

# <sub>1</sub> Chapter 1

## <sub>2</sub> Introduction



## <sup>3</sup> Chapter 2

# <sup>4</sup> GLMS and WinBUGS



## <sup>5</sup> Chapter 3

## <sup>6</sup> Closed population models



<sup>7</sup> Chapter 4

<sup>8</sup> Fully Spatial  
<sup>9</sup> capture-recapture models





<sup>10</sup> **Chapter 5**

<sup>11</sup> **Other observation models**



<sup>12</sup> **Chapter 6**

<sup>13</sup> **MCMC details**



## <sup>14</sup> Chapter 7

## <sup>15</sup> Goodness of Fit and stuff



## <sup>16</sup> Chapter 8

## <sup>17</sup> Covariate models





## 18 Chapter 9

# 19 Inhomogeneous Point 20 Process



## <sup>21</sup> Chapter 10

## <sup>22</sup> Open models



## <sup>23</sup> Chapter 11

## <sup>24</sup> Unmarked populations



## Chapter 12

# Spatial Capture-Recapture for Unmarked Populations

Traditional capture-recapture models share the fundamental assumption that each individual in a population can be uniquely identified when captured. This can often be accomplished by marking individuals with color bands, ear tags, or some other artificial mark that can be read in the field. For other species, such as tigers or marbled salamanders, individuals can be easily identified using only their natural markings. In a great number of cases, however, species do not possess sufficient natural markings and are too difficult to capture to make it practical to apply artificial marks. So we must throw up our hands and not study these species. End of chapter.

When capture-recapture methods are not a viable option, researchers often collect simple count data or even detection/non-detection data to estimate population parameters. These data are often analyzed using Poisson regression or logistic regression, perhaps with random effects; but when detection is imperfect, as it almost always is, these methods cannot be used to obtain unbiased estimates of population size or occurrence probability. Even when these data are used an index of abundance or occurrence, standard models may yield unreliable results when covariates affect both the state variable and detection probability. A classic example is the finding by Bibby and Buckland (1987) who reported that the probability of detecting songbirds in restocked conifer plantations decreased with vegetation height; whereas population density was positively related to vegetation height. This intuitive and common phenomenon has led to the development a vast number of methods to model population size or density while controlling for factors affecting detection probability. A review of these models is beyond the scope of this chapter, but we mention a few deficiencies of existing methods that warrant the exploration of alternatives.

Distance sampling, which we briefly introduced in chapter XXXX, is perhaps the most widely used method for estimating population density when individuals are unmarked and detection probability is less than one. This class of

methods is known to work impeccably when estimating the number of stakes in a field or the number of duck nests in a wetland. It can also work very well in more interesting situations; however, common issues such as animal movement and measurement error may result in substantial bias. In addition, traditional distance sampling methods assume that individuals are randomly located with respect to the observer and are available for detection (but see Johnson (2010); Chandler et al. (2011)). Most other methods, such as double-observer sampling and repeated counts, can be used to estimate population size, but as with traditional CR methods, it may be difficult to convert abundance estimates to density estimates because the effective area sampled is unknown. We mention these issues not to suggest that existing models do not have value—indeed we believe that they can be used to obtain reliable density estimates in many situations—rather our aim to highlight the need for alternative methods when the assumptions of existing methods cannot be met. Additionally, the model we develop in this chapter serves as the foundation for a broad class of SCR models in which all or some of the individuals cannot be uniquely identified.

In this chapter we highlight the work of Chandler and Royle (2012) who demonstrated that the individual recognition assumption of CR models is not a requirement of spatial capture-recapture models. The ability to fit SCR models to data from unmarked populations has important consequences in several respects. For one, it means that SCR models can be applied to data collected using methods like point counts in which observers record simple counts of animals at an array of survey points. This development also has important implications for traditional SCR studies because many resulting datasets include some individuals that cannot be identified due to poor photo quality or the indistinguishable natural markings.

