Evaluation of infection risk of two endoparasites to moose (Alces alces) in the Adirondack Park of New York

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Introduction

Moose (Alces alces) populations are in decline in parts of their range in North America such as in Minnesota, Vermont, and New Hampshire in the United States of America and Nova Scotia, Alberta, Saskatchewan, Manitoba, and Ontario in Canada (Lankester 2018; Timmermann and Rodgers 2017). Some researchers point to parasite infection from liver fluke (Fascioloides magna) and/or brainworm (Parelaphastrongylus tenuis) as contributing factors to these declines (Murray et al. 2006; Lankester 2010, 2018; Wünschmann et al. 2015), although this claim is controversial (Lenarz 2009) and in some areas may coincide with other factors such as depredation or low reproduction rates that may disguise effects of parasite infection (Barber-Meyer and David Mech 2016; Kuzyk et al. 2018). Nevertheless, brain worm and liver fluke cause morbidity and mortality in moose populations and should be monitored.

In the Adirondacks of New York, moose populations have increased since their extirpation in the late 19th century, but their populations have not seen the same growth during the 20th century as their New England counterparts (Timmermann and Rodgers 2017). Recent estimates of moose population size indicate that there are approximately 600 to 1000 moose in the Adirondacks (Wong, Fuller, and Royle, n.d.), a range which does not indicate substantial change from previous assertions of population size (Hickey 2008). Factors that may be limiting the population are not established, but the focal parasites *P. tenuis* and *F. magna* have been observed in moose necropsies with moderate frequency (REF Schuler and/or DEC).

Brain worm and liver fluke are endoparasites that infect white-tailed deer ($Odocoileus\ virginianus$) and moose. Infections by $P.\ tenuis$ are benign to white-tailed deer, but cause severe neurological disorders in moose that often end in death (Pybus 2001; Lankester 2010).

Materials and Methods

Study area

We surveyed within the Adirondack Park of the state of New York. It is an area of approximately 24,000 km², composing approximately 17% of the state of New York. Elevation ranges from 100m - 1600m. The Adirondack Park contains primarily upland forest habitat dominated by such species as American beech (Fagus grandifolia), red spruce (Picea rubens), hemlock (Tsuga canadensis), sugar maple (Acer saccharum), striped maple (Acer pensylvanicum), and balsam fir (Abies balsamea). Approximately 20% of the park is composed of wetland habitats including open river corridors, floating bogs, and large open bogs dominated by conifers (Hickey 2008).

Field data collection

White-tailed deer fecal pellet groups were collected opportunistically during the years of 2016 and 2017 for the purpose of parasite quantification during a larger survey investigating the population size of moose in the park. The moose study surveyed triangular transects approximately 3 km long, selected by cluster random sampling. The clustering design of the transects were optimized for a spatial capture-recapture survey, including 3 transects per cluster separated by approximately 2-4 km. **INSERT FIGURE FOR MOOSE TRANSECTS**

In 2018, a single-observer distance sampling protocol was implemented to quantify deer fecal group abundance in addition to measuring parasite intensity. Along each moose transects, 6 distance sampling transects each 100m long were spaced 200m apart from each other. **INSERT FIGURE FOR DEER TRANSECTS** A total of **N** transects were sampled in 2018. At the specified start point, field technicians extended a 100m tape to serve as the center line, and made observations along its length. Any number of scat pellets were considered to constitute a fecal group, and the approximate number was recorded. When a fecal group was detected, its distance perpendicular from the center line, and parallel along the center line was recorded. The

perpendicular distance is the focal variable of distance sampling, but the parallel distance provided highly precise location coordinates of the scats – because the start point of the transect was recorded with greater accuracy (due to longer GPS averaging time), the coordinates of the scats could be refined by translating the start point in space by the angle of the transect, and the parallel distance away from the start point.

Parasitological analyses

The fecal samples were analyzed by the Cornell Animal Health and Diagnostic Center using several methods. For *F. magna*, Flukefinder, a modified Baermann technique, and fecal quantitative flotation were used. For *P. tenuis*, the modified Baermann, and fecal quantitative flotation methods were implemented. After 2016, the performant method for *F. magna* was determined to be Flukefinder, and for *P. tenuis* the performant method was the modified Baermann technique. This was assessed by **HOW? ASK KRYSTEN**, and validated post-hoc by an occupancy analysis.

Flukefinder

Needs to be updated by relevant members; Flukefinder methodology not in shared folder

Modified Baermann

Fecal flotation

Statistical Model

For analysis of relative risk of infection to moose, we implemented a hierarchical Bayesian modeling framework analyzed with the R packages INLA, which uses integrated nested Laplace approximation to approximate the posterior distribution of model parameters, and inlabru which extends INLA by integrating thinning of a point process – in this context, the fecal groups are modeled as a point process thinned by imperfect detection which is modeled by distance sampling. Our measure of relative risk of infection to moose is predicted parasite abundance spatially referenced by the following model.

$$\widehat{\Lambda}(s) = \widehat{\lambda}(s) * \widehat{\delta}(s)$$

where, $\hat{\Lambda}(s)$ is the estimated parasite abundance at spatial unit s, which is the product of the per-fecal-group-sample abundance $\hat{\lambda}(s)$ (or, intensity in the epidemiological sense), and the estimated fecal group abundance $\hat{\delta}(s)$.

