# Spatial Capture-Recapture Models

A hierarchical approach

The Four Horsemen (and women)

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## 

## INTRODUCTION TO BAYESIAN ANALYSIS OF GL(M)MS USING R/WINBUGS

A major theme of this book is that spatial capture-recapture models are, for the most part, just generalized linear models (GLMs) wherein the covariate, distance between trap and home range center, is partially or fully unobserved – and therefore regarded as a random effect. Such models are usually referred to as Generalized Linear Mixed Models (GLMMs) and, therefore, SCR models can be thought of as a specialized type of GLMM. Naturally then, we should consider analysis of these slightly simpler models in order to gain some experience and, hopefully, develop a better understanding of spatial capture-recapture models.

In this chapter, we consider classes of GLM models - Poisson and binomial (i.e., logistic regression) GLMs - that will prove to be enormously useful in the analysis of capture-recapture models of all kinds. Many readers are probably familiar with these models because they represent probably the most generally useful models in all of Ecology and, as such, have received considerable attention in many introductory and advanced texts. We focus on them here in order to introduce the readers to the analysis of such models in **R** and **WinBUGS**, which we will translate directly to the analysis of SCR models in subsequent chapters.

Bayesian analysis is convenient for analyzing GLMMs because it allows us to work directly with the conditional model – i.e., the model that is conditional on the random effects, using computational methods known as Markov chain Monte Carlo (MCMC). Learning how to do Bayesian analysis of GLMs and GLMMs in **WinBUGS** is, in part, the purpose of this chapter. While we use **WinBUGS** to do the Bayesian computations, we organize and summarize our data and execute

WinBUGS from within R using the useful package R2WinBUGS (Sturtz et al., 2005). Kéry (2010), and Kery and Schaub (2011) provide excellent introductions to the basics of Bayesian analysis and GLMs at an accessible level. We don't want to be too redundant with those books and so we avoid a detailed treatement of Bayesian methodology - instead just providing a cursory overview so that we can move on and attack the problems we're most interested in related to spatial capture-recapture. In addition, there are a number of texts that provide general introductions to Bayesian analysis, MCMC, and their applications in Ecology including McCarthy (2007), Kéry (2010), Link and Barker (2009), and King (2009).

While this chapter is about Bayesian analysis of GLMMs, such models are routinely analyzed using likelihood methods too, as discussed by Royle and Dorazio (2008), and Kéry (2010). Indeed, likelihood analysis of such models is the primary focus of many applied statistics texts, a good one being Zuur et al. (2009). Later in this book, we will use likelihood methods to analyze SCR models but, for now, we concentrate on providing a basic introduction to Bayesian analysis because that is the approach we will use in a majority of cases in later chapters.

## 2.1 NOTATION

We will sometimes use conventional "bracket notation" to refer to probability distributions. If y is a random variable the [y] indicates its distribution or its probability density/mass function (pdf, pmf) depending on context. If x is another random variable then [y|x] is the conditional distribution of y given x, and [y,x] is the joint distribution of y and x. To differentiate specific distributions in some contexts we might label them g(y),  $g(y|\theta)$ , f(x), or similar. We will also write  $y \sim \text{Normal}(\mu, \sigma^2)$  to indicate that y "is distributed as" a normal random variable with parameters  $\mu$  and  $\sigma^2$ . The expected value or mean of a random variable is  $E[y] = \mu$ , and  $Var[y] = \sigma^2$  is the variance of y. To indicate specific observations we'll use an index such as "i". So,  $y_i$  for  $i = 1, 2, \ldots, n$  indicates observations for n individuals. Finally, we write Pr(y) to indicate specific probabilities, i.e., of events "y" or similar.

To illustrate these concepts and notation, suppose z is a binary outcome (e.g., species occurrence) and we might assume the model:  $z \sim \operatorname{Bern}(p)$  for observations. Under this model  $\Pr(z=1)=\psi$ , which is also the expected value  $E[z]=\psi$ . The variance is  $Var[z]=\psi*(1-\psi)$  and the probability mass function (pmf) is  $[z]=\psi^z(1-\psi)^{1-z}$ . Sometimes we write  $[z|\psi]$  when it is important to emphasize the conditional dependence of z on  $\psi$ . As another example, suppose y is a random variable denoting whether or not a species is detected if an occupied site is surveyed. In this case it might be natural to express the pmf of the observations y conditional on z. That is, [y|z]. In this case, [y|z=1] is the conditional pmf of y given that a site is occupied, and it is natural to assume that  $[y|z=1]=\operatorname{Bern}(p)$  where p is the "detection probability" - the probability that we detect the species, given that it is present. The model for the observations y is completely specified once we describe

the other conditional pmf [y|z=0]. For this conditional distribution it is sometimes reasonable to assume  $\Pr(y=1|z=0)=0$  (MacKenzie et al. (2002); see also Royle and Link (2006)). That is, if the species is absent, the probability of detection is 0. This implies that  $\Pr(y=0|z=0)=1$ . To allow for situations in which the true state z is unobserved, we assume that [z] is Bernoulli with parameter  $\psi$ . In this case, the marginal distribution of y is

$$[y] = [y|z=1]Pr(z=1) + [y|z=0]Pr(z=0)$$

because [y|z=0] is a point mass at y=0, by assumption, then

$$Pr(y=1) = p\psi$$

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$$\Pr(y = 0) = (1 - p) * \psi + (1 - \psi)$$

## 2.2 GLMS AND GLMMS

We have asserted already that SCR models work out most of the time to be variations of GLMs and GLMMs. Some of you might therefore ask: What are GLMs and GLMMs, anyhow? These models are covered extensively in many very good applied statistics books and we refer the reader elsewhere for a detailed introduction. We think Kéry (2010), Kery and Schaub (2011), and Zuur et al. (2009) are all accessible treatments of considerable merit. Here, we'll give the 1 minute treatment of GLMMs, not trying to be complete but rather only to preserve a coherent organization to the book.

The generalized linear model (GLM) is an extension of standard linear models by allowing the response variable to have some distribution from the exponential family of distributions (i.e., not just normal). This includes the normal distribution but also dozens of others such as the Poisson, binomial, gamma, exponential, and many more. In addition, GLMS allow the response variable to be related to the predictor variables (i.e., covariates) using a link function, which is usually nonlinear. Finally, GLMs typically accommodate a relationship between the mean and variance. The classical reference for GLMs is Nelder and Wedderburn (1972) and also McCullagh and Nelder (1989). The GLM consists of three components:

- 1. A probability distribution for the dependent variable y, from a class of probability distributions known as the exponential family.
- 2. A "linear predictor"  $\eta = \mathbf{X}\beta$ .
- 3. A link function g that relates E[y] to the linear predictor,  $E[y] = \mu = g^{-1}(\eta)$ . Therefore  $g(E[y]) = \eta$ .

The dependent variable y is assumed to be an outcome from a distribution of the exponential family which includes many common distributions including the

normal, gamma, Poisson, binomial, and many others. The mean of the distribution of y is assumed to depend on predictor variables x according to

$$g(E[y]) = \mathbf{x}'\beta$$

where E[y] is the expected value of y, and  $\mathbf{x}'\beta$  is termed the linear predictor, i.e., a linear function of the predictor variables with unknown parameters  $\beta$  to be estimated. The function g is the link function. In standard GLMs, the variance of y is a function V of the mean of y:  $Var(y) = V(\mu)$  (see below for examples).

A Poisson GLM posits that  $y \sim \text{Poisson}(\lambda)$  with  $E[y] = \lambda$  and usually the model for the mean is specified using the log link function by

$$log(\lambda_i) = \beta_0 + \beta_1 * x_i$$

The variance function is  $V(y_i) = \lambda_i$ . The binomial GLM posits that  $y_i \sim \text{Binomial}(K, p)$ where K is the fixed sample size parameter and  $E[y_i] = K * p_i$ . Usually the model for the mean is specified using the *logit link function* according to

$$logit(p_i) = \beta_0 + \beta_1 * x_i$$

Where logit(u) = log(u/(1-u)). The inverse-logit function,  $g^{-1}$ , is a function we will refer to as "expit", so that expit(u) = exp(u)/(1 + exp(u)).

A GLMM is the extension of GLMs to accommodate "random effects". Often this involves adding a normal random effect to the linear predictor, and so a simple example is:

$$\log(\lambda_i) = \alpha_i + \beta_1 * x_i$$

where

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$$\alpha_i \sim \text{Normal}(\mu, \sigma^2)$$

## 2.3 BAYESIAN ANALYSIS

Bayesian analysis is unfamiliar to many ecological researchers because older cohorts of ecologists were largely educated in the classical statistical paradigm of frequentist inference. But advances in technology and increasing exposure to benefits of Bayesian analysis are fast making Bayesians out of people or at least making Bayesian analysis an acceptable, general, alternative to classical, frequentist inference.

Conceptually, the main thing about Bayesian inference is that it uses probability directly to characterize uncertainty about things we don't know. "Things", in this case, are parameters of models and, just as it is natural to characterize uncertain outcomes of stochastic processes using probability, it seems natural also to characterize information about unknown "parameters" using probability. At least this seems natural to us and, we think, most ecologists either explicitly adopt that view or tend to fall into that point of view naturally. Conversely, frequentists use

probability in many different ways, but never to characterize uncertainty about parameters<sup>1</sup> Instead, frequentists use probability to characterize the behavior of procedures such as estimators or confidence intervals (see below), which can lead to some inelegant or unnatural interpretations of things. It is paradoxical that people readily adopt a philosophy of statistical inference in which the things you don't know (i.e., parameters) should not be regarded as random variables, so that, as a consequence, one cannot use probability to characterize ones state of knowledge about them.

### 2.3.1 Bayes Rule

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As its name suggests, Bayesian analysis makes use of Bayes' rule in order to make direct probability statements about model parameters. Given two random variables z and y, Bayes rule relates the two conditional probability distributions [z|y] and [y|z] by the relationship:

$$[z|y] = [y|z][z]/[y]$$

Bayes' rule itself is a mathematical fact and there is no debate in the statistical community as to its validity and relevance to many problems. Generally speaking, these distributions are characterized as follows: [y|z] is the conditional probability distribution of y given z, [z] is the marginal distribution of z and [y] is the marginal distribution of y. In the context of Bayesian inference we usually associate specific meanings in which [y|z] is thought of as "the likelihood", [z] as the "prior" and so on. We leave this for later because here the focus is on this expression of Bayes rule as a basic fact of probability.

As an example of a simple application of Bayes rule, consider the problem of determining species presence at a sample location based on imperfect survey information. Let z be a binary random variable that denotes species presence (z=1) or absence (z=0), let  $\Pr(z=1)=\psi$  where  $\psi$  is usually called occurrence probability, "occupancy" (MacKenzie et al., 2002) or "prevalence". Let y be the observed presence (y = 1) or absence (y = 0), and let p be the probability that a species is detected in a single survey at a site given that it is present. Thus,  $\Pr(y=1|z=1)=p$ . The interpretation of this is that, if the species is present, we will only observe presence with probability p. In addition, we assume here that  $\Pr(y=1|z=0)=0$ . That is, the species cannot be detected if it is not present which is a conventional view adopted in most biological sampling problems (but see Royle and Link (2006)). If we survey a site T times but never detect the species, then this clearly does not imply that the species is not present (z=0) at this site. Rather, our degree of belief in z=0 should be made with a probabilistic statement  $Pr(z=1|y_1=0,\ldots,y_T=0)$ . If the T surveys are independent so that we might regard  $y_t$  as *iid* Bernoulli trials, then the total number of detections, say y, is Binomial with probability p then we can use Bayes rule to compute the

<sup>&</sup>lt;sup>1</sup>To hear this will be shocking to some readers perhaps.

probability that it is present given that it is not detected in T samples. In words, the expression we seek is:

$$\Pr(\text{present}|\text{not detected}) = \frac{\Pr(\text{not detected}|\text{present}) \Pr(\text{present})}{\Pr(\text{detected})}$$

Mathematically, this is

$$Pr(z = 1|y = 0) = Pr(y = 0|z = 1) Pr(z = 1) / Pr(y = 0)$$
$$= [(1 - p)^{T} \psi] / [(1 - p)^{T} \psi + (1 - \psi)].$$

To apply this, suppose that T=2 surveys are done at a wetland for a species of frog, and the species is not detected there. Suppose further that  $\psi=.8$  and p=.5 are obtained from a prior study. Then the probability that the species is present at this site is .25\*.8/(.25\*.8+.2)=0.50. That is, there seems to be about a 50/50 chance that the site is occupied despite the fact that the species wasn't observed there.

In summary, Bayes' rule provides a simple linkage between the conditional probabilities [y|z] and [z|y] which is useful whenever one needs to deduce one from the other. Bayes' rule as a basic fact of probability is not disputed.

## 2.3.2 Bayesian Inference

What is controversial to some is the scope and manner in which Bayes rule is applied by Bayesian analysts. Bayesian analysts assert that Bayes rule is relevant, in general, to all statistical problems by regarding all unknown quantities of a model as realizations of random variables - this includes "data", latent variables, and also "parameters". Classical (non-Bayesian) analysts sometimes object to regarding "parameters" as outcomes of random variables. Classically, parameters are thought of as "fixed but unknown" (using the terminology of classical statistics). Of course, in Bayesian analysis they are also unknown and, in fact, there is a single data-generating value and so they are also fixed. The difference is that this fixed but unknown value is regarded as having been generated from some probability distribution. Specification of that probability distribution is necessary to carryout Bayesian analysis, but it is not required in classical frequentist inference.