In order to apply SCR models to data collected on unmarked animals, one requirement is critical—counts must be spatially correlated. Of course, this condition holds true in virtually all SCR models since animals are often detected at more than one trap. In fact, efficient SCR designs should try to ensure correlation in counts among neighboring traps because this is the primary source of information about the encounter rate parameter,  $\sigma$ .

## 12.1 Data Requirements and Survey Designs

## 12.2 Encounter Histories as Latent Variables

Just when you thought we ran out of things to treat as latent variables, we are now going to regard even the data itself as latent.

State model is the same as other SCR models.

It is natural to regard the encounter rate of an individual as a function of the Euclidean distance between the individual's activity center and the trap location,  $d_{ir} = \|\mathbf{x}_i - \mathbf{s}_r\|$ . To be precise about this, we let  $z_{irt}$  be the encounter frequency of individual  $i$  in trap  $r$  during occasion  $t$ . While we will adopt the



view that the variables  $z_{irt}$  are latent variables (see below), it will be convenient to formulate the model in terms of these variables.

Therefore, we assume that the expected encounter frequency of an individual in some trap is related to  $d_{ir}$  as follows:

$$E[z_{irt}] = \lambda_{ir} = \lambda_0 k_{ir}$$

where  $\lambda_0$  is the expected encounter rate at  $d = 0$  and  $k_{ir}$  is some positive-valued function of distance  $d_{ir}$ . We assume

$$k_{ir} = \exp(-d_{ir}^2/2\sigma^2)$$

where  $\sigma$  is a scale parameter related to home range size.  $\sigma$  also determines the degree of correlation among counts since animals with large home ranges are more likely to be detected at multiple traps relative animals with small home ranges. The phenomenon is analogous to correlation induced by averaging spatial noise, in which case there is a unique correlation between the smoothing kernel and the induced covariance function (Higdon, 2002).

We emphasize that our focus is on situations in which individuals are *not* uniquely identifiable, and therefore the encounter frequencies for each individual cannot be observed, and so they are latent variables. We assume that these latent variables are realizations from a Poisson distribution with mean  $\lambda_{ir}$ :

$$z_{irt} \sim \text{Poisson}(\lambda_{ir}). \quad (12.1)$$

In traditional SCR models,  $z_{irt}$  are the observed data, *i.e.*, the frequency of encounters of individual  $i$  at trap  $r$  on replicate survey  $t$ . However, when individual identity is not known, the observed data are the sample- and trap-specific totals, aggregated over all individuals:

$$n_{rt} = \sum_{i=1}^N z_{irt}.$$

Thus the data required by our model are a reduced-information summary of the latent encounter histories.

Under the Poisson encounter model we have that

$$n_{rt} \sim \text{Poisson}(\Lambda_r) \quad (12.2)$$

where

$$\Lambda_r = \lambda_0 \sum_i k_{ir}.$$

Further, because  $\Lambda_r$  does not depend on  $t$ , we can aggregate the replicated counts, defining  $n_{r\cdot} = \sum_t n_{rt}$  and then

$$n_{r\cdot} \sim \text{Poisson}(T\Lambda_r)$$

As such,  $T$  and  $\lambda_0$  serve equivalent roles as affecting baseline encounter rate. This formulation of the model in terms of the aggregate count simplifies computations as the latent variables  $z_{irt}$  do not need to be updated in the MCMC estimation scheme (see below). However, retaining  $z_{irt}$  in the formulation of the model is important if some individuals are uniquely marked, in which case modifying the MCMC algorithm (see below) to include both types of data is trivial. This is because uniquely identifiable individuals produce observations of some of the  $z_{irt}$  variables.

We imagine that other observation models might be possible (see Discussion) although we focus here on the Poisson encounter model because it has considerable relevance to animal surveys, and has additional methodological context related to point process models which we address in the Discussion.

