft. (A) Surveys used in the model were conducted between December and March in each year. In 1996 and 1997, no survey occurred (gray horizontal lines). Counts for 1989, 1991, 2006, and 2014 are considered unreliable as a census of the population and so were discarded (red lines). The count for 2019 was conducted via helicopter late in the season and was also discarded (red line). Georeferenced group data are unavailable between 1995 and 2001 and could not be included in the model (orange). In 1994, 2002, and 2004, georeferenced group data are only available inside of Yellowstone National Park (data from the State of Montana were not

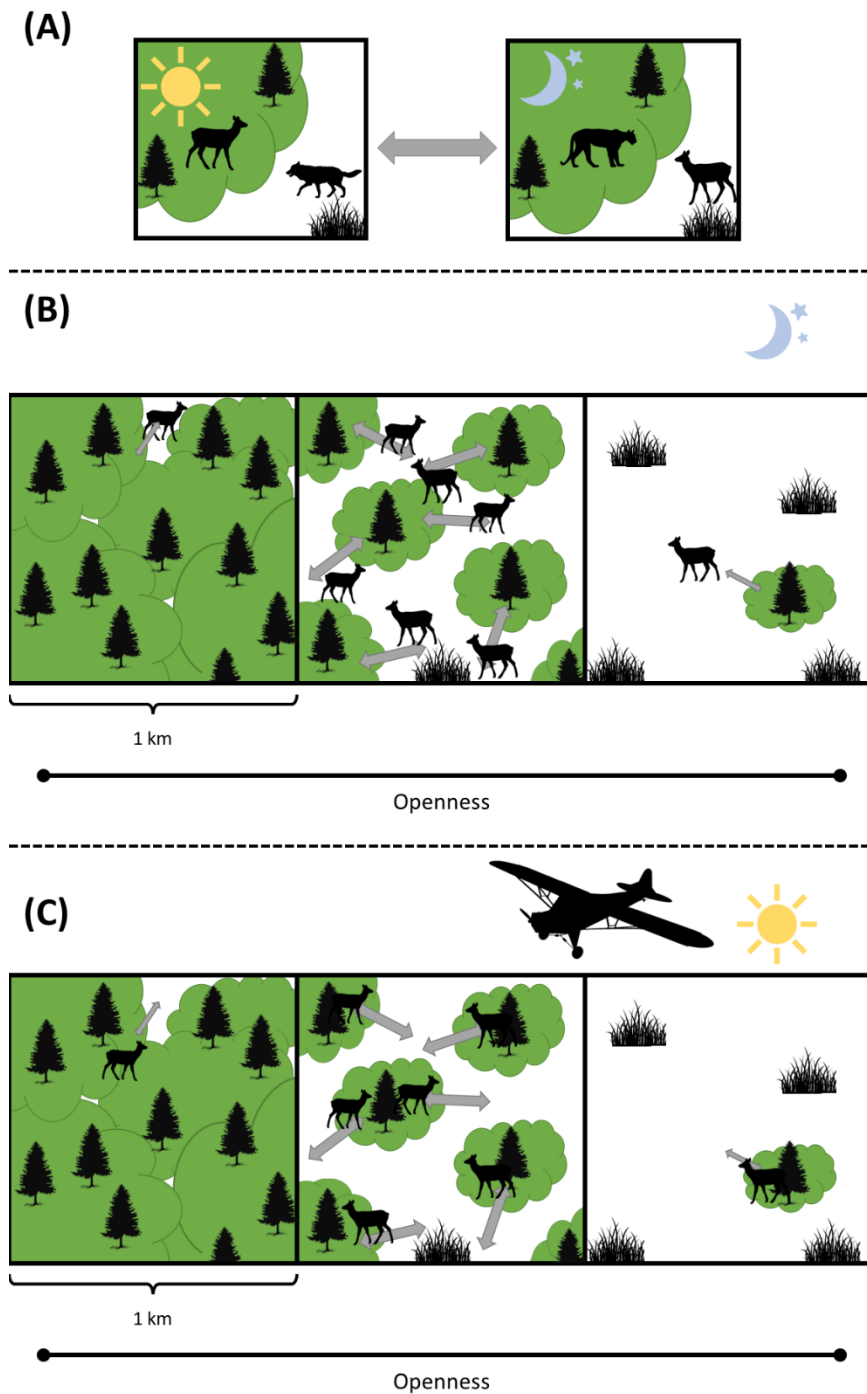
georeferenced); these data were not included for model fitting but were included for model validation (blue lines). A full census was available for 2020, but we chose to withhold it as an additional validation year with complete georeferencing (blue line). All other years were used for model fitting (green lines). All surveys used in modeling fall between December 16 and March 7 (vertical green dashed lines). (B) Surveys were conducted primarily in the morning hours, and 92% of group observations were recorded between 08:00 and 12:00.