To see the general relevance of Bayes rule in the context of statistical inference, let y denote observations - i.e., "data" - and let  $[y|\theta]$  be the observation model (often colloquially referred to as the "likelihood"). Suppose theta is a parameter of interest having (prior) probability distribution  $[\theta]$ . These are combined to obtain the posterior distribution using Bayes' rule, which is:

$$[\theta|y] = [y|\theta][\theta]/[y]$$

Asserting the general relevance of Bayes rule to all statistical problems, we can conclude that the two main features of Bayesian inference are that: (1) "parameters"

 $\theta$  are regarded as realizations of a random variable and, as a result, (2) inference is based on the probability distribution of the parameters given the data,  $[\theta|y]$ , which is called the posterior distribution. This is the result of using Bayes rule to combine "the likelihood" and the prior distribution. The key concept is regarding parameters as realizations of a random variable because, once you admit this conceptual view, this leads directly to the posterior distribution, a very natural quantity upon which to base inference about things we don't know - including parameters of statistical models. In particular,  $[\theta|y]$  is a probability distribution for  $\theta$  and therefore we can make direct probability statements to characterize uncertainty about  $\theta$ .

The denominator of our invocation of Bayes rule, [y], is the marginal distribution of the data y. We note without further remark right now that, in many practical problems, this can be an enormous pain to compute. The main reason that the Bayesian paradigm has become so popular in the last 20 years or so is because methods exist for characterizing the posterior distribution that do not require that we possess a mathematical understanding of [y], i.e., we never have to compute it or know what it looks like, or know anything specific about it.

A common misunderstanding on the distinction between Bayesian and frequentist inference goes something like this "in frequentist inference parameters are fixed but unknown but in a Bayesian analysis parameters are random." At best this is a sad caricature of the distinction and at worst it is downright wrong. What is true is that, to a Bayesian, parameters are random variables. However, a Bayesian assumes, just like a frequentist, that there was a single data-generating value of that parameter - a fixed, and unknown value that produced the given data set. The distinction between Bayesian and frequentist approaches is that Bayesians regard the parameter as a random variable, and its value as the outcome of a random value, on par with the observations. This allows Bayesians to use probability to make direct probability statements about parameters. Frequentist inference procedures do not permit direct probability statements to be made about parameter values – because parameters are not random variables!

While we can understand the conceptual basis of Bayesian inference merely by understanding Bayes rule – that's really all there is to it – it is not so easy to understand the basis of classical "frequentist" inference which is mostly like<sup>2</sup> a "basket of methods" with little coherent organization. What is mostly coherent in frequentist inference is the manner in which items in this basket of methods are evaluated – the performance of a given procedure is evaluated by "averaging over" hypothetical realizations of y, regarding the *estimator* as a random variable. For example, if  $\hat{\theta}$  is an estimator of  $\theta$  then the frequentist is interested in  $E_y[\hat{\theta}|y]$  which is used to characterize bias. If the expected value of  $\hat{\theta}$ , when averaged over realizations of y, is equal to  $\theta$ , then  $\hat{\theta}$  is unbiased.

The view of parameters as fixed constants and estimators as random variables leads to interpretations that are not so straightforward. For example confidence

<sup>&</sup>lt;sup>2</sup>Characterization from Sims REF XYZ

intervals having the interpretation "95% probability that the interval contains the true value" and p-values being "the probability of observing an outcome as extreme or more than the one observed." These are far from intuitive interpretations to most people. Moreover, this is conceptually probblematic to some because the hypothetical realizations that characterize the performance of our procedure we will never get to observe.

While we do tend to favor Bayesian inference for the conceptual simplicity (parameters are random, posterior inference), we mostly advocate for a pragamatic non-partisian approach to inference because, frankly, some of these "bucket of methods" are actually very convenient in certain situations as we will see in later chapters.

#### 2.3.3 Prior distributions

The prior distribution  $[\theta]$  is an important feature of Bayesian inference. As a conceptual matter, the prior distribution characterizes "prior beliefs" or "prior information" about a parameter. Indeed, an oft-touted benefit of Bayesian analysis is the ease with which prior information can be included in an analysis. However, more commonly, the prior is chosen to express a lack of prior information, even if previous studies have been done and even if the investigator does in fact know quite a bit about a parameter. This is because the manner in which prior information is embodied in a prior (and the amount of information) is usually very subjective and thus the result can wind up being very contentious, e.g., if different investigators might report different results based on subjective assessments of things. Thus it is usually better to "let the data speak" and use priors that reflect absence of information beyond the data set being analyzed.

But still the need occasionally arises to embody prior information or beliefs about a parameter formally into the estimation scheme. In SCR models we often have a parameter that is closely linked to "home range radius" and thus auxiliary information on the home range size of a species can be used as prior information (e.g., see Chandler and Royle (2012); also chapter XYZ).

### XXXXXXXX

noninformative prior on one scale is informative on another scale. e.g., flat prior on logit(p) is very different from uniform(0,1) on p... show graphic.....

reference to non-invariance of prior distributions to transformation.....

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### 2.3.4 Posterior Inference

In Bayesian inference, we are not focusing on estimating a single point or interval but rather on characterizing a whole distribution – the posterior distribution – from which one can report any summary of interest. A point estimate might be the posterior mean, median, mode, etc.. In many applications in this book, we will compute 95% Bayesian intervals using the 2.5% and 97.5% quantiles of the posterior distribution. For such intervals, it is correct to say  $\Pr(L < \theta < U) = 0.95$ . That is, "the probability that  $\theta$  is between L and U is 0.95". It is not a subtle thing that this cannot be said using frequentist methods - although people tend to say it anyway and not really understand why it is wrong or even that it is wrong. This is actually a failing of frequentist ideas and the inability of frequentists to get people to overcome their natural tendency to use probability - which is something that, as a frequentist, you simply cannot do in the manner that you would like to.

Posterior inference is the main practical element of Bayesian analysis. We get to make an inference conditional on the data that we actually observed - i.e., what we actually know. To us, this seems logical - to condition on what we know. Conversely, frequentist inference is based on considering average performance over hypothetical unobserved data sets (i.e., the "relative frequency" interpretation of probability). Frequentists know that their procedures work well when averaged over all hypothetical, unobserved, data sets but no one ever really knows how well they work for the specific data set analyzed. That seems like a relevant question to biologists who oftentimes only have their one, extremely valuable, data set. This distinction comes into play a lot in exposing philosophical biases in the peer review of statistical analyses in ecology in the sense that, despite these opposing conceptual views to inference (i.e. conditional on the data you have, or averaged over hypothetical realizations), those who conduct a Bayesian analysis are often (in ecology, almost always) required to provide a frequentist evaluation of their Bayesian procedure.

## 2.3.5 Small sample inference

Using Bayesian inference, we obtain an estimate of the posterior distribution which is an exhaustive summary of the state-of-knowledge about an unknown quantity. It is the posterior distribution - not an estimate of that thing. It is also not, usually, an approximation except to within Monte Carlo error (in cases where we use simulation to calculate it). One of the great virtues of Bayesian analysis which is not really appreciated is that it is completely valid for any particular sample size. i.e., it is  $[\theta|y]$ , as precise as we claim it to be based on our ability to do calculations, for the particular sample size and observations that we have even if we have only a single datum y. The same cannot be said for almost all frequentist procedures in which estimates or variances are very often (almost always in practice) based on "asymptotic approximations" to the procedure which is actually being employed.

There seems to be a prevailing view in statistical ecology that classical likelihoodbased procedures are virtuous because of the availability of simple formulas and procedures for carrying out inference, such as calculating standard errors, doing model selection by AIC, and assessing goodness-of-fit. In large samples, this may be an important practical benefit, but the theoretical validity of these procedures cannot be asserted in most situations involving small samples. This is not a minor

issue because it is typical in many wildlife sampling problems - especially in surveys of carnivores or rare/endangered species - to wind up with a small, sometimes extremely small, data set. For example, a recent paper on the fossa (Cryptoprocta ferox), an endangered carnivore in Madagascar, estimated an adult density of 0.18 adults / km sq based on 20 animals captured over 3 years (Hawkins and Racey, 2005). A similar paper on the endangered southern river otter (Lontra provocax) estimated a density of 0.25 animals per river km based on 12 individuals captured over 3 years (Sepúlveda et al., 2007). Gardner et al. (2010) analyzed data from a study of the Pampas cat, a species for which very little is known, wherein only 22 individual cats were captured .during the two year period. Trolle and Kéry (2005) reported only 9 individual ocelots captured and Jackson et al. (2006) captured 6 individual snow leopards using camera trapping. Thus, studies of rare and/or secretive carnivores necessarily and flagrantly violate one of Le Cam's Basic Principles, that of "If you need to use asymptotic arguments, do not forget to let your number of observations tend to infinity." (Le Cam, 1990).

The biologist thus faces a dilemma with such data. On one hand, these datasets, and the resulting inference, are often criticized as being poor and unreliable. Or, even worse<sup>3</sup>, "the data set is so small, this is a poor analysis." On the other hand, such data may be all that is available for species that are extraordinarily important for conservation and management. The Bayesian framework for inference provides a valid, rigorous, and flexible framework that is theoretically justifiable in arbitrary sample sizes. This is not to say that one will obtain precise estimates of density or other parameters, just that your inference is coherent and justifiable from a conceptual and technical statistical point of view. That is, we report the posterior probability  $\Pr(D|data)$  which is easily interpretable and just what it is advertised to be and we don't need to do a simulation study to evaluate how well some approximate  $\Pr(D|data)$  deviates from the actual  $\Pr(D|data)$  because they are precisely the same quantity.

## 2.4 CHARACTERIZING POSTERIOR DISTRIBUTIONS BY MCMC SIMULATION

In practice, it is not really feasible to ever compute the marginal probability distribution Pr(y), the denominator resulting from application of Bayes' rule. For decades this impeded the adoption of Bayesian methods by practitioners. Or, the few Bayesian analyses done were based on asymptotic normal approximations to the posterior distribution. While this was useful stuff from a theoretical and technical standpoint and, practically, it allowed people to make the probability statements that they naturally would like to make, it was kind of a bad joke around the Bayesian water-cooler to, on one hand, criticize classical statistics for being, essentially, completely ad hoc in their approach to things but then, on the other hand,

<sup>&</sup>lt;sup>3</sup>Actual quote from a referee

have to devise various approximations to what they were trying to characterize. The advent of Markov chain Monte Carlo (MCMC) methods has made it easier to calculate posterior distributions for just about any problem to arbitrary levels of precision.

Broadly speaking, MCMC is a class of methods for drawing random numbers (sampling or simulating) from the target posterior distribution. Thus, even though we might not recognize the posterior as a named distribution or be able to analyze its features analytically, e.g., devise mathematical expressions for the mean and variance, we can use these MCMC methods to obtain a large sample from the posterior and then use that sample to characterize features of the posterior. What we do with the sample depends on our intentions – typically we obtain the mean or median for use as a point estimate, and take a confidence interval based on Monte Carlo estimates of the quantiles. These are estimates, but not like frequentist estimates. Rather, they are Monte Carlo estimates with an associated Monte Carlo error which is largely determined arbitrarily by the analyst. They are not estimates qualified by a sampling distribution as in classical statistics. If we run our MCMC long enough then our reported value of  $E[\theta|y]$  or any feature of the posterior distribution is precisely what we say it is. There is no "sampling variation" in the frequentist sense of the word. In summary, the MCMC samples provide a Monte Carlo characterization of the posterior distribution.

## 2.5 WHAT GOES ON UNDER THE MCMC HOOD

We will develop and apply MCMC methods in some detail for spatial capturerecapture models in chapter 10. Here we provide a simple illustration of some basic ideas related to the practice of MCMC.