## 12.3 Estimation by MCMC

We adopt a Bayesian framework for inference allowing estimation of  $N$  while retaining the formulation of the model that is conditional on the latent activity centers  $\mathbf{s}_i$ . Specifically, we employ Markov chain Monte Carlo (MCMC) to simulate posterior distributions of the parameters. However, the fact that  $N$  is unknown presents a technical challenge because the size of the parameter space can change with each MC iteration. To resolve this, we adopt the formulation of data augmentation in Royle et al. (2007) who used a specific prior construction for  $N$  in terms of individual level Bernoulli trials. In particular, we assume  $N \sim \text{Unif}(0, M)$  for some large integer  $M$ . We construct this prior by assuming  $N|M, \phi \sim \text{Bin}(M, \phi)$  and  $\phi \sim \text{DUnif}(0, 1)$  which implies, marginally, that  $N$  has the requisite  $\text{DUnif}(0, M)$  distribution. However the hierarchical formulation of the prior suggests an implementation in which we introduce a set of latent indicator variables  $w_i \sim \text{Bern}(\phi)$  and, furthermore, the model implies that  $z_{irt}$  are obtained from the specified distribution (Eq. 12.1) if  $w_i = 1$ , or if  $w_i = 0$ ,  $z_{irt} = 0$  with probability 1. In effect, extending the model in this way induces a reparameterization for the latent counts that is a zero-inflated version of the original conditional-on- $N$  model. Specifically, the model under data augmentation becomes

$$\begin{aligned} z_{irt}|w_i &\sim \text{Poisson}(\lambda_{ir}w_i) \\ w_i &\sim \text{Bern}(\phi) \end{aligned}$$

Under this formulation  $N = \sum_{i=1}^M w_i$ , and population density is simply  $D = N/A(\mathcal{S})$  where  $A(\mathcal{S})$  is the area of the point process state-space  $\mathcal{S}$ .

We developed two distinct MCMC implementations for this model (??). In the first, we devised an algorithm for the model conditional on the latent variables  $z_{irt}$ . This formulation is useful for problems in which one or more individual identities are available, in which case the  $z_{irt}$  are observable for those individuals. The unobserved  $z_{irt}$  are easily updated using their full-conditional distribution which is multinomial with sample size  $n_{rt}$ . The remaining parameters are updated using Metropolis-Hastings steps (see ??). In the second

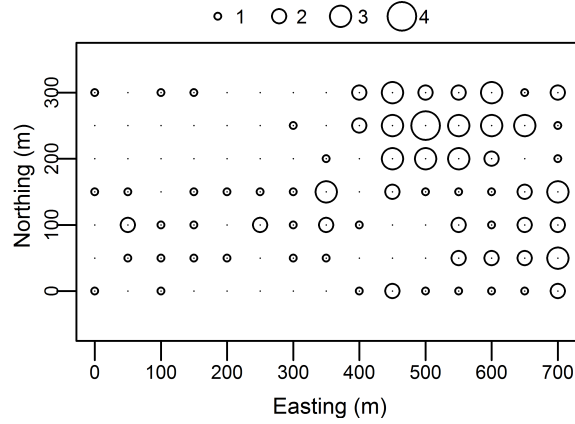


Figure 12.1: Spatially-correlated counts of northern parula on a 50-m grid. The size of the circle represents the total number of detections at each point.

163 formulation of the algorithm we applied the Metropolis-Hastings algorithm to  
 164 the model *unconditional* on the  $z_{irt}$  variables. In that case, the marginal distri-  
 165 bution for  $n_{rt}$  is precisely Eq. 12.2. This algorithm is slightly more convenient  
 166 because it avoids having to update the  $z_{irt}$  variables of which there are many.

## 167 12.4 Northern Parula Example

168 To apply our model to data collected in the field, we designed a point count  
 169 study of the northern parula (*Parula americana*), a Neotropical-Nearctic migra-  
 170 tory passerine. This species defends well-defined territories during the breeding  
 171 season (?), and thus our modeling effort was focused on estimating the number  
 172 and location of territory centers. Points were located on a 50-m grid to ensure  
 173 spatial correlation. This small grid spacing contrasts with the conventional prac-  
 174 tice of spacing points by  $> 200$  m to obtain *i.i.d.* counts. Figure 12.1 depicts  
 175 the spatially-correlated counts ( $n_r$ ) from the 105 point count locations surveyed  
 176 three times each during June 2006 at the Patuxent Wildlife Research Center in  
 177 Laurel Maryland, USA. A total of 226 detections were made with a maximum  
 178 count of 4 during a single survey. At 38 points, no warblers were detected. All  
 179 but one of the detections were of singing males, and this one observation was  
 180 not included in the analysis.