**Figure S3.** Residuals by survey month



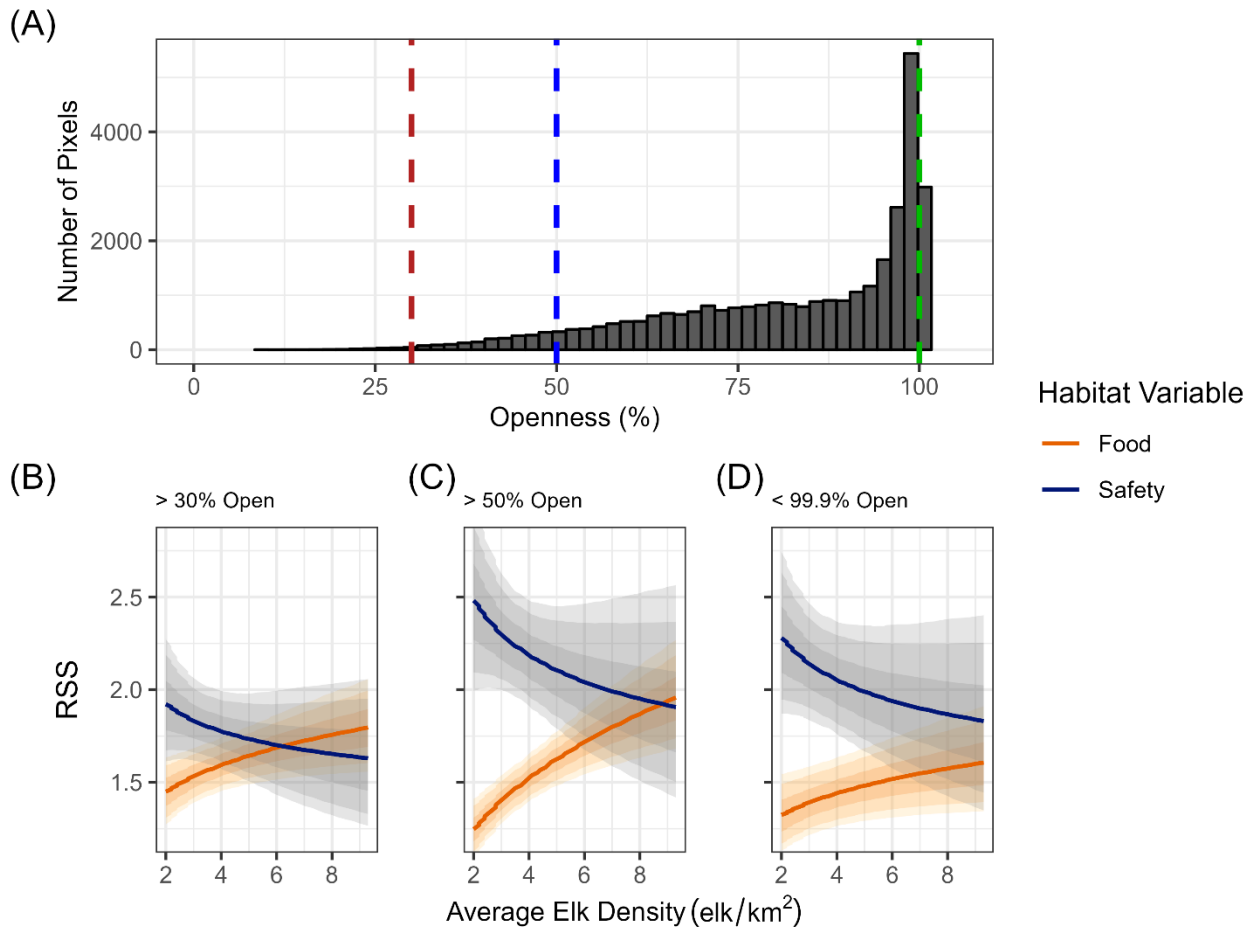
**Figure S3.** Ordinary model residuals by survey month. We checked that variable survey dates (Fig. S2A) did not introduce a bias in our model. We plotted the model's ordinary residuals as a function of survey date, and visually checked for any pattern by fitting a very flexible GAM smoothing line, which could have picked up subtle non-linear differences by date (blue line). The fitted line shows no obvious pattern, which was confirmed by an ANOVA and Tukey's HSD test (Table S2).

**Figure S4. Scale of analysis**



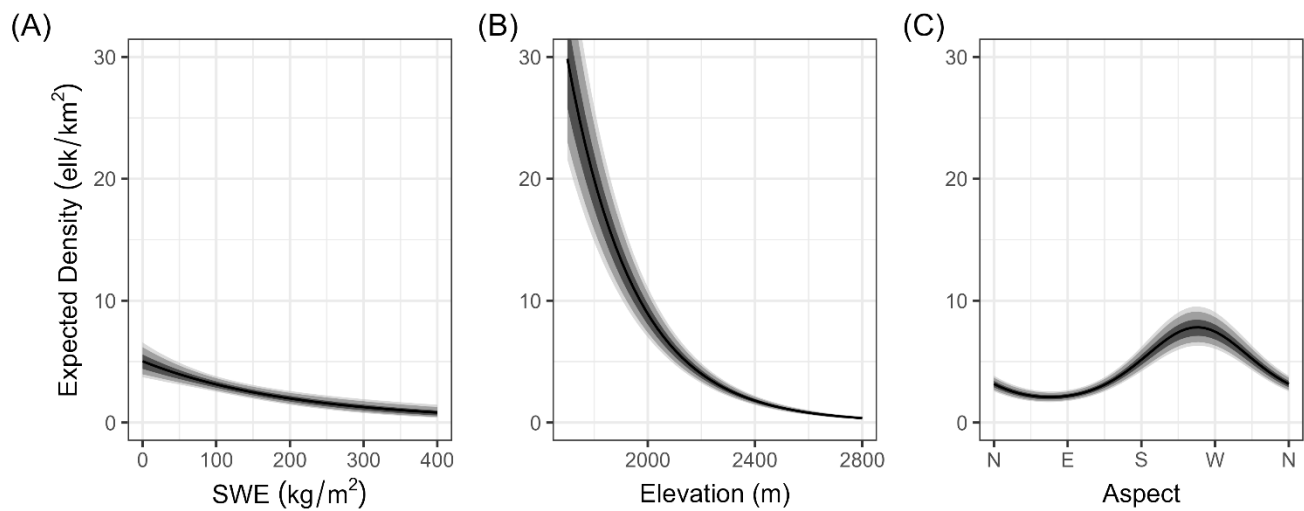
**Figure S4.** Spatial scale of analysis. We rasterized all counts and covariates on the same 1 km x 1 km raster grid. We chose this coarse spatial scale to average over daily elk movements and capture the population-level distribution. Elk are known to move between habitats characterized by openness (depicted here) and roughness (not depicted) at different times of day to manage their risk from both wolves and cougars. In terms of openness, risk from wolves is greatest in habitats with little or no tree cover, whereas risk from cougars is greatest in forested habitats. (A) Elk spend daytime hours, when wolves are more active, in forested habitats (left panel), whereas they spend nighttime hours, when cougars are more active, in open habitats (right panel). (B and C) In one-km pixels across a gradient of varying openness, elk are limited by the heterogeneity of the pixel, with gray bidirectional arrows representing diel shuttling. The gray arrows are identical between (B) and (C), but (B) shows the nighttime position of elk and (C) shows the daytime position of elk. At the 1-km scale, we hypothesized that elk density would be greatest in habitats with intermediate openness (middle panel of B and C), which provide multiple patches that facilitate this diel shuttling behavior. Aerial surveys occur during daytime (C).