A type of MCMC method relevant to most problems is Gibbs sampling (REF XYZ XYZ), which is based on the idea of iterative simulation from the "full conditional" distributions (also called conditional posterior distributions). The full conditional distribution for an unknown quantity is the conditional distribution of that quantity given every other random variable in the model - the data and all other parameters. For example, for a normal regression model with  $y \sim \text{Normal}(\alpha + \beta x, 1)$  then the two full conditionals are, in symbolic terms,

 $[\alpha|y,\beta]$ 

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 $[\beta|y,\alpha].$ 

We might use our knowledge of probability to identify these mathematically. In particular, by Bayes' Rule,  $[\alpha|y,\beta] = [y|\alpha,\beta][\alpha|\beta]/[y|\beta]$  and similarly for  $[\beta|y,\alpha]$ . For example, if we have priors for  $[\alpha]$  and  $[\beta]$  which are also normal distributions, some algebra reveals that XXXX COPY NOTATION FFROM CH. 6 XXXXX

 $[\alpha|y,\beta] = Normal(ybar, ...weightedvariancehere...).$ 

Similarly,

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[\beta|y,\alpha]isnormal(....)
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The MCMC algorithm for this model has us simulate in succession, repeatedly, from those two distributions. See Gelman et al. (2004) for more examples of Gibbs sampling for the normal model. A conceptual representation of the MCMC algorithm for this simple model is therefore: XXXX Check out ALGORITHM environment XXXXX

```
Algorithm
```

O. Initialize \$\alpha\$ and \$\beta\$

```
Repeat{
```

}

- 1. Draw a new value of \$\alpha\$ from Eq. \ref{xyz}
- 2. Draw a new value of \$\beta\$ from Eq. \ref{xyz}

As we just saw for this simple "normal-normal" model it is sometimes possible to specify the full conditional distributions analytically. In general, when certain so-called conjugate prior distributions are chosen, the form of full conditional distributions is similar to that of the observation model. In this normal-normal case, the normal distribution for the mean parameters is the conjugate prior under the normal model, and thus the full-conditional distributions are also normal. This is convenient because, in such cases, we can simulate directly from them using standard methods (or R functions). But, in practice, we don't really ever need to know such things because most of the time we can get by using a simple algorithm, called the Metropolis-Hastings (henceforth "MH") algorithm, to obtain samples from these full conditional distributions without having to recognize them as specific, named, distributions. This gives us enormous freedom in developing models and analyzing them without having to resolve them mathematically because to implement the MH algorithm we need only identify the full conditional distribution up to a constant of proportionality, that being the marginal distribution in the denominator (e.g.,  $[y|\beta]$  above).

We will talk about the Metropolis-Hastings algorithm shortly, and we will use it extensively in the analysis of SCR models (e.g., chapter 10).

#### 2.5.1 Rules for constructing full conditional distributions

The basic strategy for constructing full-conditional distributions for devising MCMC algorithms can be reduced conceptually to a couple of basic steps summarized as follows:

(step 1) Collect all stochastic components of the model;

(step 2) Recognize and express the full conditional in question as proportional to the product of all components;

(step 3) Remove the ones that don't have the focal parameter in them.

(step 4) Do some algebra on the result in order to identify the resulting pdf or pmf.

Of the 4 steps, the last of those is the main step that requires quite a bit of statistical experience and intuition because various algebraic tricks can be used to reshape the mess into something noticeable - i.e., a standard, named distribution. But step 4 is not necessary if we decide instead to use the Metrpolis-Hastings algorithm as described below.

To illustrate for computing  $[\alpha|y,\beta]$  we first apply step 1 and identify the model components as:  $[y|\alpha,\beta]$ ,  $[\alpha]$  and  $[\beta]$ . Step 2 has us write  $[\alpha|y,\beta] \propto [y|\alpha,\beta][\alpha][\beta]$ . Step 3: We note that  $[\beta]$  is not a function of alpha and therefore we remove it to obtain  $[\alpha|y,\beta] \propto [y|\alpha,\beta][\alpha]$ . Similarly we obtain  $[\beta|y,\alpha] \propto [y|\alpha,\beta][\beta]$ . We apply step 4 and manipulate these algebraically to arrive at the result or, alternatively, we can sample them indirectly using the Metropolis-Hastings algorithm (see below).

## 2.5.2 Metropolis-Hastings algorithm

The Metropolis-Hastings (MH) algorithm is a completely generic method for sampling from any distribution, say  $f(\theta)$ . In our applications,  $f(\theta)$  will typically be the full conditional distribution of  $\theta$ . While we sometimes use Gibbs sampling, we seldom use "pure" Gibbs sampling because we might use MH to sample from one or more of the full conditional distributions. When the MH algorithm is used to sample from full conditional distributions of a Gibbs sampler the resulting hybrid algorithm is called Metropolized Gibbs Simple or more commonly Simple Sim

The MH algorithm generates candidates from some proposal or candidate-generating distribution, that may be conditional on the current value of the parameter, denoted by  $h(\theta^*|\theta^t)$ . Here,  $\theta^*$  is the candidate or proposed value and  $\theta^t$  is the current value, i.e., at iteration t of the MCMC algorithm. The proposed value is accepted with probability XXXX check notation with Rahel XXXXXX

$$r = \frac{f(\theta^*)h(\theta^t|\theta^*)}{f(\theta^t)h(\theta^*|\theta^t)}$$

which we call the MH acceptance probability. This ratio can sometimes be > 1 in which case we set it equal to 1. It is useful to note that h() can be anything at all. Absolutely anything! You can generate candidate values from a normal(0,1) distribution, from a uniform(-3455,3455) distribution, or anything of proper support. Note, however, that good choices of h() are those that approximate the posterior distribution. Obviously if  $h() = f(\theta|y)$  (i.e., the posterior) then you always accept the draw, and it stands to reason that proposals that are more similar to  $f(\theta|y)$ 

will lead to higher acceptance probabilities. No matter the choice of h(), we can evaluate this ratio numerically because the marginal f(y) cancels from both the numerator and denominator, which is the magic of the MH algorithm.

A special kind of h() are those that are symmetric, which means that h(a|b) = h(b|a) in which case h(a|b) and h(b|a) just cancel out from the MH acceptance probability and r is then just the ratio of the target density evaluated at the candidate value to that evaluated at the current value. A type of symmetric proposal useful in many situations is the so-called random-walk proposal distribution where candidate values are drawn from a normal distribution with mean equal to the current value and some standard deviation, say  $\delta$ , which is prescribed by the user. For parameters that have support on the real line, e.g.,  $\alpha$  in our example above, the random walk proposal generator has us generate  $\alpha^* \sim \text{Normal}(\alpha^t, \delta)$ . If we set  $\delta$  very small we have a high probability of accepting the proposal and vice versa. In practice, we "tune" delta to achieve a compromise between acceptance rate and efficient mixing of the Markov chains (see below for an example) normally assessed by autocorrelation. Low  $\delta$  increases the acceptance rate but will tend to produce Markov chains with high autocorrelation, and vice versa.

**Parameters with bounded support**: Many models contain parameters that have bounded support. E.g., variance parameters live on  $[0,\infty]$ , parameters that represent probabilities live on [0,1], etc.. In that case it is sometimes convenient to use a random walk proposal distribution that can generate any real number (e.g., a normal random walk proposal). In that case, we can just reject parameters that are outside of the parameter space (XXXX REF FOR THIS XXXX).

## 2.6 PRACTICAL BAYESIAN ANALYSIS AND MCMC

There are a number of really important practical issues to be considered in any Bayesian analysis and we cover some of these briefly here.

## 2.6.1 Choice of prior distributions

## XXX integrate this material with previous section on prior distributions XXXXXX

Bayesian analysis requires that we choose prior distributions for all of the structural parameters of the model (we use the term structural parameter to mean all parameters that aren't customary thought of as latent variables). We will strive to use priors that are meant to express little or no prior information - default or customary "non-informative" or diffuse priors. This will be  $\mathrm{Unif}(a,b)$  priors for parameters that have a natural bounded support and, for parameters that live on the real line we use either (1) diffuse normal priors; (2) "improper" uniform priors or (3) sometimes even a bounded  $\mathrm{Unif}(a,b)$  prior if that greatly improves the performance of **WinBUGS** or other software doing the MCMC for us. In **WinBUGS** a prior with low "precision",  $\tau$ , where  $\tau = 1/\sigma^2$ , such as  $\mathrm{Norm}(0,.01)$  will typically

be used. Of course  $\tau=0.01$  ( $\sigma^2=100$ ) might be very informative for a regression parameter that has a high variance. Therefore, we recommend that predictor variables *always* be standardized. Clearly there are a lot of choices for ostensibly non-informative priors, and the degree of non-informativeness depends on the parameterization. For example, a natural non-informative prior for the intercept of a logistic regression

$$logit(p_i) = \alpha + \beta x_i$$

Would be  $[\alpha] = \text{const}$  which is the same as saying  $a \sim Unif(\infty, infty)$ , the custom-ary improper uniform prior. However, we might also use a prior on the parameter  $p0 = logit^{-1}(a)$ , which is Pr(y = 1) for the value x = 0. Since p0 is a probability a natural choice is  $p0 \sim Unif(0,1)$ . These two priors can affect results (see Chapter 3.XYZ), yet they are both sensible non-informative priors. Choice of priors and parameterization is very much problem-specific and often largely subjective. More-over, it also affects the behavior of MCMC algorithms and therefore the analyst needs to pay some attention to this issue and possibly try different things out. XXX REFS on prior distributions XXXXXX

## 2.6.2 Convergence and so-forth

Once we have carried-out an analysis by MCMC, there are many other practical issues that we have to confront. One of the most important is "have the chains converged?" Most MCMC algorithms only guarantee that, eventually, the samples being generated will be from the target posterior distribution. So-called "convergence" of the Markov chain is achieved when that happens. Typically a period of transience is observed in the early part of the MCMC algorithm, and this is usually discarded as the "burn-in" period.

The quick diagnostic to whether convergence has been achieved is that your Markov chains look "grassy" – see Fig. 2.5 below. Another way to check convergence is to update the parameters some more and see if the posterior changes. It is good to confirm convergence using the "R-hat" statistic  $(\hat{R})$  or Brooks-Gelman-Rubin statistic (Gelman et al., 1996)) which should be close to 1 if the Markov chains have converged and sufficient posterior samples have been obtained. In practice,  $\hat{R} = 1.2$  is probably good enough for some problems. For some models you can't actually realize a low  $\hat{R}$ . E.g., if the posterior is a discrete mixture of distributions then you can be misled into thinking that your Markov chains have not converged when in fact the chains are just jumping back and forth in the posterior state-space. So, for example, using model selection methods (section XYZ) sometimes suggests non-convergence. Another situation is when one of the parameters is on the boundary of the parameter space which might appear to be very poor mixing, but all within some extreme region of the parameter space.<sup>4</sup> This

 $<sup>^4</sup>$ it would be nice if we could compile examples of this later in the book and reference back to this point

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kind of stuff is normally ok and you need to think really hard about the context of the model and the problem before you conclude that your MCMC algorithm is ill-behaved.

Some models exhibit "poor mixing" of the Markov chains or what people might also say "have not coverged" (or "slow convergence") which is a term we would disagree with because the samples might well be from the posterior (i.e., the Markov chains have converged to the proper stationary distribution) but simply mix around the posterior rather slowly. Anyway, poor mixing can happen for a huge number of reasons – when parameters are highly correlated (even confounded), or barely identified from the data, or the algorithms are very terrible and probably many other reasons. Slow mixing equates to high autocorrelation in the Markov chain the successive draws are highly correlated, and thus we need to run the MCMC algorithm much longer to get an effective sample size that is sufficient for estimation - or to reduce the MC error to a tolerable level. A strategy often used to reduce autocorrelation is "thinning" - i.e., keep every  $m^{th}$  value of the Markov chain output. However, thinning is necessarily inefficient from the stand point of inference - you can always get more precise posterior estimates by using all of the MCMC output regardless of the level of autocorrelation (MacEachern and Berliner, 1994). Practical considerations might necessitate thinning, even though it is statistically inefficient. For example, in models with many parameters or other unknowns being tabulated, the output files might be enormous and unwieldy to work with. In such cases, thinning is perfectly reasonable. In many cases, how well the Markov chains mix is strongly influenced by parameterization, standardization of covariates, and the prior distributions being used. Some things work better than others, and the investigator should experiment with different settings and remain calm when things don't work out perfectly. MCMC is an art, and a science.

Is the posterior sample large enough? A good rule of thumb is that you should never report MCMC results to more than 2 decimal places - because they will always be different! Look at the MC error which is printed by default in summaries of BUGS output. You want that to be smallish relative to the magnitude of the parameter and this might depend on the purpose of the analysis. For a preliminary analysis you might settle for a few percent whereas for a final analysis then certainly less than 1% is called for, but you can run your MCMC algorithm as long as it takes. Note that MC error in summaries of the posterior is not the same as having an "approximate" solution in a standard likelihood analysis or similar. The approximate SE in likelihood inference is actually wrong in its actual value....

## 2.6.3 Bayesian confidence intervals

The 95% Bayesian interval based on percentiles of the posterior is not a unique interval - there are many of them - and the so-called "highest posterior density" (HPD) interval is the narrowest interval. We might compute that frequently because

it is easy to do with an integer parameter which N is (See the next chapter). The 95 % HPD is not often exactly 95% but usually slightly more conservative than nominal because it is the narrowest interval that contains at least 95% of the posterior mass.

## 2.6.4 Estimating functions of parameters

A benefit of analysis by MCMC is that we can seamlessly estimate functions of parameters by simply tabulating the desired function of the simulated posterior draws. For example, if  $\theta$  is the parameter of interest and let  $\theta^{(i)}$  for  $i=1,2,\ldots,M$ be the posterior samples of  $\theta$ . Let  $\eta = exp(\theta)$ , then a posterior sample of  $\eta$  can be obtained simply by computing  $exp(\theta^{(i)})$  for i = 1, 2, ..., M. We give another example in section 2.7.2 below and throughout this book. Almost all SCR models in this book involve at least 1 derived parameter. For example, density D is a derived parameter, being a function of population size N and the area A of the underlying state-space of the point process (see chapter 5).