181 In our analysis of the parula data, we defined the point process state-space  
 182 by buffering the grid of point count locations by 250 m and used  $M = 300$ .  
 183 We simulated posterior distributions using three Markov chains, each consisting  
 184 of 300000 iterations after discarding the initial 10000 draws. Convergence was  
 185 satisfactory, as indicated by an  $\hat{R}$  statistic of  $< 1.02$  (Gelman and Rubin, 1992).

186 One benefit of a Bayesian analysis is that it can accommodate prior in-  
 187 formation on the home range size and encounter rate parameters, which are

Table 12.1: Posterior summary statistics for spatial Poisson-count model applied to the northern parula data. Two sets of priors were considered.  $M = 300$  was used in both cases. Parulas/ha,  $D$ , is a derived parameter.

Par	Prior	Mean	SD	Mode	q0.025	q0.50	q0.975
$\sigma$	$U(0, \infty)$	2.154	1.222	1.230	0.896	1.665	5.170
$\lambda_0$	$U(0, \infty)$	0.284	0.149	0.212	0.084	0.256	0.665
$N$	$U(0, M)$	40.953	38.072	4.000	3.000	31.000	143.000
$D$	–	0.427	0.397	0.0417	0.0313	0.323	1.490
$\sigma$	$G(13, 10)$	1.301	0.258	1.230	0.889	1.266	1.908
$\lambda_0$	$U(0, \infty)$	0.298	0.132	0.240	0.098	0.279	0.603
$N$	$U(0, M)$	59.321	36.489	36.000	18.000	50.000	157.000
$D$	–	0.618	0.380	0.375	0.188	0.521	1.635

readily available for many species. To illustrate, we analyzed the parula data using two sets of priors. In the first set, all priors were improper, customary non-informative priors (see Table 12.1). Uniform priors were also used in the second set, with the exception of an informative prior for the scale parameter  $\sigma \sim \text{Gamma}(13, 10)$ . We arrived at this prior using the methods described by Royle et al. (2011) and published information on the warbler’s home range size and detection probability (Simons et al., 2009). More details on this derivation are found in ???. We briefly note here that this prior includes the biologically-plausible range of values from  $\sigma$  suggested by the published literature.

The posterior distribution for  $N$  was highly skewed with a long right tail resulting in a wide 95% credible interval (Table 12.1). Nonetheless, the interval for density,  $D$ , includes estimates reported from more intensive field studies (?). This was true when considering both sets of priors, although posterior precision was higher under the informative set of priors. Specifically, the use of prior information reduced posterior density at high, biologically implausible, values of  $\sigma$ , and hence decreased the posterior mass for low values of  $N$  (Fig. 12.2).

In addition to estimating density, our model can be used to produce density surface maps, which are often used in applied ecological research to direct management efforts and develop hypotheses regarding the factors influencing abundance. Density surface maps can be produced by discretized the state-space and tallying the number of activity centers occurring in each pixel during each MCMC iteration. Parula density was highest near the northeastern corner of the study plot, which may correspond to important habitat features such as suitable nest site locations (Fig. 12.3). We anticipate future model extensions to directly model the point process intensity using habitat covariates.

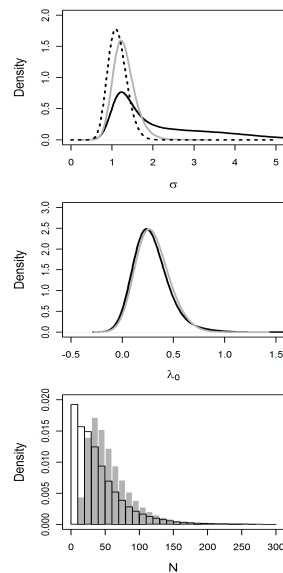


Figure 12.2: Effects of  $\sigma \sim \text{Gamma}(13, 10)$  prior on the posterior distributions from the northern parula model. Posteriors from model with uniform priors are shown in black, and posteriors from the informative prior model are shown in gray. The prior itself is shown as dotted line in the upper panel.