**Figure S5. Openness sensitivity analysis**



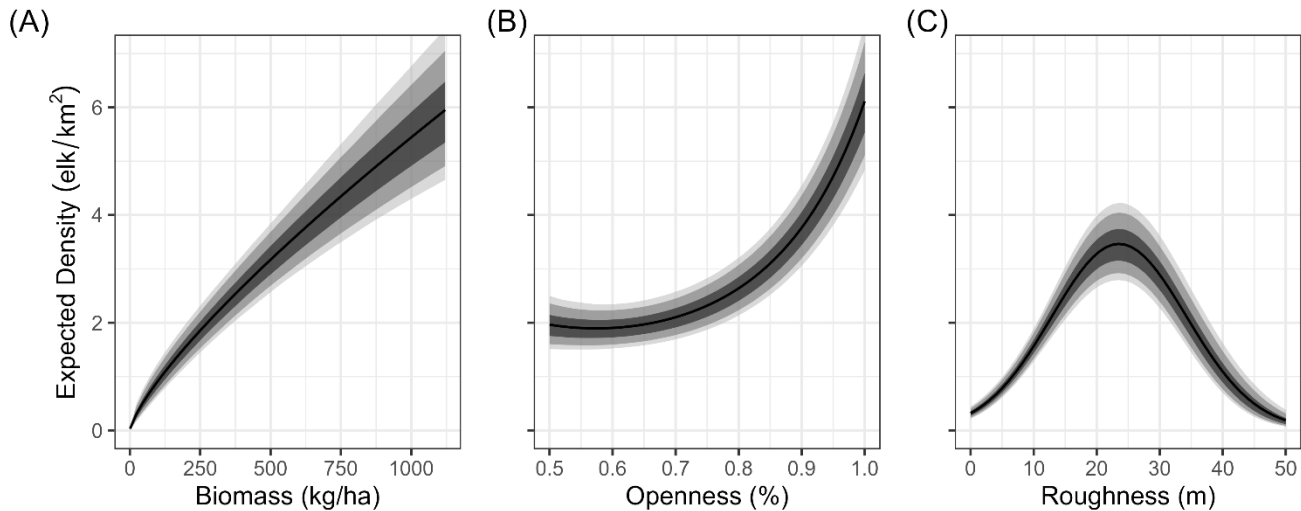
**Figure S5.** We conducted a sensitivity analysis to assess the potential impact of imperfect detection of elk due to low openness (high forest canopy cover) on our results. We discarded any pixels from the landscape that fell outside a particular openness cutoff in any year and refitted the model. (A) We defined three openness cutoffs, where the pixels retained for model fitting were  $>30\%$  open (95% of original data retained; red dashed line),  $>50\%$  open (70% of original data retained; blue dashed line), and  $<99.9\%$  open (73% of original data retained; green dashed line). (B) Results for the  $>30\%$  cutoff, (C)  $>50\%$  cutoff, and (D)  $<99.9\%$  cutoff were qualitatively similar to the results presented in the main text: we found positive DDHS for food, negative DDHS for safety, and a switch in their relative importance. DDHS for food equals the RSS for a 1-SD change in herbaceous biomass, and DDHS for safety equals a 0.5-SD change in openness and a 0.5-SD change in roughness. See main text for details. In panels (B) – (D), solid lines are posterior mean estimates and shaded envelopes are 50%, 80%, and 90% credible intervals.

**Figure S6. Mean effect of conditions**



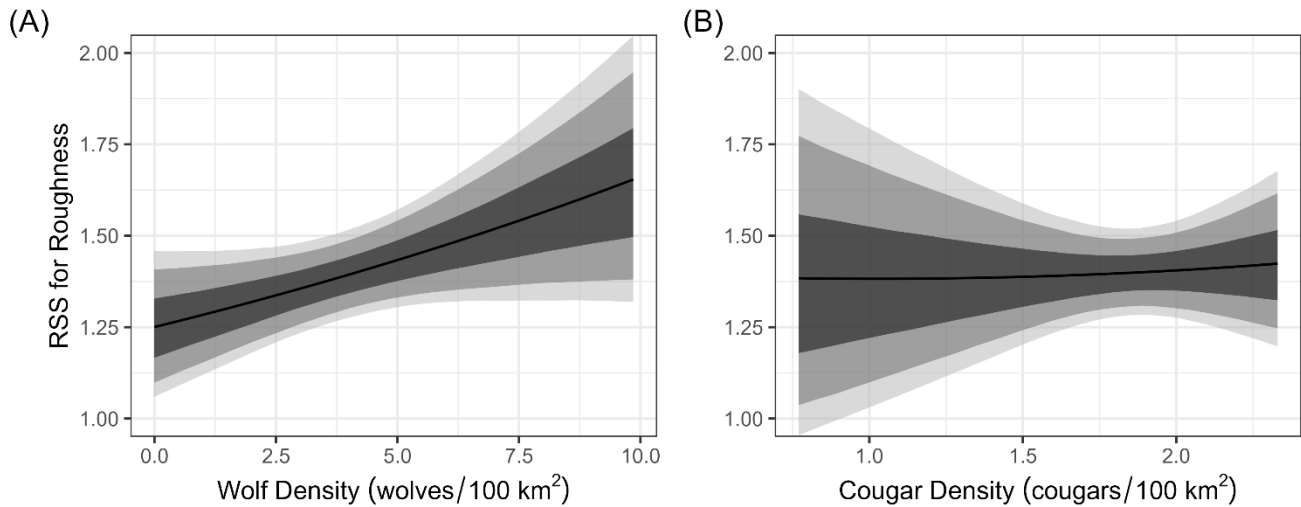
**Figure S6.** Mean effect of snow-water equivalent (SWE), elevation, and aspect on expected elk density during winter (December–February), with all other variables held at their mean. (A) Elk density was greatest in pixels with low snow, (B) low elevation, and (C) southwest aspects. Solid black lines show the mean effects, and the shaded gray envelopes show the 50%, 80%, and 90% credible intervals.