Parasite intensity model

Let $S = \{s_i, i = 1, ..., n\} \subset \Omega$ be the *n* distinct locations where white-tailed deer fecal groups were observed within the observation region $\Omega \subseteq \mathbb{R}^2$, and let y_i be the observed parasite count within the fecal sample.

Parasites observed within each scat sample are ≥ 0 , and due to the high prevalence of 0's, we considered a negative binomial distribution in addition to the Poisson. On any bounded region $B \subseteq \Omega$, observations are thus modeled as follows:

$$y_i \sim \text{Poisson}(\mu_i)$$

$$\log(\mu_i) = \beta_0 + \beta_1 * x_1(s_i) + \dots + \beta_p * x_p(s_i) + \Psi_1(s_i)$$

$$\Psi_1(s_i) \sim GP(0, \Sigma)$$
(1)

Where, y_i represent the parasites observed per fecal group sample, μ_i is the mean function modeled dependent upon spatial covariates $x_j(s)$, and $\Psi_1(s)$ is a random Gaussian process with mean 0 and covariance matrix Σ . The covariance function used is the Matern:

$$cov[\Psi(s), \Psi(s')] = \sigma_{\Psi}^2 \kappa ||s' - s|| \mathcal{K}_1(\kappa ||s' - s||)$$

where τ, κ are variance and range scaling parameters, respectively, $\sigma_{\Psi}^2 = 1/(4\pi\kappa^2\tau^2)$ is the marginal variance, and \mathcal{K}_1 is the modified Bessel function of the second kind and order 1 (See Lindgren and Rue (2015) for the specific representation and estimation of the spatial random effects within INLA). The spatial random field is optional and we test models with and without it to assess support for modeling unstructured spatial dependence.

Priors

Covariates on parasite intensity

In our model, we consider the following covariate effects on parasite intensity. Coordinate data (UTM Easting and Northing) are included to model broad-scale spatial trend. We considered a derived variable of precipitation representing the total accumulation of precipitation in the year prior to the data collection date, measured in inches. Precipitation affects the survival of P. tenuis and F. magna outside of a host (i.e. as L1 larvae, and eggs in feces, respectively), and so it is anticipated that deer would be more likely to be infected in areas of higher rainfall (Pybus 2001). Additionally, gastropod activity is inhibited by drought, and so higher rainfall would suggest higher risk of infection due to a higher availability of gastropods to be ingested in the case of P. tenuis (Lankester 2018). We used the spatially interpolated PRISM dataset for precipitation (n.d.). We considered another derived variable for snowfall accumulation in the snowfall season prior to the data collection date (that is, from September until April prior to the date of survey). Similar to precipitation, we anticipate that areas that experience low snowfall prior to the survey date will exhibit a higher parasite intensity due to increased parasite survival, and increased activity of gastropod intermediate hosts. The National Weather Service's National Snowfall Analysis dataset was used to obtain spatially-referenced snowfall amounts. Distance-to-wetland was considered as a potentially important effect particularly for F. magna, as the life cycle of the parasite has a period during which metacercariae encyst on aquatic vegetation and once matured infect aquatic snail intermediate hosts. We hypothesize that intensity of infection for F. magna will be greater nearer to wetland systems.

Distance sampling model

The white tailed deer fecal groups are treated as an log Gaussian Cox point process which are thinned by imperfect observation, and the rate of detection is a parameter to be estimated. Within any bounded region $B \subseteq \Omega$, let $\Lambda(s)$ be the mean intensity function under the following distribution:

$$\begin{split} N(B) \sim \text{Poisson}(\Lambda(s)*|B|) \\ \log \Lambda(s) &= \beta_{p+1} * x_{p+1}(s) + \ldots + \beta_m * x_m(s) + \Psi_2(s) \\ &\Psi_2(s) \sim GP(0,\Sigma) \end{split} \tag{2}$$

where, N(B) is the realization of the total abundance of fecal groups within $B, x_j(s) \in x_{p+1}(s), ..., x_m(s)$ are spatial covariates that may be the same as those used in the parasite intensity model, $\Psi_2(s)$ is a Gaussian process distinct to $\Psi_1(s)$.

Covariates on deer fecal group intensity

Deer fecal group abundance is assumed to be proportional to deer density, and so factors affecting deer density should correlate with deer fecal group abundance. We do not attempt to quantify deer density per se because the density of infected fecal groups within an area more directly influences the probability that intermediate gastropod hosts will become infected, and thus risk of infection to moose.

Results

Discussion

References

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