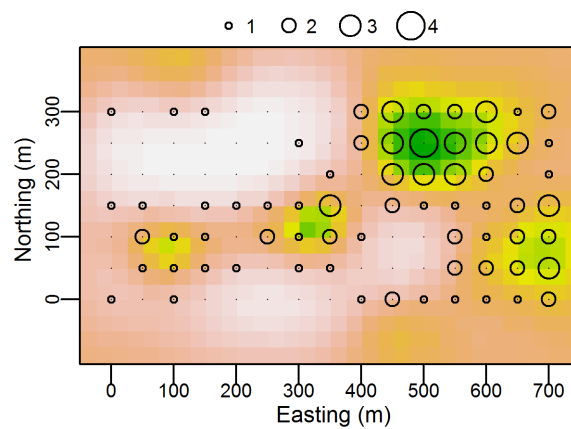


Figure 12.3: Estimated density surface of northern parula activity centers. The grid of point count locations with count totals is superimposed. See Fig. 1 for additional details.

## 213 12.5 On (Im)precision

## 214 12.6 How Much Correlation Is Enough?

## 215 12.7 Mutants

### 216 12.7.1 Other observation models

### 217 12.7.2 Linear designs

## 218 12.8 Summary

219 In this paper, we confronted one of the most difficult challenges faced in wildlife  
 220 sampling — estimation of density in the absence of data to distinguish among  
 221 individuals. To do so, we developed a novel class of spatially-explicit models that  
 222 applies to spatially organized counts, where the count locations or devices are  
 223 located sufficiently close together so that individuals are exposed to encounter  
 224 at multiple devices. This design yields correlation in the observed counts, and  
 225 this correlation proves to be informative about encounter probability paramet-  
 226 ers and hence density. We note that sample locations in count-based studies  
 227 are typically *not* organized close together in space because conventional wisdom  
 228 and standard practice dictate that independence of sample units is necessary  
 229 (Hurlbert, 1984). Our model suggests that in some cases it might be advan-  
 230 tageous to deviate from the conventional wisdom if one is interested in direct  
 231 inference about density. Of course, this is also known in the application of  
 232 standard spatial capture-recapture models (Borchers and Efford, 2008) where  
 233 individual identity is preserved across trap encounters, but it is seldom, if ever,  
 234 considered in the design of more traditional count surveys.

235 Our model has broad relevance to an incredible number of animal sampling  
 236 problems. Our motivating problem involved bird point counts where individual  
 237 identity is typically not available. The model also applies to other standard  
 238 methods used to sample unmarked populations, such as camera traps or even  
 239 methods that yield sign (*e.g.* scat, track) counts indexed by space. However,  
 240 results of our simulation study reveal some important limitations of the basic  
 241 estimator applied to situations in which none of the individuals can be uniquely  
 242 identified. In particular, posterior distributions are highly skewed in typical  
 243 small to moderate sample size situations and posterior precision is low.

244 Several modifications of the model can lead to improved performance of the  
 245 estimator. Our simulation results demonstrate that marking a subset of indi-  
 246 viduals can yield substantial increases in posterior precision. Marking a subset  
 247 of individuals is commonplace in animal studies such as when a small number of  
 248 individuals are radio-collared in conjunction with a count-based survey (Bart-  
 249 mann et al., 1987). In many other situations a subset of individuals can be  
 250 identified by natural marks alone, and thus our model could be applied to data  
 251 from camera-trapping studies of species such as mountain lions, deer, coyotes for

which traditional SCR methods are not effective (Kelly et al., 2008). Thus, the ability to study partially-marked populations adds flexibility to existing SCR methods, and also creates new opportunities for designing efficient SCR studies since the costs of marking all individuals in a population can be prohibitive.