#### 2.7 BAYESIAN ANALYSIS USING WINBUGS

We won't be too concerned with devising our own MCMC algorithms for every analysis although we will do that a few times for fun. More often, we will rely on the freely available software package WinBUGS or JAGS for doing this. We will always execute these BUGS engines from within R using the R2WinBUGS (REF XYZ XYZ) or rjags pacages. WinBUGS and JAGS are MCMC black boxes that takes a pseudo-code description (i.e., written in the BUGS language) of all of the relevant stochastic and deterministic elements of a model and generates an MCMC algorithm for that model. But you never get to see the algorithm. Instead, WinBUGS/JAGS will run the algorithm and just return the Markov chain output - the posterior samples of model parameters.

The great thing about using the **BUGS** language is that it forces you to become intimate with your statistical model - you have to write each element of the model down, admit (explicitly) all of the various assumptions, understand what the actual probability assumptions are and how data relate to latent variables and data and latent variables relate to parameters, and how parameters relate to one another.

While we normally use **WinBUGS** or **JAGS** in this book, we note that **OpenBUGS** is the current active development tree of the **BUGS** language. See Kéry (2010, ch.xyz) and Kery and Schaub (2011, appendix xyz) for more on practial analysis in **WinBUGS**. That book should also be consulted for a more comprehensive introduction to using **WinBUGS**. In this example, we're going to accelerate pretty fast.

## 2.7.1 Linear Regression in WinBUGS

We provide a brief introductory example of a normal regression model using a small simulated data set. The following commands are executed from within your R

workspace, the command line being indicated by ">". First, simulate a covariate x and observations y having prescribed intercept, slope and variance:

```
779 > x<-rnorm(10)

780 > mu<- -3.2+ 1.5*x

781 > y<-rnorm(10,mu,sd=4)
```

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The **BUGS** model specification for a normal regression model is written within **R**as a character string input to the command cat() and then dumped to a text file
named normal.txt:

```
> cat("
785
   model {
       for (i in 1:10){
787
          y[i]~dnorm(mu[i],tau)
                                           # the "likelihood"
                                           # the linear predictor
          mu[i]<- beta0 + beta1*x[i]</pre>
789
         }
       beta0~dnorm(0,.01)
                                           # prior distributions
791
       beta1~dnorm(0,.01)
792
       sigma~dunif(0,100)
793
       tau<-1/(sigma*sigma)
                                           # tau is a derived parameter
794
   }
795
    ",file="normal.txt")
796
```

Alternatively, you can write the model specifications directly within a text file and save it in your current working directory, but we do not usually take that approach in this book.

Remarks: 1. WinBUGS parameterizes the normal in terms of the mean and inverse-variance, called the precision. Thus,  $\mathtt{dnorm}(0,.01)$  implies a variance of 100; 2. We typically use diffuse normal priors for mean parameters,  $\beta_0$  and  $\beta_1$  in this case, but sometimes we might use uniform priors with suitable bounds -B and +B. 3. We typically use a  $\mathtt{Unif}(0,B)$  prior on standard deviation parameters (Gelman XXX 2006 XXXX). But sometimes we might use a gamma prior on the precision parameter  $\tau$ . 4. In a **WinBUGS** model file, every variable referenced in the model description has to be either data, which will be input (see below), a random variable which must have a probability distribution associated with it using the "~", or it has to be a derived parameter connected to variables and data using "<-".

To fit the model, we need to describe various data objects to **WinBUGS**. In particular, we create an  $\mathbf{R}$  list object called **data** which are the data objects identified in the BUGS model file. In the example, the data consist of two objects which exist as y and x in the  $\mathbf{R}$  workspace and also in the **WinBUGS** model definition. We also have to create an  $\mathbf{R}$  function that produces a list of starting values **inits** that get sent to **WinBUGS**. Finally, we identify the names of the parameters (labeled correspondingly in the **WinBUGS** model specification) that

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we want **WinBUGS** to save the MCMC output for. In this example, we will "monitor" the parameters  $\beta_0$ ,  $\beta_1$ ,  $\sigma$  and  $\tau$ . **WinBUGS** is executed using the **R** command bugs(). We set the option debug=TRUE if we want the **WinBUGS** GUI to stay open (useful for analyzing MCMC output and looking at the **WinBUGS** error log). Also, we set working.dir=getwd() so that **WinBUGS** output files and the log file are saved in the current **R** working directory. All of these activities look like this:

```
library("R2WinBUGS") # "attach" the R2WinBUGS library

data <- list ( "y","x")

inits <- function()

list ( beta1=rnorm(1),beta0=rnorm(1),sigma=runif(1,0,2) )

parameters <- c("beta0","beta1","sigma","tau")

out<-bugs (data, inits, parameters, "normal.txt", n.thin=2, n.chains=2,

n.burnin=2000, n.iter=6000, debug=TRUE,working.dir=getwd())
```

Remarks: A common question is "how should my data be formatted?" That depends on how you describe the model in the BUGS language, how your data are input into R and subsequently formatted. There is no unique way to describe any particular model and so you have some flexibility. We talk about data format further in the context of capture-recapture models and SCR models in chapter 5 and elsewhere. In general, starting values are optional but we recommend to always provide reasonable starting values for structural parameters, but are not always necessary for random effects. Note that the previously created objects defining data, initial values and parameters to monitor are passed to the function bugs(). In addition, various other things are declared: The number of Markov chains (n.chains), the thinning rate (n.thin), the number of burn-in iterations (n.burnin) and the total number of iterations (n.iter). To develop a detailed understanding of the various parameters and settings used for MCMC, consult a basic reference such as Kéry (2010).

You should execute all of the commands given above and then look at the resulting output. Kill the **WinBUGS** GUI and the data will be read back into **R** (or specify debug=FALSE). We don't want to give instructions on how to navigate and use the GUI - see XYZ REF (XYZ) for that. The object out prints important summaries by default (this is slightly edited):

```
> print(out,digits=2)
851
    Inference for Bugs model at "normal.txt", fit using WinBUGS,
852
     2 chains, each with 6000 iterations (first 2000 discarded), n.thin = 2
853
     n.sims = 4000 iterations saved
854
                      sd 2.5%
                                 25%
                                              75% 97.5% Rhat n.eff
              mean
                                        50%
    beta0
             -2.43 1.84 -6.21 -3.50 -2.42 -1.34
                                                   1.27
                                                               4000
    beta1
              2.62 1.54 -0.42
                                1.68
                                       2.62
                                             3.57
                                                   5.67
                                                            1
                                                               4000
857
              5.29 1.66
                          3.11
                                4.14
                                       4.95
                                             6.05
                                                   9.39
                                                               4000
858
    sigma
              0.05 0.02 0.01 0.03
                                      0.04
                                             0.06
                                                   0.10
                                                            1
                                                               4000
859
    deviance 59.85 3.24 56.18 57.47 59.00 61.37 68.32
                                                                840
```

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869 870

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```
For each parameter, n.eff is a crude measure of effective sample size, and Rhat is the potential scale reduction factor (at convergence, Rhat=1).

BIC info (using the rule, pD = Dbar-Dhat)

pD = 2.6 and DIC = 62.4
```

**Remarks:** (1) convergence is assessed using the  $\hat{R}$  statistic – which we might sometimes write "Rhat". A value of Rhat near 1 indicates convergence; (2) DIC is the "deviance information criterion" (Spiegelhalter et al., 2002) (see section 2.8) which some people use in a manner similar to AIC although it is recognized to have some problems in hierarchical models (Millar, 2009). We evaluate this in the context of SCR models in chapter XYZ XYZ.

## 2.7.2 Inference about functions of model parameters

Using the MCMC draws for a given model we can easily obtain the posterior distribution of any function of model parameters. We showed this in the above example by providing the posterior of  $\tau$  when the model was parameterized in terms of standar deviation  $\sigma$ . As another example, suppose that the normal regression model above had a quadratic response function of the form

$$E(y_i) = \beta_0 + \beta_1 x_i + \beta_2 x_i^2$$

Then the optimum value of x, i.e., that corresponding to the optimal expected response, can be found by setting the derivative of this function to 0 and solving for x. We find that

$$df/dx = \beta_1 + 2 * \beta_2 x = 0$$

yields that  $x_{opt} = -\beta_1/(2 * \beta_2)$ . We can just take our posterior draws for  $beta_1$  and  $beta_2$  and obtain a posterior sample of  $x_{opt}$  by this simple calculation. As an exercise, take the normal model above and simulate a quadratic response and then describe the posterior distribution of  $x_{opt}$ .

### 2.8 MODEL CHECKING AND SELECTION

In general terms model checking - or assessing the adequacy of the model - and model selection are quite thorny issues and, despite contrary and, sometimes, strongly held belief among practitioners, there are not really definitive, general solutions to either problem. We're against dogma on these issues and think people need to be open-minded about such things and recognize that models can be useful whether or not they pass certain statistical tests. Some models are intrinsically better than others because they make more biological sense or foster understanding or achieve some objective that some bootstrap or other goodness-of-fit test can't

decide for you. That said, it gives you some confidence if your model seems adequate and we try to provide some fit assessment in most real applications of SCR models We provide a very brief overview of concepts here, but provide more detailed coverage in chapter 12. See also Kéry (2010, ch. xyz) and Link and Barker (2009, ch. xyz) for specific context related to Bayesian model checking and selection.

#### 2.8.1 Goodness-of-fit

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Goodness-of-fit testing is an important element of any analysis because our model represents a general set of hypotheses about the ecological and observation processes that generated our data. Thus, if our model "fits" in some statistical or scientific sense, then we believe it to be consistent with the hypotheses that went into the model. More formally, we would conclude that the data are not inconsistent with the hypotheses, or that the model appears adequate. If we have enough data, then of course we will reject any set of statistical hypotheses. Conversely, we can always come up with a model that fits by making the model extremely complex. Despite this paradox, it seems to us that simple models that you can understand should usually be preferred even if they don't fit, for example if they embody essential mechanisms central to our understanding of things, or if we think that some contributing factors to lack-of-fit are minor or irrelevant to the scientific context and intended use of the model. In other words, models can be useful irrespective of whether they fit according to some formal statistical test of fit. Yet the tension is there to obtain fitting models, and this comes naturally at the expense of models that can be easily interpreted and studied and effectively used. Moreover, conducting goodness-of-fit tests is not always so easy to do. Moreover, it is never really easy (or especially convenient) to decide if your goodness-of-fit test is worth anything. It might have 0 power! Despite this, we recommend attempting to assess model fit in real applications, as a general rule, and we provide some basic guidance here and some more specific to SCR models in chapter 12.

To evaluate goodness-of-fit in Bayesian analyses, we will most often use the Bayesian p-value (Gelman et al., 1996). The basic idea is to define a fit statistic or "discrepancy measure" and compare the posterior distribution of that statistic to the posterior predictive distribution of that statistic for hypothetical perfect data sets for which the model is known to be correct. For example, with count frequency data, a standard measure of fit is the sum of squares of the "Pearson residuals",

$$D(y_i, \theta) = \frac{(y_i - E(y_i))^2}{Var(y_i)}$$

The fit statistic based on the squared residuals is

$$FIT = \sum_{i} D(y_i, \theta)^2$$

which can be computed at each iteration of a MCMC algorithm given the current values of parameters that determine the response distribution. At the same time (i.e., at each MCMC iteration), the equivalent statistic is computed for a "new" data set, simulated using the current parameter values. The Bayesian p-value is simply the posterior probability  $Pr(Fit > Fit_{new})^5$  which should be close to 0.50 for a good model – one that "fits" in the sense that the observed data set is consistent with realizations simulated under the model being fitted to the observed data. In practice we judge "close to 0.50" as being "not too close to 0 or 1" and, as always, closeness is somewhat subjective. We're happy with anything > .1 and < .9 but might settle for > .05 and < 0.95. In summary, the Bayesian p-value seems like a bootstrap idea, is easy to compute, and widely used as a result.

Another useful fit statistic is the Freeman-Tukey statistic<sup>6</sup>, in which

$$D(\mathbf{y}, \theta) = \sum_{i} (\sqrt{y_i} - \sqrt{e_i})^2$$

(Brooks et al., 2000), where  $y_i$  is the observed value of observation i and  $e_i$  its expected value. In contrast to a chi-square discrepancy, the Freeman-Tukey statistic removes the need to pool cells with small expected values.

#### 2.8.2 Model Selection

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For model selection we typically use three different methods: First is, let's say, common sense. If a parameter has posterior mass concentrated away from 0 then it seems like it should be regarded as important - that is, it is "significant." This approach seems to have fallen out of favor with all of the interest over the last 10 or 15 years on model selection in ecology. It seems reasonable to us.

For regression problems we sometimes use the factor weighting idea which is to introduce a set of binary variables  $w_k$  for variable k, and express the model as, e.g., for a single covariate model:

$$E(y_i) = \alpha + w\beta x_i$$

where w is given a Bernoulli prior distribution with some prescribed probability. E.g.,  $w \sim Bern(0.50)$  to provide a prior probability of 0.50 that variable x should be an element of the linear predictor. The posterior probability of the event w=1 is a gauge of the importance of the variable x. i.e., high values of Pr(w=1) indicate stronger evidence to support that "x is in the model" whereas values of Pr(w=1) close to 0 suggest that x is less important.

This idea seems to be due to Kuo and Mallick (1998)<sup>7</sup> and see Royle and Dorazio (2008, ch. XXXX) for an example in the context of logistic regression. This

<sup>&</sup>lt;sup>5</sup>Check this definition!

<sup>&</sup>lt;sup>6</sup>Ref for this?

<sup>&</sup>lt;sup>7</sup> Is this also what people call Zellner's G-priors?

approach seems to even work sometimes with fairly complex hierarchical models of a certain form. E.g., Royle (2008) applied it to a random effects model to evaluate the importance of the random effect component of the model. The main problem with this approach is that its effectiveness and results will typically be highly sensitive to the prior distribution on the structural parameters (e.g., see Royle and Dorazio (2008, table xyz)). The reason for this is obvious: If w=0 for the current iteration of the MCMC algorithm, so that  $\beta$  is sampled from the prior distribution, and the prior distribution is very diffuse, then extreme values of  $\beta$  are likely. Consequently, when the current value of  $\beta$  is far away from the mass of the posterior when w=1, then the Markov chain may only jump from w=0 to w=1 infrequently. One seemingly reasonable solution to this problem (Aitken XYZ FIND THIS XXXXX<sup>8</sup>) is to fit the full model to obtain posterior distributions for all parameters, and then use those as prior distributions in a "model selection" run of the MCMC algorithm. This seems preferable to more-or-less arbitrary restriction of the prior support to improve the performance of the MCMC algorithm.

A third method that that we advocate is subject-matter context. It seems that there are some situations – some models – where one should not have to do model selection because it is necessitated by the specific context of the problem, thus rendering a formal hypotesis test pointless (Johnson, 1999). SCR models are such an example. In SCR models, we will see that "spatial location" of individuals is an element of the model. The simpler, reduced, model is an ordinary capture-recapture model which is not spatially explicit (i.e., chapter 11), but it seems silly and pointless to think about actually using the reduced model even if we could concoct some statistical test to refute the more complex model. The simpler model is manifestly wrong but, more importantly, not even a plausible data-generating model! Other examples are when effort, area or sample rate is used as a covariate. One might prefer to have such things in models regardless of whether or not they pass some statistical litmus test (although one can always find referees to argue for pedantic procedure over thinking).

Many problems can be approached using one of these methods but there are also broad classes of problems that can't and, for those, you're on your own. In later chapters we will address model selection in specific contexts and we hope those will prove useful for a majority of the situations you encounter.

## 2.9 POISSON GLMS

The Poisson GLM (also known as "Poisson regression") is probably the most relevant and important class of models in all of ecology. The basic model assumes observations  $y_i; i = 1, 2, ..., n$  follow a Poisson distribution with mean  $\lambda$  which we write

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

<sup>&</sup>lt;sup>8</sup>see Royle 2008 paper for reference

Commonly  $y_i$  is a count of animals or plants at some point in space and lambda might depend on i. For example, i might index point count locations in a forest, BBS route centers, or sample quadrats, or similar. If covariates are available it is typical to model them as linear effects on the log mean. If x(i) is some measured covariate associated with observation i. Then,

$$log(x(i)) = \alpha + \beta * x(i)$$

While we only specify the mean of the Poisson model directly, the Poisson model (and all GLMs) has a "built-in" variance which is directly related to the mean. In this case,  $Var(y) = E(y) = \lambda$ . Thus the model accommodates a linear increase in variance with the mean.

## 2.9.1 Important properties of the Poisson distribution

There are two properties of the Poisson distribution that make it extremely useful in ecology. First is the property of compound additivity. If  $y_1$  and  $y_2$  are Poisson random variables with means  $\lambda_1$  and  $\lambda_2$ , then their sum  $N=y_1+y_2$  is Poisson with mean  $\lambda_1+\lambda_2$ . Thus, if the observations can be viewed as an aggregate of counts over some finer unit of measurement, then the mean aggregates in a corresponding manner. Secondly, the Poisson distribution has a direct relationship to the multinomial. If  $y_1$  and  $y_2$  are iid Poisson then, conditional on their sum  $N=y_1+y_2$ , their joint distribution is multinomial with sample size N and cell probabilities  $\lambda_1/(\lambda_1+\lambda_2)$  and  $\lambda_2/(\lambda_1+\lambda_2)$ . As a result of this, most multinomial models can be analyzed as a Poisson GLM and vice versa.

## 2.9.2 Example: Breeding Bird Survey Data

As an example we consider a classical situation in ecology where counts of an organism are made at a collection of spatial locations. In this particular example, we have mourning dove counts made along North American Breeding Bird Survey (BBS) routes in Pennsylvania, USA. A route consists of 50 stops separated by 0.5 mile. For the purposes here we are defining  $y_i$  = route total count and he sample location will be marked by the center point of the BBS route. The survey is run annually and the data set we have is 1966-1998. BBS data can be obtained online at http://www.pwrc.usgs.gov/bbs/. We will make use of the whole data set shortly but for now we're going to focus on a specific year of counts – 1990 – for the sake of building a simple model. For 1990 there were 77 active routes. We have the data stored in a .csv file<sup>9</sup> where rows index the unique route, column 1 is the route ID, columns 2-3 are the route coordinates (longitude/latitude), column 4 is a habitat covariate "forest cover" (standardized, see below) and the remaining columns are the yearly counts. Years for which a route was not run are coded as "NA" in the data matrix. We imagine that this will be a typical format for many ecological

<sup>&</sup>lt;sup>9</sup>check this data format

studies, perhaps with more columns representing covariates. To read in the data and display the first few elements of this matrix, do this:

It is useful to display the spatial pattern in the observed counts. For that we use a spatial dot plot - where we plot the coordinates of the observations and mark the color of the plotting symbol based on the magnitude of the count. We have a special plotting function for that which is called  $\mathtt{spatial.plot}()$  and it is available with the supplemental  $\mathbf R$  package. Actually, what we want to do here is plot the log-count (+1 of course) which (Fig. 2.1) displays a notable pattern that could be related to something. The  $\mathbf R$  commands for obtaining this figure are:

```
data<-read.csv("pa-bbsdovedata-all.csv")

y<-data[,29] # pick out 1990

notna<-!is.na(y)

y<-y[notna]

spatial.plot(data[notna,2:3],y)
```

We can ponder the potential effects that might lead to dove counts being high....corn fields, telephone wires, barn roofs along with misidentification of pigeons, these could all correlated reasonably well with the observed count of mourning doves. Unfortunately we don't have any of that information.

We do have a measure of forest cover in the vicinity of each point which is contained in the data set (variable "habitat"). This was derived from a larger GIS coverage of the state (provided in the data file "pahabdata.csv") which can be plotted using the spatial.plot function using the following commands

```
1060 > map('state',regions="penn",lwd=2)
1061 > spatial.plot(pahabdata[,2:3],pahabdata[,"dfor"],cx=2)
1062 > map('state',regions="penn",lwd=2,add=TRUE)
```

where the result appears in Fig. 2.2. We see a prominent pattern that indicates high forest coverage in the central part of the state and low forest cover in the SE. Inspecting the previous figure of log-counts suggests a relationship between counts and forest cover which is perhaps not surprising.

## 2.9.3 Doing it in WinBUGS

Here we demonstrate how to fit a Poisson GLM in **WinBUGS** using the covariate  $x_i$  = forest cover. It is advisable that  $x_i$  be standardized in most cases as this will improve mixing of the Markov chains. Recall that the data we have stored include a standardized covariate (forest cover) and so we don't have to worry about that here. To read the BBS data into **R** and get things set up for **WinBUGS** we issue the following commands:

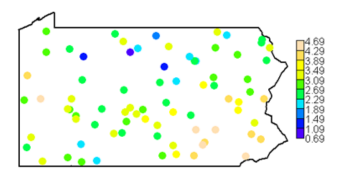


Figure 2.1. Needs a caption

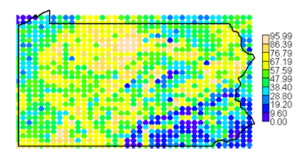


Figure 2.2. Needs a caption

```
data<-read.csv("pa-bbsdovedata-all.csv")
1074
    v<-data[,29]
                                            # pick out 1990
1075
    notna<-!is.na(y)
1076
    y<-y[notna]
                                            # discard missing
1077
    habitat <- data[notna, 4]
                                              get habitat data
1078
    library("R2WinBUGS")
                                              load R2WinBUGS
1079
    data <- list ( "y", "M", "habitat")</pre>
                                            # bundle data for WinBUGS
1080
        Now we write out the Poisson model specification in WinBUGS pseudo-code.
1081
    provide initial values, identify parameters to be monitored and then execute Win-
1082
    BUGS:
1083
    cat("
1084
    model {
1085
         for (i in 1:M){
1086
           y[i]~dpois(lam[i])
1087
           log(lam[i])<- beta0+beta1*habitat[i]</pre>
1088
1089
1090
     beta0~dunif(-5,5)
     beta1~dunif(-5,5)
1091
    }
1092
    ",file="PoissonGLM.txt")
1093
    inits <- function() list ( beta0=rnorm(1),beta1=rnorm(1))</pre>
    parameters <- c("beta0", "beta1")
1096
    out <- bugs (data, inits, parameters, "PoissonGLM.txt", n.thin=2,n.chains=2,
1097
                      n.burnin=2000,n.iter=6000,debug=TRUE,working.dir=getwd())
1098
        Remarks: (1) Note the close correspondence in how the model is specified
1099
    here compared with the normal regression model previously. As an exercise you
    should discuss the specific differences between the BUGS model specifications for
    the normal and Poisson models.
1102
    > print(out,digits=3)
1103
    Inference for Bugs model at
1104
    "'PoissonGLM.txt", fit using WinBUGS,
1105
     2 chains, each with 4000 iterations (first 1000 discarded), n.thin = 2
     n.sims = 3000 iterations saved
1107
                  mean
                            sd
                                    2.5%
                                               25%
                                                         50%
                                                                   75%
                                                                           97.5% Rhat n.eff
1108
    beta0
                 3.151
                         0.025
                                   3.102
                                             3.135
                                                       3.151
                                                                 3.168
                                                                           3.199 1.001
1109
    beta1
                -0.498
                         0.021
                                  -0.539
                                            -0.512
                                                      -0.498
                                                                -0.484
                                                                          -0.457 1.001
                                                                                         3000
1110
    fit
               869 930 19 856
                                 835 500
                                           855.700
                                                     868 600
                                                               881 900
                                                                        913.602 1.002
                                                                                         1600
1111
                                            68.107
                76.709 12.519
                                  54.098
                                                      76.215
                                                                84.510
                                                                        102,602 1,001
                                                                                         3000
1112
    fitnew
    deviance 1116.605 2.014 1115.000 1115.000 1116.000 1117.000 1122.000 1.001
                                                                                         3000
1113
```

We might wonder whether this model provides an adequate fit to our data. To evaluate that, we used a Bayesian p-value analysis with fit statistic based on the Freeman-Tukey residual by replacing the model specification above with this:

1114

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1144

1145

1146