*Figure S7. Mean effect of food and safety variables*



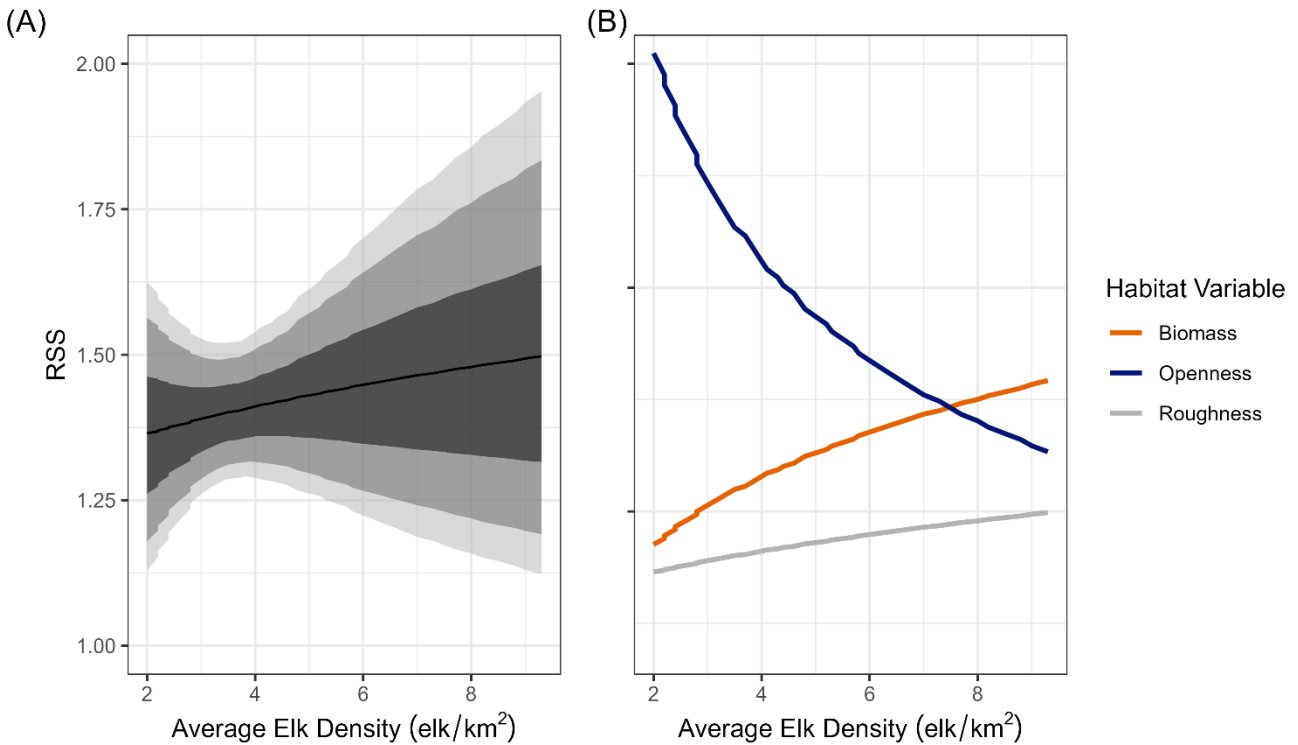
**Figure S7.** Mean effects of food and safety variables on expected elk density, with all other variables held at their overall mean. (A) Expected elk density increased with herbaceous biomass, consistent with our assertion that it is an important food resource for elk. (B) Contrary to our prediction that elk would most prefer an intermediate openness, expected elk density was greatest at 100% openness, and the relationship between openness and elk density was largely monotonic for the observed range of openness. (C) Expected elk density was greatest for a roughness of 23.5 m (intermediate, as expected). Solid black lines show the mean effects, and the shaded gray envelopes show the 50%, 80%, and 90% credible intervals.

*Figure S8. Effect of predator density on RSS for roughness*



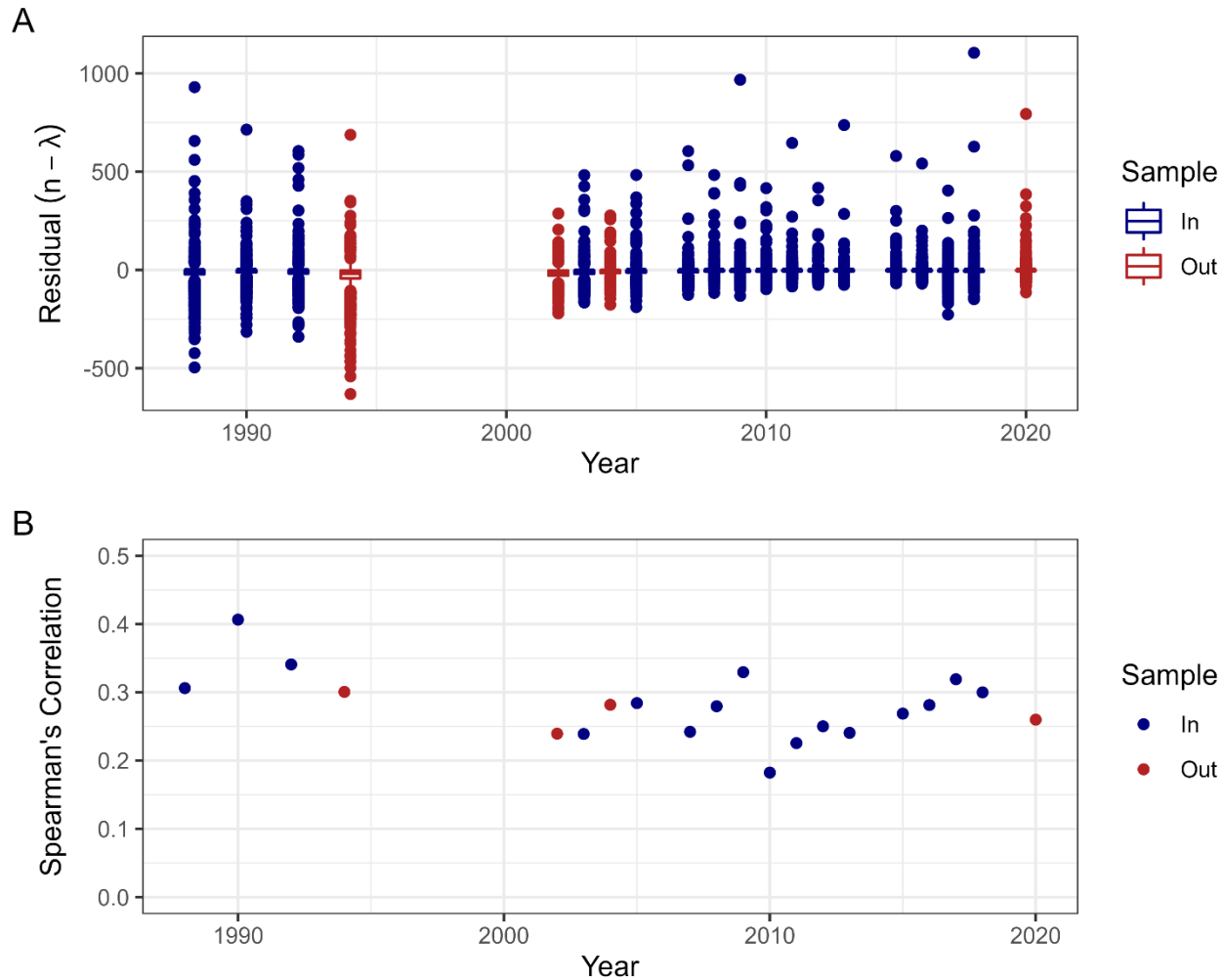
**Figure S8.** Mean effect of predator density on relative selection strength (RSS) for roughness, with all other variables held at their mean. (A) RSS for roughness increased with wolf density, in agreement with our prediction that elk would move away from the wolf habitat domain. (B) RSS for roughness did not change with cougar density, in agreement with our prediction that the wolf effect would be stronger than the cougar effect during the daytime aerial elk surveys. Solid black lines are mean effects, and shaded gray envelopes are 50%, 80%, and 90% credible intervals.