We note the existence of traditional approaches to combining data on marked and unmarked animals based on either the Lincoln-Peterson estimator or so-called “mark-resight” methods. (Bartmann et al., 1987; ?; ?). In their simplest form, mark-resight methods involve fitting standard closed-population mark-recapture models to the data on marked individuals, and the resultant estimate of detection probability ( $\hat{p}$ ) is used to estimate population size as  $\hat{N} = m + u/\hat{p}$  where  $m$  and  $u$  are the number of marked and unmarked individual, respectively. In this case, the unmarked individuals provide no information about the encounter rate parameters, and thus mark-resight methods cannot be used unless a large sample of marked individuals is available. This contrasts with our approach which can be used even when all individuals are unmarked.

In some cases, such as in point counts of birds, it may not be practical to mark individuals. An alternative to increasing posterior precision is to utilize prior information on home range size. Indeed, extensive information on home range size has been compiled for many species in diverse habitats (*e.g.*, DeGraaf and Yamasaki, 2001). It is easy to embody this information in a prior distribution as we demonstrated for the parula data.

An additional design extension that could increase precision is to use multiple sampling methods, in which one method generates encounter frequencies and the other method generates individuality. For example, camera traps are now commonly used with surveys for sign (scat or tracks), or hair snares for sampling bear populations. These distinct methods would have different basal detection rates but share an underlying spatial model describing the organization of individuals in space. Our models show promise for using these disparate data types efficiently for estimating density.

### 12.8.1 $N$ -mixture models

Parallel developments which appear ostensibly orthogonal to SCR models have addressed the problem of estimating population size when individuals are unmarked. So-called  $N$ -mixture models (Royle, 2004a,b; Royle et al., 2004) can be applied to a repeated-measures type of data structure wherein data are collected at  $R$  sites, with  $J$  replicate surveys are conducted at each.  $N$ -mixture models regard abundance at each site ( $N_r$ ) as an *i.i.d.* realization of a discrete distribution such as the Poisson or negative binomial with expectation  $\theta$ . In the standard binomial  $N$ -mixture model, the observed counts are treated as binomial outcomes with  $N_r$  “trials” and detection probability  $p$ .

Although these models have proven useful for studies of factors that affect variation in abundance, interpretation of model parameters is strongly dependent on the assumption that populations are closed with respect to demographic processes and movement. The closure assumption can be an important practical limitation (but see Dail and Madsen, 1988; Chandler et al., 2011). Furthermore

the *i.i.d.* assumption is violated if spatial correlation exists among sites, such as if animals move among plots. Although we formulated the model developed in our paper as an extension of spatially explicit capture-recapture models, it clearly can also be viewed as a spatially explicit extension of  $N$ -mixture models where the local population sizes  $N_r$  are dependent owing to the nature of the sampling design.

Thus, two recently developed methodological frameworks, spatial capture-recapture and  $N$ -mixture models, address different problems that arise in sampling animal populations. SCR models address non-closure by accommodating information on the spatial organization of individuals and juxtaposition of individuals with traps, and  $N$ -mixture models address inability to uniquely identify individuals. Our model unifies these two modeling frameworks by addressing both issues simultaneously.

### 12.8.2 Alternative Observation Models

Several aspects of our “spatial  $N$ -mixture model” can be modified to accommodate alternative sampling designs or parametric distributions. We considered situations where an individual can be detected more than once at a trap during a single occasion, but under some designs this is not possible. When collecting DNA samples, for instance, an individual can often be detected at most once during an occasion, because multiple samples of biological material cannot be attributed to distinct episodes. Therefore, rather than  $z_{irt} \sim \text{Poisson}(\lambda_{ir})$  we have  $z_{irt} \sim \text{Bernoulli}(p_{ir})$  where, for example,  $p_{ir} = p_0 \exp(-d_{ir}^2/(2\sigma^2))$ , and  $p_0$  is the probability of detecting an individual whose home range is centered on trap  $r$ . This Bernoulli model is a focus of ongoing investigations.