```
cat("
1117
     model {
1118
          for (i in 1:M){
1119
            y[i]~dpois(lam[i])
            log(lam[i])<- beta0+beta1*habitat[i]</pre>
1121
            d[i] \leftarrow pow(pow(y[i], 0.5) - pow(lam[i], 0.5), 2)
1122
1123
            ynew[i]~dpois(lam[i])
1124
            dnew[i] <-pow( pow(ynew[i],0.5)-pow(lam[i],0.5),2)</pre>
1125
1126
           }
1127
1128
      fit<-sum(d[])
       fitnew<-sum(dnew[])
1129
      beta0~dunif(-5,5)
1130
      beta1~dunif(-5,5)
1131
     }
1132
     ",file="PoissonGLM.txt")
1133
```

The Bayesian p-value is the proportion of times fitnew > fit which, for this data set, is 0, which was 1.0 in this case (calculation omitted). This suggests that the basic Poisson model does not fit well.

# 2.9.4 Constructing your own MCMC algorithm

It might be helpful to suffer through a couple examples building custom MCMC algorithms. Here, we develop an MCMC algorithm for the Poisson regression model, using a Metropolis-within-Gibbs sampling framework.

We will assume that the two parameters have diffuse normal priors, say  $[\alpha] = \text{Norm}(0, 100)$  and  $[\beta] = \text{Norm}(0, 100)$  where each has standard deviation 100 (recall that **WinBUGS** parameterizes the normal in terms of  $1/\sigma^2$ ). We need to assemble the relevant elements of the model which are these two prior distributions and the likelihood  $[\mathbf{y}|\alpha,\beta] = \prod_i [y_i|\alpha\beta]$  which is, mathematically, the product of the Poisson pmf evaluated at each  $y_i$ , given particular values of  $\alpha$  and  $\beta$ . Next, we need to identify the full conditionals  $[\alpha|\beta,\mathbf{y}]$  and  $[\beta|\alpha,\mathbf{y}]$ . We use the all-purpose rule for constructing full conditionals (section 2.5.1) to discover that:

$$[\alpha|\beta, \mathbf{y}] \propto \left\{ \prod_{i} [y_i|\alpha, \beta] \right\} [\alpha]$$

1149 and

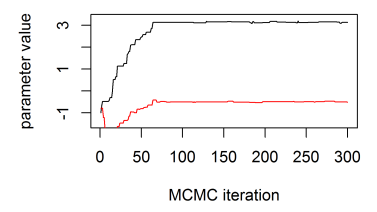
$$[\beta|\alpha, \mathbf{y}] \propto \left\{ \prod_{i} [y_i|\alpha, \beta] \right\} [\beta]$$

Remember, we could replace the " $\propto$ " with "=" if we put  $[y|\beta]$  or  $[y|\alpha]$  in the denominator. But, in general,  $[y|\alpha]$  or  $[y|\beta]$  will be quite a pain to compute and, more

importantly, it is a constant as far as the operative parameters ( $\alpha$  or  $\beta$ , respectively) 1152 are concerned. Therefore, the MH acceptance probability will be the ratio of the 1153 ful-conditional evaluated at a candidate draw to that evaluated at the current draw, 1154 and so the denominator required to change  $\propto$  to = winds up canceling from the MH acceptance probability. Here we will use the random walk candidate generator 1156 so that, for example,  $\alpha^* \sim \text{Normal}(\alpha^t, \delta)$  where  $\delta$  is the standard-deviation of the 1157 proposal distribution, which is just a tuning parameter<sup>10</sup>. We remark also that cal-1158 culations are often done on the log-scale to preserve numerical integrity of things 1159 when quantities evaluate to small or large numbers, so keep in mind, for example, 1160 a\*b = exp(log(a) + log(b)). The "Metropolis within Gibbs" algorithm for a Poisson 1161 regression turns out to be remarkably simple: 1162

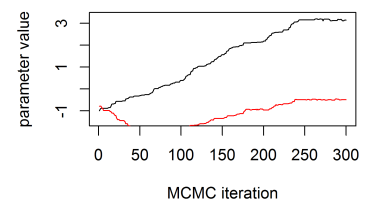
```
set.seed(2013)
1163
     out <- matrix (NA, nrow=1000, ncol=2)
                                              # matrix to store the output
1165
     alpha<- -1
                                              # starting values
1166
     beta <- -.8
1167
1168
     # begin the MCMC loop; do 1000 iterations
1169
     for(i in 1:1000){
1170
     # update the alpha parameter
1172
     lambda<- exp(alpha+beta*habitat)</pre>
1173
     lik.curr<- sum(log(dpois(y,lambda)))</pre>
1174
     prior.curr<- log(dnorm(alpha,0,100))</pre>
1175
     alpha.cand<-rnorm(1,alpha,.25)
                                                   # generate candidate
1176
     lambda.cand<- exp(alpha.cand + beta*habitat)</pre>
1177
     lik.cand<- sum(log(dpois(y,lambda.cand)))</pre>
1178
    prior.cand<- log(dnorm(alpha.cand,0,100))</pre>
1179
    mhratio<- exp(lik.cand +prior.cand - lik.curr-prior.curr)</pre>
1180
     if(runif(1)< mhratio)</pre>
1181
          alpha<-alpha.cand
1182
1183
     # update the beta parameter
     lik.curr<- sum(log(dpois(y,exp(alpha+beta*habitat))))</pre>
1185
     prior.curr <- log(dnorm(beta,0,100))
1186
     beta.cand<-rnorm(1,beta,.25)
1187
     lambda.cand<- exp(alpha+beta.cand*habitat)</pre>
1188
     lik.cand<- sum(log(dpois(y,lambda.cand)))</pre>
1189
     prior.cand<- log(dnorm(beta.cand,0,100))</pre>
1190
     mhratio <- exp(lik.cand + prior.cand - lik.curr - prior.curr)
1191
     if(runif(1)< mhratio)</pre>
1192
          beta<-beta.cand
1193
1194
```

 $<sup>^{10}</sup>$  It would help lots of people out to see a non-symmetric proposal distribution, and the extra step needed to account for it.



**Figure 2.3.** MCMC output for Poisson regression parameters (top trace: intercept  $\alpha$ ; bottom trace: slope  $\beta$ ). This is for  $\delta = 0.25$ .

The first 300 iterations of the MCMC history of each parameter is shown in Fig. 2.3. The appearance of this is not very appealing but a couple of things are evident: First, the Markov chains clearly stabilize - "burn-in" – after about 60 or 70 iterations. They also appear to mix very slowly once convergence is achieved, although this is not so clear given the scale of the y-axis. We decreased the standard deviation of the candidate generating distribution from  $\delta=0.25$  to  $\delta=0.05$  and re-ran re-ran the MCMC algorithm producing the output shown in Fig. 2.4. We see that the burn-in takes longer but it seems to mix better although it takes slightly longer to burn-in. Using this value of  $\delta$  we generated 10,000 posterior samples, discarding the first 500 as burn-in, and the result is shown in Fig. 2.5, this time seperate panels for each parameter. The "grassy" look of the MCMC history is diagnostic of Markov chains that are well-mixing and we would generally be very



**Figure 2.4.** Same as previous fig but with  $\delta = 0.05$ .

satisfied with results that look like this.

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**Remarks:** We used a specific set of starting values for these simulations. It should be clear that starting values closer to the mass of the posterior distribution might cause burn-in to occur faster. As an exercise, evaluate that. (2) Clearly the influence of the proposal standard deviation term is important. Small values lead to much better mixing but it should be noted that values that are too small will slow down burn-in and also lead to high correlation. This suggests there is an optimal value of the Metropolis-Hastings tuning parameter<sup>11</sup>. As an exercise you should contemplate finding that optimal value for this problem<sup>12</sup> (3) For the flat normal prior distributions here we could leave the prior contribution out of the full conditional evaluation since it is locally constant, i.e., constant in the vicinity of the posterior mass, and thus has no practical effect. Removing the prior contribution from the MH acceptance probability is equivalent to saying that the parameters have an improper uniform prior, i.e.,  $\alpha \sim \text{const}$ , which is commonly used for mean parameters in practice. Note also that we have used a different prior than in our WinBUGS model specification given previously. As an exercise, evaluate whether this seems to affect the result.

<sup>&</sup>lt;sup>11</sup>Defined previously?????

<sup>&</sup>lt;sup>12</sup>effective sample size definition?

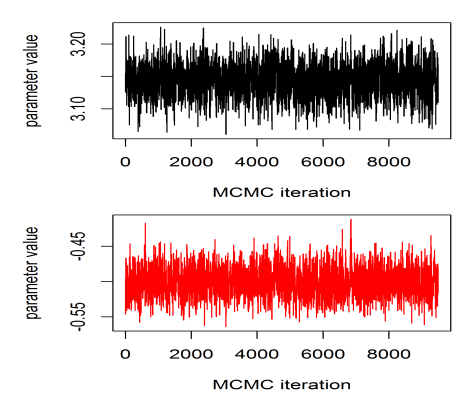


Figure 2.5. nice grassy mcmc output, longer run of previous with  $\delta=0.05$ .

#### 2.10 POISSON GLM WITH RANDOM EFFECTS

What we will be doing in most of this book is dealing with random effects in GLM-like models - similar to what are usually referred to as generalized linear mixed models (GLMMs). We provide a brief introduction by way of example, extending our Poisson regression model to include a random effect.

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The Log-Normal mixture: The classical situation involves a GLM with a normally distributed random effect that is additive on the linear predictor. For the Poisson case, we have:

$$log(\lambda_i) = \alpha + \beta x_i + \eta_i$$

where  $\eta_i \sim \text{Normal}(0, \sigma^2)$ . A natural alternative is to have multiplicative gamma-1239 distributed noise,  $exp(\eta_i) \sim \text{Gamma}(a,b)$  which would correspond to a negative 1240 binomial kind of over-dispersion, implying a different mean/variance relationship to the log-normal mixture (the interested reader should work that out). Choosing between such possibilities is not a topic we will get into here because it doesn't seem 1243 possible to provide general guidance on it. For this model we carried-out a goodness-1244 of-fit evaluation using the Bayesian p-value based on a Pearson residual statistic. 1245 See also (Kéry, 2010, ch. 18) for an example involving a binomial mixed model<sup>13</sup>. 1246 Anyhow, it is really amazingly simple to express this model in WinBUGS and 1247 have WinBUGS draw samples from the posterior distribution using the following 1248 code for the BBS dove counts:

```
data<-read.csv("pa-bbsdovedata-all.csv")
1250
     locs<-data[,2:3]
1251
     habitat <-data[,4]
1252
     y<-data[,29]
                          # grab year 1990
1253
     M<-length(y)
1254
1255
     set.seed(2013)
1256
1257
     cat("
1258
     model {
1259
       for (i in 1:M){
1260
           y[i]~dpois(lam[i])
1261
           log(lam[i])<- alpha+ beta*habitat[i] + eta[i]</pre>
1262
           frog[i] <-beta*habitat[i] + eta[i]</pre>
1263
           eta[i] ~ dnorm(0,tau)
1264
           d[i] \leftarrow pow(pow(y[i], 0.5) - pow(lam[i], 0.5), 2)
1265
1266
           ynew[i]~dpois(lam[i])
1267
           dnew[i] <- pow(pow(ynew[i],0.5)-pow(lam[i],0.5),2)</pre>
1268
1269
         }
1270
      fit<-sum(d[])
```

<sup>&</sup>lt;sup>13</sup>Kery has noticed that such tests probably have 0 power. Should use the marginal frequency of the data

```
fitnew<-sum(dnew[])
1271
1272
      alpha~dunif(-5,5)
1273
      beta~dunif(-5,5)
1274
      sigma~dunif(0,10)
1275
      tau<-1/(sigma*sigma)
1276
1277
1278
     ",file="model.txt")
1279
     data <- list ( "y", "M", "habitat")</pre>
1280
     inits <- function()</pre>
1281
       list ( alpha=rnorm(1),beta=rnorm(1),sigma=runif(1,0,4))
1282
     parameters <- c("alpha","beta","sigma","tau","fit","fitnew")</pre>
1283
     library("R2WinBUGS")
1284
1285
     out<-bugs (data, inits, parameters, "model.txt", n.thin=2,n.chains=2,
1286
      n.burnin=1000,n.iter=5000,debug=TRUE)
1287
        This produces the following posterior summary statistics:
1288
     > print(out,digits=2)
1289
     Inference for Bugs model at "model.txt", fit using WinBUGS,
1290
      2 chains, each with 5000 iterations (first 1000 discarded), n.thin = 2
1291
      n.sims = 4000 iterations saved
1292
                                        25%
                                                50%
                                                             97.5% Rhat n.eff
1293
                 mean
                          sd
                               2.5%
                                                        75%
     alpha
                 2.98
                       0.08
                               2.82
                                       2.93
                                               2.98
                                                      3.03
                                                              3.12 1.00
                                                                          1400
1294
                                                      -0.49
     beta
                -0.53
                       0.07
                              -0.68
                                      -0.58
                                              -0.53
                                                             -0.38 1.01
                                                                            350
1295
                                                      0.64
     sigma
                 0.60
                       0.06
                               0.49
                                       0.56
                                               0.59
                                                              0.73 1.00
                                                                           2000
1296
                 2.88
                       0.57
                               1.88
                                       2.47
                                               2.86
                                                      3.24
                                                              4.12 1.00
                                                                           2000
    t.au
1297
                26.58
                                      23.96
                                                     29.01
                                                             34 46 1 00
1298
    fit.
                       3.72
                              19.87
                                              26.37
                                                                           4000
               26.83 3.90 19.60 24.12 26.68
                                                     29.36
                                                             35.04 1.00
                                                                           4000
    fitnew
     deviance 445.94 12.18 424.00 437.40 445.20 453.90 471.50 1.00
1301
     [... some output deleted ...]
1302
        The Bayesian p-value for this model is
1303
     > mean(out$sims.list$fit>out$sims.list$fitnew)
1304
     [1] 0.4815
1305
    indicating a pretty good fit. Given the site-level random effect, it would be surpris-
1306
     ing for this model to not fit! One thing we notice is that the posterior standard
     deviations of the regression parameters are much higher, a result of the excess vari-
     ation. Www would also notice much less precise predictions of hypothetical new
1309
     observations.
1310
1311
```

ANDY STOPPED HERE.

#### 2.11 BINOMIAL GLMS

Another extremely important class of models in ecology are binomial models. We use binomial models for count data whenever the observations are counts or frequencies and it is natural to condition on a "sample size", say K, the maximum frequency possible in a sample. The random variable,  $y \leq K$ , is then the frequency of occurrences out of K "trials". The parameter of the binomial models is p, often called "success probability" which is related to the expected value of y by E(y) = pK. Usually we are interested in modeling covariates that affect the parameter p, and such models are called binomial GLMs , binomial regression models or logistic regression, although logistic regression really only applies when the logistic link is used to model the relationship between p and covariates (see below).

One of the most typical binomial GLMs occurs when the sample size equals 1 and the outcome, y, is "presence" (y=1) or "absence" (y=0) of a species. This is a classical "species distribution" modeling situation. A special situation occurs when presence/absence is observed with error (MacKenzie et al., 2002; Tyre et al., 2003). In that case, K>1 samples are usually needed for effective estimation of model parameters.

In standard binomial regression problems the sample size is fixed by design but interesting models also arise when the sample size is itself a random variable. These are the N-mixture models (Royle, 2004a; Kéry et al., 2005; Royle and Dorazio, 2008; Kéry, 2010) and related models (in this case, N being the sample size, which we labeled K above)<sup>14</sup>. Another situation in which the binomial sample size is "fixed" is closed population capture-recapture models in which a population of individuals is sampled K times. The number of times each individual is encountered is a binomial outcome with parameter - encounter probability – p, based on a sample of size K. In addition, the total number of unique individuals observed, n, is also a binomial random variable based onpopulation size N. We consider such models in the chapter 11.

# 2.11.1 Binomial regression

In binomial models, covariates are modeled on a suitable transformation (the link function) of the binomial success probability, p. Let  $x_i$  denote some measured covariate for sample unit i and let  $p_i$  be the success probability for unit i. The standard choice is the "logit" link function which is:

$$log(p_i/(1-p_i)) = \alpha + \beta * x_i.$$

 $<sup>^{14}\</sup>mathrm{Some}$  of the jargon is actually a little bit confusing here because the binomial index is customarily referred to as "sample size" but in the context of N-mixture models N is actually the "population size"

The inverse-logit (or "expit") is

$$p_i = \operatorname{expit}(\alpha + \beta * x_i) = \frac{\exp(\alpha + \beta * x_i)}{1 + \exp(\alpha + \beta * x_i)}$$

There are many other possible link functions. However, ecologists seem to adopt the logit link function without question in most applications <sup>15</sup>. We sometimes use the "complementary log-log" (= "cloglog") link function in ecological applications because it arises naturally in many situations (Royle and Dorazio, 2008, p. 150). For example, consider the "probability of observing a count greater than 0" under a Poisson model:  $\Pr(y > 0) = 1 - \exp(-\lambda)$ . In that case,

$$cloglog(p) = log(-log(1-p)) = log(\lambda)$$

So that if you have covariates in your linear predictor for E(y) under a Poisson model then they are linear on the complementary log-log link of p. In models of species occurrence it seems natural to view occupancy as being derived from local abundance N (Royle and Nichols, 2003; Royle and Dorazio, 2006; Dorazio, 2007). Therefore, models of local abundance in which  $N \sim \text{Poisson}(A\lambda)$  for a habitat patch of area A implies a model for occupancy  $\psi$  of the form

$$cloglog(\psi) = log(A) + log(\lambda).$$

We will use the cloglog link in some analyses of SCR models in chapter 5 and elsewhere.

#### 2.11.2 Example: Waterfowl Banding Data

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It would be easy to consider a standard "distribution modeling" application where K=1 and the outcome is occurrence (y=1) or not (y=0) of some species. Such examples abound in books (e.g., Royle and Dorazio (2008, ch. 3); Kéry (2010, ch. 21); Kery and Schaub (2011, ch. 13)) and in the literature. Instead, we will consider an example involving band returns of waterfowl which were analyzed by Royle and Dubovsky  $(2001)^{16}$ .

For these data,  $y_i$  is the number of waterfowl bands recovered out of  $B_i$  birds banded at some location  $\mathbf{s}_i$ . In this case  $B_i$  is fixed. Thinking about recovery rate as being proportional to harvest rate, we use these data to explore geographic gradients in recovery rate resulting from variability in harvest pressure experienced by populations depending on their migration ecology. As such, we fit a basic binomial GLM with a linear response to geographic coordinates (including an interaction term). The data are provided with the  $\mathbf{R}$  package scrbook. Here we provide the part of the script for creating the model and fitting the model in **WinBUGS** using

<sup>&</sup>lt;sup>15</sup>a notable exception is distance sampling, which is all about choosing among link functions <sup>16</sup>I hate this example. Anyone got a better one thats not distribution modeling?

the bugs function. There are few structural differences between this model and the Poisson GLM fitted previously. The main things are due to the data structure (we have a matrix here instead of a vector) and otherwise we change the main distributional assumption to binomial (specified with dbin) and then use the logit function to relate the parameter  $p_{it}$  to the covariates. Here is the script:

```
load("mallarddata") # not sure how this will look
1379
1380
    sink("model.txt")
1381
    cat("
1382
    model {
1383
     for(t in 1:5){
         for (i in 1:nobs){
1385
            y[i,t] ~ dbin(p[i,t], B[i,t])
1386
            logit(p[i,t]) \leftarrow alpha0[t] + alpha1*X[i,1] + alpha2*X[i,2] + alpha3*X[i,1]*X[i,2]
1387
1388
    }
1389
     alpha1~dnorm(0,.001)
1390
     alpha2~dnorm(0,.001)
1391
    alpha3~dnorm(0,.001)
1392
    for(t in 1:5){
1393
       alpha0[t] ~ dnorm(0,.001)
1394
     }
1395
    }
     ",fill=TRUE)
1397
    sink()
1398
1399
          <- list(B=mallard.bandings, y=mallard.recoveries,</pre>
1400
                   nobs=nrow(banding.locs),X=banding.locs)
1401
     inits <- function(){</pre>
1402
           list(alpha0=rnorm(5),alpha1=0,alpha2=0,alpha3=0) }
    parms <- list('alpha0', 'alpha1', 'alpha2', 'alpha3')</pre>
1404
           <- bugs(data,inits, parms, "model.txt", n.chains=3,</pre>
1405
       n.iter=2000,n.burnin=1000, n.thin=2,debug=TRUE)
1406
        Posterior summaries of model parameters are as follows:
1407
    > print(out,digits=3)
1408
     Inference for Bugs model at "model.txt", fit using WinBUGS,
1409
     3 chains, each with 2000 iterations (first 1000 discarded), n.thin = 2
1410
     n.sims = 1500 iterations saved
1411
                                                                             97.5% Rhat n.eff
                                     2.5%
                                                 25%
                                                           50%
                                                                     75%
                    mean
                             sd
1412
                                                        -2.346
     alpha0[1]
                  -2.346 0.036
                                   -2.417
                                             -2.370
                                                                  -2.323
                                                                            -2.277 1.001
                                                                                            1500
1413
                                                        -2.356
1414
     alpha0[2]
                  -2.356 0.032
                                   -2.420
                                             -2.379
                                                                  -2.335
                                                                            -2.292 1.001
                                                                                            1500
     alpha0[3]
                  -2.220 0.035
                                   -2.291
                                             -2.244
                                                        -2.219
                                                                  -2.197
                                                                            -2.153 1.001
                                                                                            1500
1415
    alpha0[4]
                  -2.144 0.039
                                   -2.225
                                             -2.169
                                                        -2.143
                                                                  -2.116
                                                                            -2.068 1.000
1416
    alpha0[5]
                  -1.925 0.034
                                   -1.990
                                             -1.949
                                                       -1.924
                                                                  -1.901
                                                                            -1.856 1.004
                                                                                             570
```

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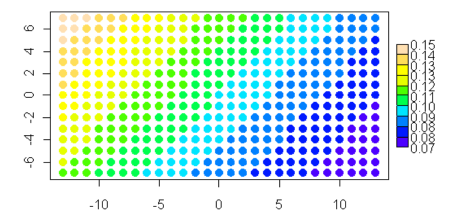


Figure 2.6. Predicted recovery rate of bands.