*Figure S9. DDHS for roughness*



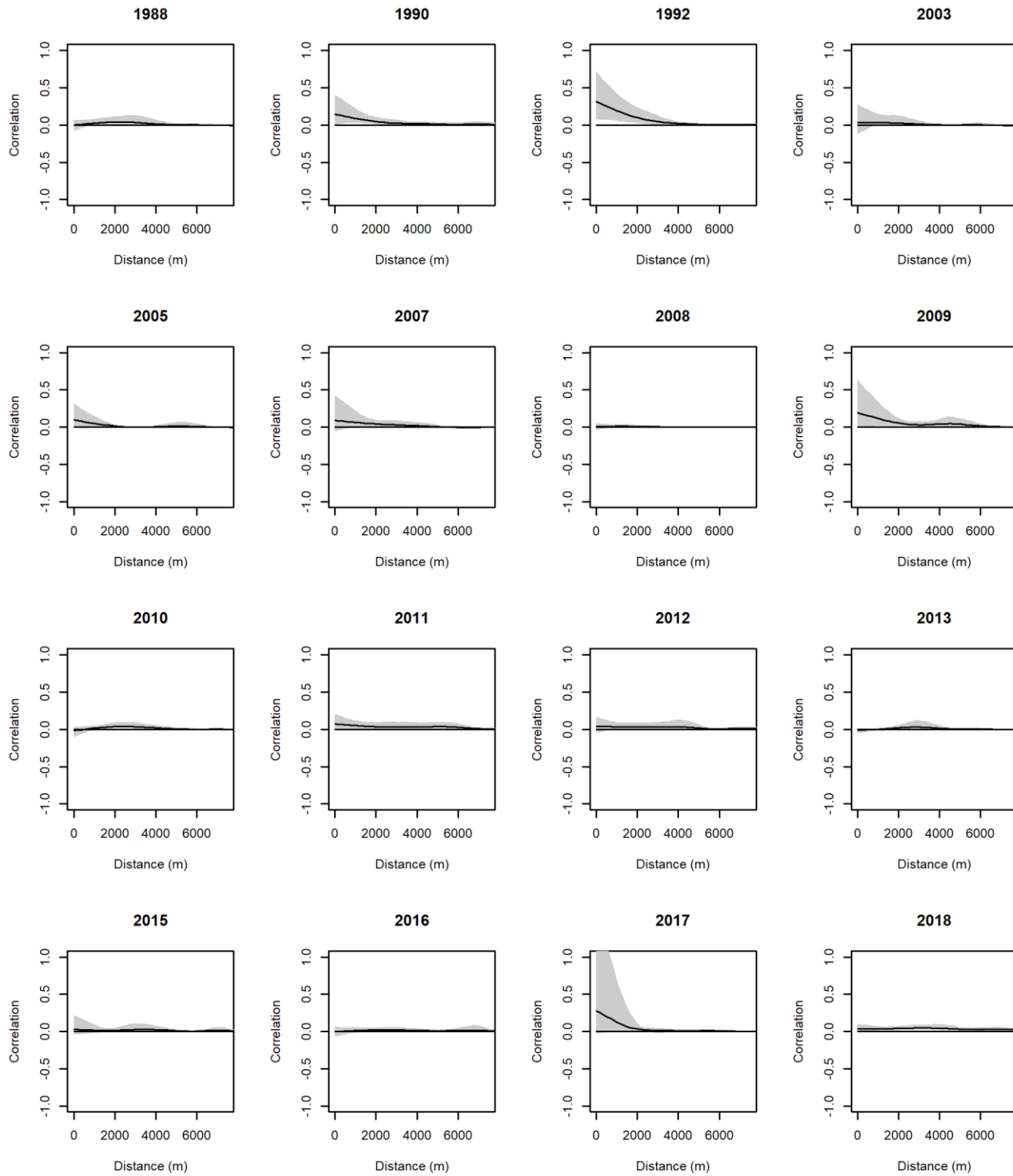
**Figure S9.** Density-dependent habitat selection (DDHS) for food and safety variables decomposed. (A) Average elk density versus relative selection strength (RSS) for a 1-SD change in roughness. The mean effect is weakly positive, but the large uncertainty indicates no support for DDHS for roughness. RSS was calculated using samples from the entire posterior distribution. Solid black line is the mean effect, and the shaded gray envelopes are the 50%, 80%, and 90% credible intervals. (B) Comparison of RSS for a 1-SD change in all food and safety variables.