Both the Poisson and the Bernoulli models produce count observations when aggregated over individuals to form trap-specific totals; however, ecologists often collect so-called “detection/non-detection” data because it can be easier to determine if “at least one” individual was present rather than enumerating all individuals in a location. In this case, the underlying  $z_{irt}$  array is the same as the above cases, but we observe  $y_{rt} = I(\sum_{i=1}^N z_{irt} > 0)$  where  $I$  is the indicator function. This “Poisson-binary model” is a spatially explicit extension of the model of Royle and Nichols (2003) in which the underlying abundance state is inferred from binary data. We have investigated this model to a limited extent but do not report on those results here.

### 12.8.3 Spatial point process models

Our model has some direct linkages to existing point process models. We note that the observation intensity function (i.e., corresponding to the observation locations) is a compound Gaussian kernel similar to that of the Thomas process (??, pp. 61-62). Also, the Poisson-Gamma Convolution models (Wolpert and Ickstadt, 1998) are structurally similar (see also Higdon (1998) and Best et al. (2000)). In particular, our model is such a model but with a *constant* basal encounter rate  $\lambda_0$  and *unknown* number and location of “support points”, which



in our case are the animal activity centers,  $\mathbf{s}_i$ . We can thus regard our model as a model for *estimating* the location and local density of support points in such models, which we believe could be useful in the application of convolution models. Best et al. (2000) devise an MCMC algorithm for the Poisson-Gamma model based on data augmentation, which is similar to the component of our algorithm for updating the  $z$  variables in the conditional-on- $z$  formulation of the model. We emphasize that our model is distinct from these Poisson-Gamma models in that the number *and* location of such support points are estimated.

If individuals were perfectly observable then the resulting point process of locations is clearly a standard Poisson or Binomial (fixed  $N$ ) cluster process or Neyman-Scott process. If detection is uniform over space but imperfect, then the basic process is unaffected by this random thinning. Our model can therefore be viewed formally as a Poisson (or Binomial) cluster process model but one in which the thinning is non-uniform, governed by the encounter model which dictates that thinning rate increases with distance from the observation points. In addition, our inference objective is, essentially, to estimate the number of parents in the underlying Poisson cluster process, where the observations are biased by an incomplete sampling apparatus (points in space).

As a model of a thinned point process, our model has much in common with classical distance sampling models (Buckland, 2001). The main distinction is that our data structure does *not* include observed distances, although the underlying observation model is fundamentally the same as in distance sampling if there is only a single replicate sample and  $\mathbf{s}_i$  is defined as an individual's location at an instant in time. For replicate samples, our model preserves (latent) individuality across samples and traps which is not a feature of distance sampling. We note that error in measurement of distance is not a relevant consideration in our model, and we explicitly do not require the standard distance sampling assumption that the probability of detection is 1 if an individual occurs at the survey point. More importantly, distance sampling models cannot be applied to data from many of the sampling designs for which our model is relevant. For example, many rare and endangered species can only be effectively surveyed using methods such as hair snares and camera traps that do not produce distance data (O'Connell et al., 2010).

## 12.9 Conclusion

Concerns about "statistical independence" have prompted ecologists to design count-based studies such that observed random variables can be regarded as *i.i.d.* outcomes (Hurlbert, 1984). Interestingly, this often proves impossible in practice, and elaborate methods have been devised to model spatial dependence as a nuisance parameter. Our paper presents a modeling framework that directly confronts this view by demonstrating that spatial correlation carries information about the locations of individuals, which can be used to estimate density even when individuals are unmarked and distance-related heterogeneity exists in encounter probability.