*Figure S10. Model evaluation*



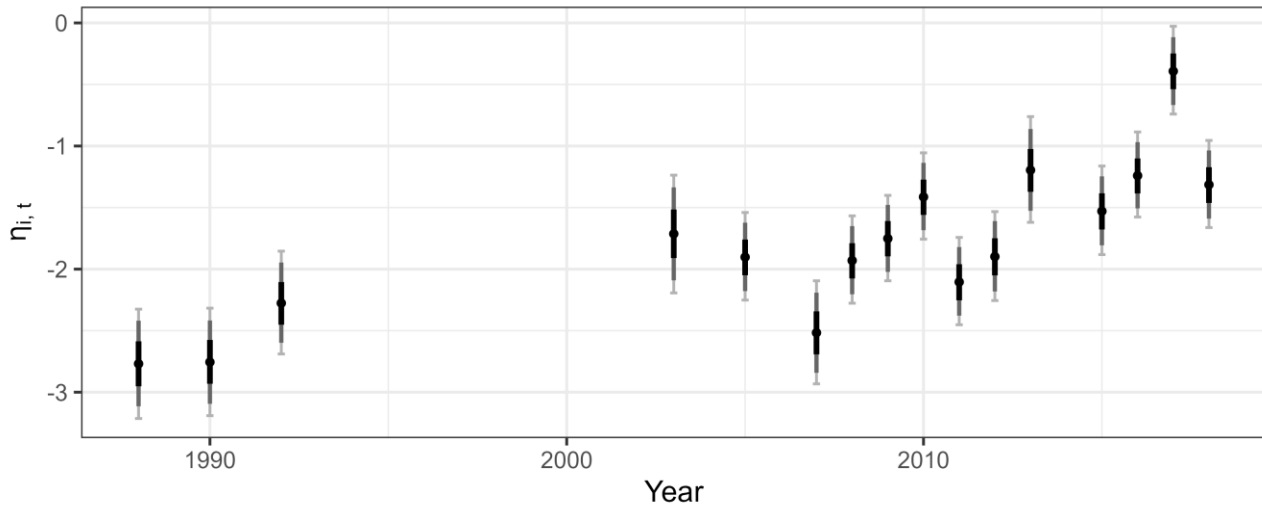
**Figure S10.** Model evaluation. We compared (A) ordinary residuals and (B) Spearman's correlation between expected and observed densities for both in-sample data (blue) and out-of-sample validation data (red). Residuals have mean near 0 in all years, showing good accuracy, but a wide spread, showing low precision. Spearman's correlation is moderate in all years (mean in-sample = 0.28, mean out-of-sample = 0.27), which shows the model has moderate power to rank pixels by abundance in both training and testing data.

**Figure S11. Residual spatial autocorrelation**



**Figure S11.** Non-parametric spline correlograms used to assess residual spatial autocorrelation. We estimated correlograms using Pearson's residuals for each pixel in each year. The scale of spatial autocorrelation is estimated as the distance where the confidence envelope first intersects the x-axis (correlation = 0). There was little to no residual spatial autocorrelation in all years, i.e., the confidence envelope crosses the x-axis near 0 m in nearly all years (1992 was the lone exception).

*Figure S12. Temporal random effect posterior estimates*



**Figure S12.** Temporal random effect ( $\eta_{i,t}$ ) over time. We included the temporal random effect to account for changing patterns of elk distribution inside versus outside Yellowstone National Park (YNP), along with a change in the late hunt outside of YNP after 2010. The temporal random effect affects pixels outside of YNP, so negative values indicate that pixels outside of YNP have lower expected elk density than would be predicted based on fixed effects alone. Points show posterior mean for each coefficient and bars show credible intervals. Black bars show 50% credible intervals, dark gray bars show 80% credible intervals, and light gray bars with end caps show 90% credible intervals.

Figure S13. Spatial random effect posterior mean

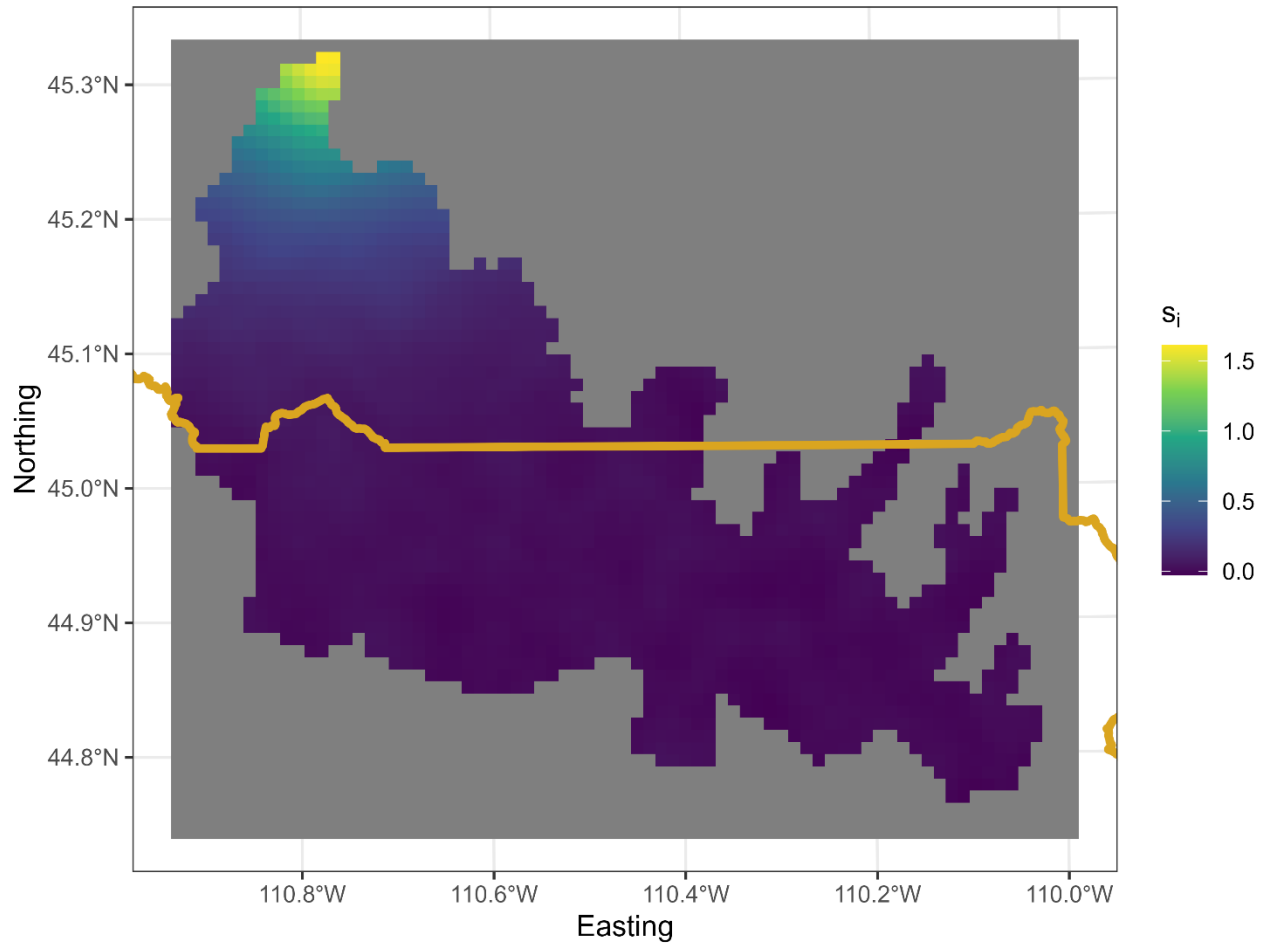
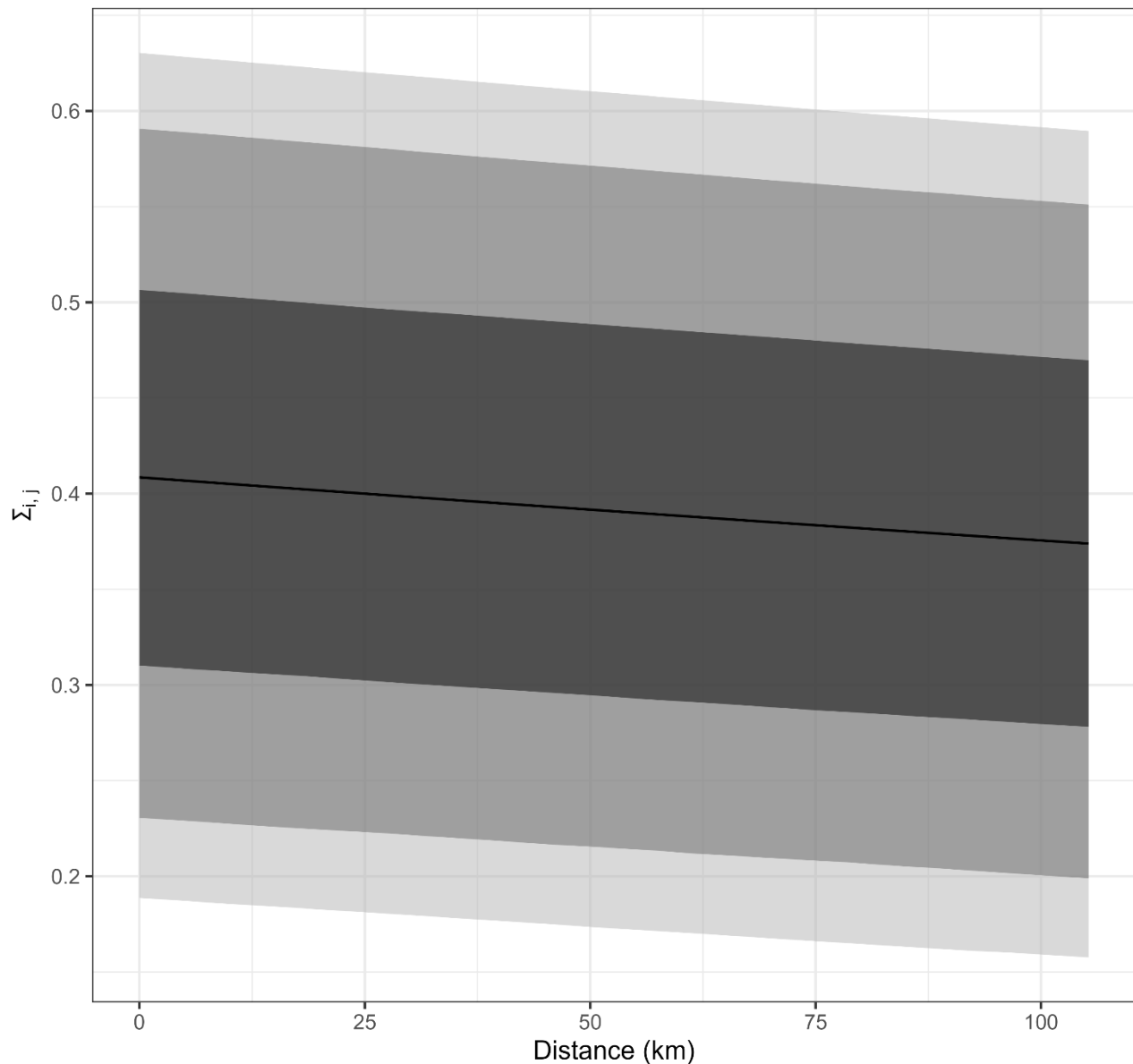


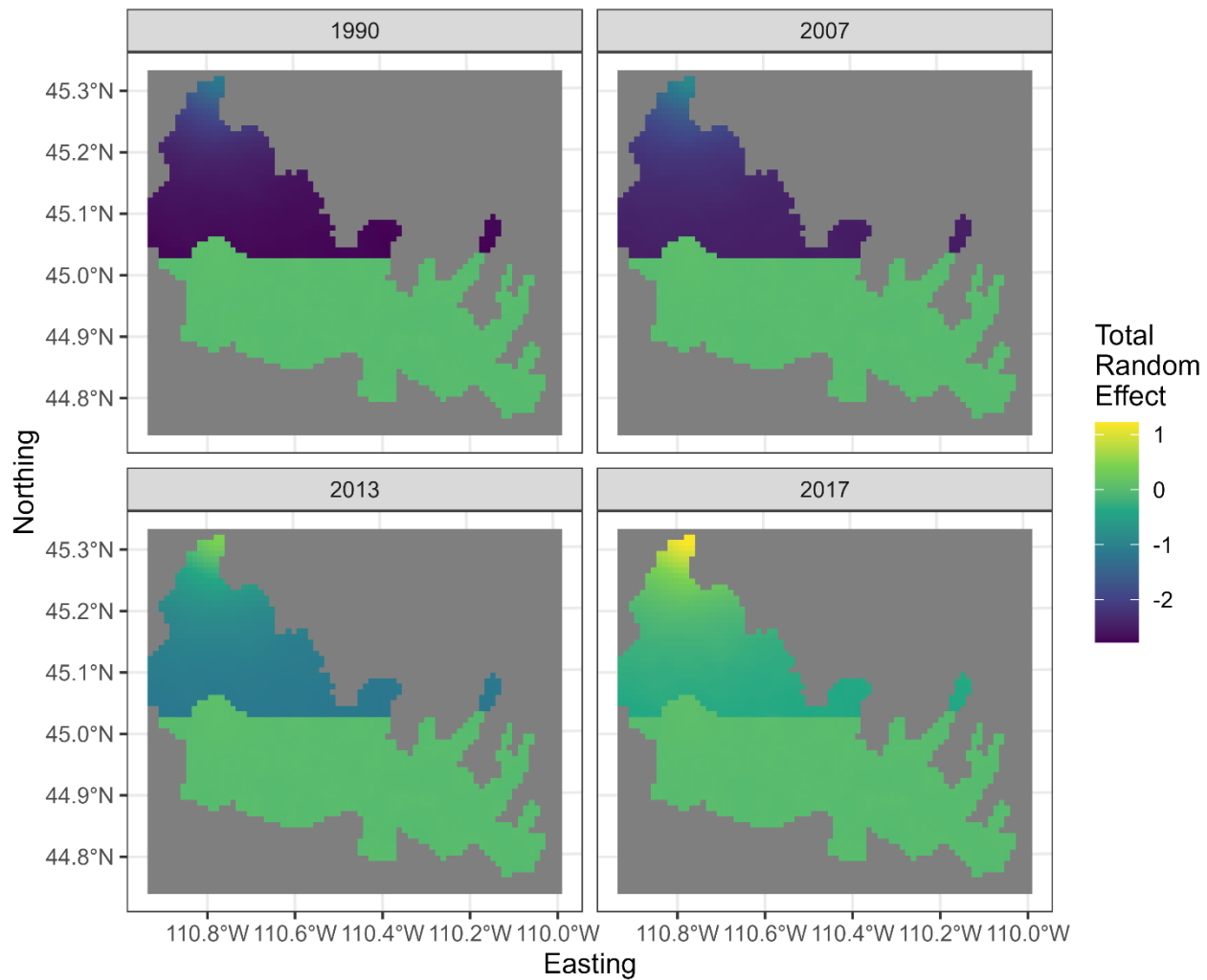
Figure S13. Posterior mean of estimated spatial random effect,  $s$ .