# Bibliography

- Bartmann, R. M., White, G. C., Carpenter, L. H., and Garrott, R. A. (1987),  
“Aerial Mark-Recapture Estimates of Confined Mule Deer in Pinyon-Juniper  
Woodland,” *The Journal of Wildlife Management*, 51, 41–46.
- Best, N. G., Ickstadt, K., and Wolpert, R. L. (2000), “Spatial Poisson Regression  
for Health and Exposure Data Measured at Disparate Resolutions,” *Journal  
of the American Statistical Association*, 95, 1076.
- Bibby, C. and Buckland, S. (1987), “Bias of bird census results due to detectabil-  
ity varying with habitat,” *Acta Ecologica*, 8, 103–112.
- Borchers, D. and Efford, M. (2008), “Spatially explicit maximum likelihood  
methods for capture–recapture studies,” *Biometrics*, 64, 377–385.
- Buckland, S. T. (2001), *Introduction to distance sampling: estimating abundance  
of biological populations*, Oxford, UK: Oxford University Press.
- Chandler, R. and Royle, J. (2012), “Spatially-explicit models for inference about  
density in unmarked populations,” *Biometrics (in review)*.
- Chandler, R., Royle, J., and King, D. (2011), “Inference about density and  
temporary emigration in unmarked populations,” *Ecology*, 92, 1429–1435.
- Dail, D. and Madsen, L. (????), “Web Supplementary Materials Models for  
Estimating Population Size from Repeated Counts of an Open Population,”  
1, 1–7.
- DeGraaf, R. M. and Yamasaki, M. (2001), *New England wildlife: habitat, nat-  
ural history, and distribution*, University Press of New England.
- Gelman, A. and Rubin, D. B. (1992), “Inference from iterative simulation using  
multiple sequences,” *Statistical Science*, 7, 457–511.
- Higdon, D. (1998), “A process-convolution approach to modelling temperatures  
in the North Atlantic Ocean,” *Environmental and Ecological Statistics*, 5,  
173–190.

- (2002), “Space and Space-Time Modeling using Process Convolutions,” in *Quantitative methods for current environmental issues*, eds. Anderson, C., Barnett, V., Chatwin, P., and El-Shaarawi, A., Springer Verlag, p. 37.
- Hurlbert, S. (1984), “Pseudoreplication and the design of ecological field experiments,” *Ecological monographs*, 54, 187–211.
- Johnson (2010), “A Model-Based Approach for Making Ecological Inference from Distance Sampling Data,” *Biometrics*, 66, 310318.
- Kelly, M., Noss, A., Di Bitetti, M., Maffei, L., Arispe, R., Paviolo, A., De Angelo, C., and Di Blanco, Y. (2008), “Estimating puma densities from camera trapping across three study sites: Bolivia, Argentina, and Belize,” *Journal of Mammalogy*, 89, 408418.
- O’Connell, A. F., Nichols, J. D., and Karanth, U. K. (2010), *Camera traps in animal ecology: Methods and analyses*, Springer.
- Royle, J., Dorazio, R., and Link, W. (2007), “Analysis of multinomial models with unknown index using data augmentation,” *Journal of Computational and Graphical Statistics*, 16, 67–85.
- Royle, J. and Nichols, J. (2003), “Estimating abundance from repeated presence-absence data or point counts,” *Ecology*, 84, 777–790.
- Royle, J. A. (2004a), “Generalized estimators of avian abundance from count survey data,” *Animal Biodiversity and Conservation*, 27, 375–386.
- (2004b), “N-mixture models for estimating population size from spatially replicated counts,” *Biometrics*, 60, 108–115.
- Royle, J. A., Dawson, D. K., and Bates, S. (2004), “Modeling abundance effects in distance sampling,” *Ecology*, 85, 1591–1597.
- Royle, J. A., Kéry, M., and Guélat, J. (2011), “Spatial capture-recapture models for search-encounter data,” *Methods in Ecology and Evolution*, 1–10.
- Simons, T. R., Pollock, K. H., Wettroth, J. M., Alldredge, M. W., Pacifici, K., and Brewster, J. (2009), “Sources of Measurement Error, Misclassification Error, and Bias in Auditory Avian Point Count Data,” in *Modeling Demographic Processes In Marked Populations*, eds. Thomson, D. L., Cooch, E. G., and Conroy, M. J., Boston, MA: Springer US, vol. 3, pp. 237–254.
- Wolpert, R. L. and Ickstadt, K. (1998), “Poisson/gamma random field models for spatial statistics,” *Biometrika*, 85, 251–267.