*Figure S14. Fitted spatial covariance function*



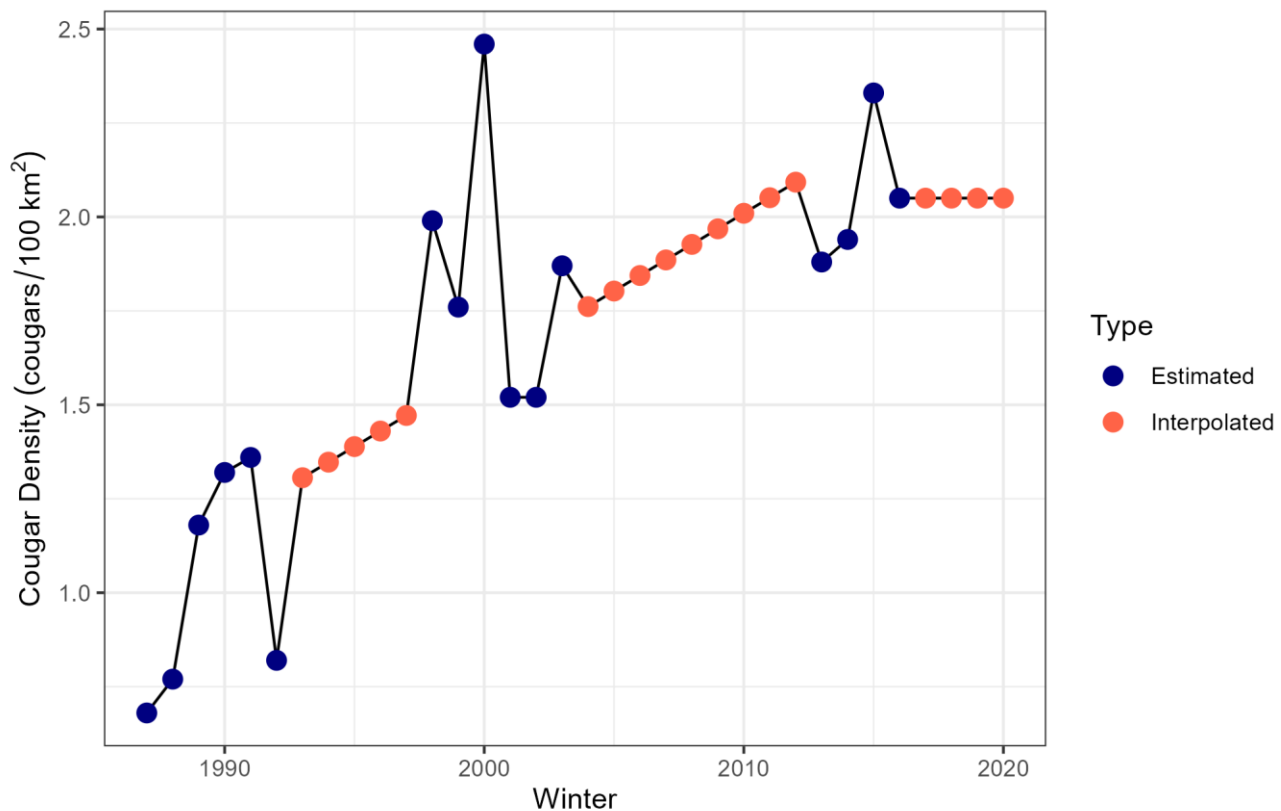
**Figure S14.** Fitted spatial covariance as a function of distance for the spatial random effect.

*Figure S15. Combined temporal and spatial random effects.*



**Figure S15.** The temporal random effect was defined as operating outside Yellowstone National Park (YNP) and the estimated values were always negative. The spatial random effect could only take on positive values, but it could operate anywhere in the study area. The estimated values were largest outside of YNP. The negative effect of the temporal random effect and the positive effect of the spatial random effect thus acted in opposite directions and almost entirely outside YNP, with expected elk density inside YNP almost completely captured by fixed effects. Examples of the combined effect for 1990, 2007, 2013, and 2017. See Fig. S12 for temporal random effect alone in each year and Fig. S13 for spatial random effect alone.

*Figure S16. Estimated and interpolated cougar densities.*



**Figure S16.** Cougar densities were estimated from field surveys in three phases: Phase 1 (1987 – 1993; Ruth *et al.* 2019), Phase 2 (1998 – 2004; Ruth *et al.* 2019), and Phase 3 (2014 – 2017; Anton 2020). Values between Phases 1 and 2 and between Phases 2 and 3 were linearly interpolated from the entire dataset. Values after Phase 3 (2018 – 2020) were assigned the mean of Phase 3.

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