```
alpha1
                  -0.023 0.003
                                  -0.028
                                            -0.025
                                                      -0.023
                                                                -0.022
                                                                         -0.018 1.001
1418
     alpha2
                   0.020 0.006
                                   0.009
                                             0.016
                                                       0.020
                                                                0.024
                                                                          0.031 1.001
                                                                                        1500
1419
                                                                          0.002 1.001
     alpha3
                   0.000 0.001
                                  -0.002
                                            -0.001
                                                       0.000
                                                                 0.000
                                                                                        1500
               1716.001 4.091 1710.000 1713.000 1715.000 1718.000 1726.000 1.001
1421
     deviance
                                                                                        1500
     [... some output deleted ...]
1423
```

The basic result suggests a negative east-west gradient and a positive south to north gradient but no interaction. A map of the response surface is shown in Fig. 2.6. We did an additional MCMC run where we saved the binomial parameter p and computed the Bayesian p-value (double use of "p" here is confusing, but I guess that happens sometimes!) using a fit statistic based on the Freeman-Tukey statistic (see Section XXX above). The result indicates that the linear response surface model does not provide an adequate fit of the data. The reader should contemplate whether this invalidates the basic interpretation of the result.

# 2.12 SUMMARY AND OUTLOOK

GLMs and GLMMs are the most useful statistical methods in all of ecology. The principles and procedures underlying these methods are relevant to nearly all modeling and analysis problems in every branch of ecology. Moreover, understanding how to analyze these models is crucial in a huge number of diverse problems. If you

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understand and can conduct classical likelihood and Bayesian analysis of Poisson and binomial GLM(M)s, then you will be successful analyzing and understanding more complex classes of models that arise. We will see shortly that spatial capture-recapture models are a type of GLMM and thus having a basic understanding of the conceptual origins and formulation of GLM(M)s and their analysis is extremely useful.

We note that GLM(M)s are routinely analyzed by likelihood methods but we have focused on Bayesian analysis here in order to develop the tools that are less familiar to most ecologists. In particular, Bayesian analysis of models with random effects is relatively straightforward because the models are easy to analyze conditional on the random effect, using methods of MCMC. Thus, we will often analyze SCR models in later chapters by MCMC, explicitly adopting a Bayesian inference framework. In that regard, the various BUGS engines (WinBUGS, OpenBUGS, JAGS) are enormously useful because they provide an accessible platform for carrying out analyses by MCMC by just describing the model, and not having to worry about how to actually build MCMC algorithms. That said, the **BUGS** language is more important than just to the extent that it enables one to do MCMC - it is useful as a modeling tool because it fosters understanding, in the sense that it forces you to become intimate with your model. You have to write down all of the probability assumptions, the relationships between observations and latent variables and parameters. This is really a great learning paradigm that you can grow with.

While we have emphasized Bayesian analysis in this chapter, and make primary use of it through the book, we we will provide an introduction to likelihood analysis in chapter 9 and use those methods also from time to time. Before getting to that, however, it will be useful to talk about more basic, conventional closed population capture-recapture models and these are the topic of the next chapter.

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465	CLOSED POPULATION MODELS	

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# ESTIMATING THE SIZE OF A CLOSED POPULATION

In this chapter we will consider ordinary capture-recapture (CR) models for estimating population size in closed populations. We will see that such models are closely related to binomial (or logistic) regression type models. In fact, when Nis known, they are precisely such models. We consider some important extensions of ordinary closed population models that accommodate various types of "individual effects" — either in the form of explicit covariates (sex, age, body mass) or unstructured "heterogeneity" in the form of an individual random effect. In general, these models are variations of generalized linear or generalized linear mixed models (GLMMs). Because of the paramount importance of this concept, we focus mainly on fairly simple models in which the observations are individual encounter frequencies,  $y_i$  = the number of encounters of individual i out of K replicate samples of the population which, for the models we consider here, is the outcome of a binomial random variable. Along the way, we consider the spatial context of capture-recapture data and models and demonstrate that density cannot be formally estimated when spatial information is ignored. We also review some of the informal methods of estimating density using CR methods, and consider some of their limitations. We will be exposed to our first primitive spatial capture-recapture models which arise as relatively minor variations of so-called "individual covariate models" (of the Huggins (1989) and Alho (1990) variety). In a sense, the point of this chapter is to establish that linkage in a direct and concise manner beginning with the basic "Model M0" and extensions of that model to include individual heterogeneity and also individual covariates. A special type of individual covariate models is distance sampling, which could be thought of as the most primitive spatial capture-recapture model. In later chapters we further develop and extend

ideas introduced in this chapter.

We emphasize Bayesian analysis of capture-recapture models and we accomplish this using a method related to classical "data augmentation" from the statistics literature Tanner and Wong (e.g., 1987)). This is a general concept in statistics but, in the context of capture-recapture models where N is unknown, it has a consistent implementation across classes of capture-recapture models and one that is really convenient from the standpoint of doing MCMC (Royle et al., 2007). We use data augmentation throughout this book and thus emphasize its conceptual and technical origins and demonstrate applications to closed population models. We refer the reader to Kery and Schaub (2011, ch. 6) for an accessible and complimentary development of ordinary closed population models.

#### 4.1 THE SIMPLEST CLOSED POPULATION MODEL: MODEL MO

We suppose that there exists a population of N individuals which we subject to repeated sampling, say over K nights, where individuals are captured, marked, and subsequently recaptured. We suppose that individual encounter histories are obtained, and these are of the form of a sequence of 0's and 1's indicating capture (y = 1) or not (y = 0) during any sampling occasion ("sample"). As an example, suppose K = 5 sampling occasions, then an individual captured during sample 2 and 3 but not otherwise would have an encounter history of the form  $\mathbf{y} = (0, 1, 1, 0, 0)$ . Thus, the observation  $\mathbf{y}_i$  for each individual (i) is a vector having elements denoted by  $y_{ik}$  for k = 1, 2, ..., K. Usually this is organized as a row of a matrix with elements  $y_{ik}$ , see Table 4.1. Except where noted explicitly, we suppose that observations are independent within individuals and among individuals. Formally, this allows us to say that  $y_{ik}$  are Bernoulli random variables and we may write  $y_{ik} \sim \text{Bern}(p)$ . Consequently, for this very simple model in which p is in fact constant, then we can declare that the individual encounter frequencies (total captures),  $y_i = \sum_k y_{ik}$ , have a binomial distribution based on a sample of size K. That is

$$y_i = \sum_k y_{ik} \sim \text{Bin}(p, K)$$

for every individual in the population. This is a remarkably simple model that forms the cornerstone of almost all of classical capture-recapture models, including most spatial capture-recapture models discussed throughout this book. Evidently, the basic capture-recapture model structure is precisely a simplistic version of a logistic-regression model with only an intercept term (logit(p) = constant). To say that all capture-recapture models are just logistic regressions is only slightly inaccurate. In fact, we are proceeding here "conditional on N", i.e., as if we knew N. In practice we don't, of course, and that is kind of the point of capture-recapture models as estimating N is the central objective. But, by proceeding conditional on N, we can specify a simple model and then deal with the fact that N is unknown

**Table 4.1.** a capture-recapture data set with n=6 observed individuals and K=5 samples.

	Sample occasion						
indiv $i$	1	2	3	4	5	$y_i$	
1	1	0	0	1	0	2	
2	0	1	0	0	1	2	
3	1	0	0	1	0	2	
4	1	0	1	0	1	3	
5	0	1	0	0	0	1	
n = 6	1	0	0	0	0	1	

using standard methods that you are already familiar with (i.e., GLMs - see chapter 2).

Assuming individuals of the population are observed independently, the joint probability distribution of the observations is the product of N binomials

$$Pr(y_1, \dots, y_N | p) = \prod_{i=1}^N Bin(y_i | K, p)$$
$$= \prod_{k=0}^K \pi(k)^{n_k}$$

where  $\pi(k) = \text{Bin}(k|K,p)$  and where  $n_k = \sum_{i=1}^N I(y_i = k)$  denotes the number of individuals captured k times in K surveys. We emphasize that this is conditional on N, in which case we get to observe the y = 0 observations and the resulting data are just iid binomial counts. Because this is a binomial regression model of the variety described in chapter 2, fitting this model using a BUGS engine poses no difficulty.

The essential problem in capture-recapture, however, is that N is not known because the number of uncaptured/missing individuals (i.e., those in the zero cell that occur with probability  $\pi(0)$ ) is unknown. Consequently, the observed capture frequencies  $n_k$  are no longer independent. Instead, their joint distribution is multinomial (e.g., see Illian (2008a, p. xyz)):

$$n_1, n_2, \dots, n_K \sim \text{Multin}(N, \pi(1), \pi(2), \dots, \pi(K))$$
 (4.1.1)

Note that in our notation the number of uncaptured/missing individuals is denoted by  $n_0 = N - n$ , where  $n = \sum_{k=1}^{K} n_k$  denotes the total number of distinct individuals seen in the K samples.

To fit the model in which N is unknown, we can regard N as a parameter and maximize the multinomial likelihood directly. While direct likelihood analysis of the multinomial model is straightforward, that does not prove to be too useful in practice because we seldom are concerned with models for the aggregated encounter history frequencies. In many instances, including for spatial capture-recapture (SCR)

models, we require a formulation of the model that can accommodate individual level covariates which we address subsequently in this chapter.

#### 4.1.1 The Spatial Context of Capture-Recapture

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A common assumption made is that of population "closure" which is really just a colloquial way of saying (in part) the Bernoulli assumptions stated explicitly above. In the biological context, closure means, strictly, no additions or subtractions from the population during study. This is manifest by the statement that the encounters are independent and identically distributed (iid) Bernoulli trials. In practice, closure is usually interpreted by the manner in which potential violations of that assumption arise. In particular, two important elements of the closure assumption are "demographic" and "geographic" closure. If an individual dies then subsequent values of  $y_{ik}$  are clearly no longer Bernoulli trials with the same parameter p. If there is no mortality or recruitment in the population, then we say that demographic closure is satisfied. Similarly, animals may emigrate or immigrate. If they do not, then geographic closure is satisfied. Sometimes a distinction is made between temporary and permanent emigration or immigration. That is a relevant distinction in spatial capture-recapture models, because SCR models explicitly accommodate "temporary emigration" of a certain type, due to individuals moving about their home range. The demographic closure assumption can also be relaxed using SCR models, but we will save that discussion for chapter 5.

# 4.1.2 Conditional likelihood

We saw that a basic closed population model is a simple logistic regression model if N is known and, when N is unknown, the model is multinomial with index or sample size parameter N. This multinomial model, being conditional on N, is sometimes referred to as the "joint likelihood" the "full likelihood" or the "unconditional likelihood" (or model in place of likelihood). This formulation differs from the so-called "conditional likelihood" approach in which the likelihood of the observed encounter histories is devised conditional on the event that an individual is captured at least once. To construct this likelihood, we have to recognize that individuals appear or not in the sample based on the value of the random variable  $y_i$ , that is, we capture them if and only if  $y_i > 0$ . The observation model is therefore based on Pr(y|y>0). For the simple case of Model M0, the resulting conditional distribution is a "zero truncated" binomial distribution which accounts for the fact that we cannot observe the value y=0 in the data set (see Royle and Dorazio. 2008, section XYZ). Both the conditional or unconditional models are legitimate modes of analysis in all capture-recapture types of studies, and they provide equally valid descriptions of the data and for many practical purposes provide equivalent inferences, at least in large sample sizes (Sanathanan, 1972).

Mode of analysis	parameters in model	statistical model
Joint likelihood	p, N	multinomial with index $N$
Conditional likelihood	p	zero-truncated binomial
Data augmentation	$p,\ \psi$	zero-inflated binomial

Table 4.2. Modes of analysis of capture-recapture models.

In this book we emphasize Bayesian analysis of capture-recapture models using data augmentation (discussed subsequently), which produces yet a third distinct formulation of capture recapture-models based on the zero-inflated binomial distribution that we describe in the next section. Thus, there are 3 distinct formulations of the model – or models of analysis – for analyzing all capture-recapture models based on the (1) binomial model for the joint or unconditional specification; (2) zero-truncated binomial that arises "conditional on n"; and (3) the zero-inflated binomial that arises under data augmentation. Each formulation has a distinct complement of model parameters (shown in Table 4.2 for Model M0).

#### 4.2 DATA AUGMENTATION

We consider a method of analyzing closed population models using data augmentation (DA) which is useful for Bayesian analysis and, in particular, analysis of models using the various BUGS engines and other software. Data augmentation is a general statistical concept that is widely used in statistics in many different settings. The classical reference is Tanner and Wong (1987) but see also Liu and Wu (1999). Data augmentation can be adapted to provide a very generic framework for Bayesian analysis of capture-recapture models with unknown N. This idea was introduced for closed populations by Royle et al. (2007), and has subsequently been applied to a number of different contexts including individual covariate models (Royle, 2009), open population models (Royle and Dorazio, 2008, 2010; Gardner et al., 2010), spatial capture-recapture models (Royle and Young, 2008; Royle, 2010; Gardner, 2009), and many others.

Conceptually, data augmentation takes the data you wish you had - that is, the data set with N rows - the known-N data set - and embeds that data set into a larger data set having M>N rows. <sup>1</sup> It is always possible, in practice, to choose M pretty easily for a given problem and context. Then, under data augmentation, analysis is focused on the "augmented data set." That is, we analyze the bigger data set - the one having M rows - with an appropriate model that accounts for the augmentation. Inference is focused directly on estimating the proportion  $\psi=E[N]/M$ , instead of directly on N, where  $\psi$  is the "data augmentation parameter."

<sup>&</sup>lt;sup>1</sup> RC: Might be just me, but I find that formulation a little confusing... I think it's the 'data you wish you had because that's effectively data you don't have. I think it might be easier to grasp if this were explained with the data you do have - based on n.

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#### 4.2.1 DA links occupancy models and closed population models

We provide a heuristic description of data augmentation based on the close correspondence between so-called "occupancy" models and closed population models following Royle and Dorazio (2008, sec. xyz).

In occupancy models (MacKenzie et al., 2002; Tyre et al., 2003) the sampling situation is that M sites, or patches, are sampled multiple times to assess whether a species occurs at each site. This yields encounter data such as that illustrated in the left panel of Table 4.3. The important problem is that a species may occur at a site, but go undetected, yielding the "all-zero" encounter histories which are observed. However, some of the all-zeros may well correspond to sites where the species in fact does not occur. Thus, while the zeros are observed, there are too many of them and, in a sense, the inference problem is to allocate the zeros into "structural" (fixed) and "sampling" (or stochastic) zeros. More formally, inference is focused on the parameter  $\psi$ , the probability that a site is occupied. In contrast, in classical closed population studies, we observe a data set as in the middle panel of Table 4.3 where no zeros are observed. The inference problem is, essentially, to estimate how many sampling zeros there are - or should be - in a "complete" data set. The inference objective (how many sampling zeros?) is precisely the same for both types of problems if an upper limit M is specified for the closed population model. The only distinction being that, in occupancy models, M is set by design (i.e., the number of sites to visit) whereas a natural choice of M for capture-recapture models may not be obvious. However, by assuming a uniform prior for N on the integers [0, M], this upper bound is induced (Royle et al., 2007). Then, one can analyze capture-recapture models by adding M-n all-zero encounter histories to the data set and regarding the augmented data set, essentially, as a site-occupancy data set.

Thus, the heuristic motivation of data augmentation is to fix the size of the data set by adding too many all-zero encounter histories to create the data set shown in the right panel of Table 4.3 - and then analyze the augmented data set using an occupancy type model which includes both "unoccupied sites" as well as "occupied sites" at which detections did not occur. We call these M-n all-zero histories "potential individuals" because they exist to be recruited (in a non-biological sense) into the population, for example during an analysis by MCMC.

To analyze the augmented data set, we recognize that it is a zero-inflated version of the known-N data set. That is, some of the augmented all-zeros are sampling zeros (corresponding to actual individuals that were missed) and some are "structural" zeros, which do not correspond to individuals in the population. For a basic closed-population model, the resulting likelihood under data augmentation - that is, for the data set of size M – is a simple zero-inflated binomial likelihood. The zero-inflated binomial model can be described "hierarchically", by introducing a set of binary latent variables,  $z_1, z_2, \ldots, z_M$ , to indicate whether each individual i is  $(z_i = 1)$  or is not  $(z_i = 0)$  a member of the population of N individuals exposed

to sampling. We assume that  $z_i \sim \mathrm{Bern}(\psi)$  where  $\psi$  is the probability that an individual in the data set of size M is a member of the sampled population - in the sense that  $1-\psi$  is the probability of realizing a "structural zero" in the augmented data set. The zero-inflated binomial model which arises under data augmentation can be formally expressed bythe following set of assumptions:

$$\begin{aligned} y_i|z_i &= 1 \sim \operatorname{Bin}(K,p) \\ y_i|z_i &= 0 \sim \delta(0) \\ z_i &\stackrel{iid}{\sim} \operatorname{Bern}(\psi) \\ \psi &\sim \operatorname{Unif}(0,1) \\ p &\sim \operatorname{Unif}(0,1) \end{aligned}$$

for i = 1, ..., M, where  $\delta(0)$  is a point mass at y = 0.

We note that N is no longer an explicit parameter of this model. Instead, we estimate  $\psi$  and functions of the latent variables. In particular, under the assumptions of the zero-inflated model,  $z_i \stackrel{iid}{\sim} \mathrm{Bern}(\psi)$ ; therefore, N is a function of these latent variables:

$$N = \sum_{i=1}^{M} z_i.$$

Further, we note that the latent  $z_i$  parameters can be removed from the model by integration, in which case the joint probability of the data is

$$\Pr(y_1, \dots, y_M | p, \psi) = \prod_{i=1}^M \psi \text{Bin}(y_i | K, p) + I(y_i = 0)(1 - \psi)$$
 (4.2.1)

Which can be maximized directly to obtain the MLEs of the structural parameters  $\psi$  and p or those of other more complex models (e.g., see Royle, 2006). We could estimate these parameters and then use them to obtain an estimator of N using the so-called "Best unbiased predictor" (see Royle and Dorazio, 2011).

#### 4.2.2 Model $M_0$ in BUGS

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For model  $M_0$  in which we can aggregate the encounter data to individual-specific encounter frequencies, the augmented data are given by the vector of frequencies  $(y_1, \ldots, y_n, 0, 0, \ldots, 0)$ . The zero-inflated model of the augmented data combines the model of the latent variables,  $z_i \sim \text{Bern}(\psi)$  with the conditional-on-z binomial model:

$$y_i|z_i = 0 \sim \delta(0)$$
  
$$y_i|z_i = 1 \sim \text{Bin}(K, p)$$

**Table 4.3.** Hypothetical occupancy data set (left), capture-recapture data in standard form (center), and capture-recapture data augmented with all-zero capture histories (right).

Occupancy data		Ca	apture-	recapt	ure	Augmented C-R			-R		
site	k=1	k=2	k=3	ind	k=1	k=2	k=3	ind	k=1	k=2	k=3
1	0	1	0	1	0	1	0	1	0	1	0
2	1	0	1	2	1	0	1	2	1	0	1
3	0	1	0		0	1	0	3	1	0	1
4	1	0	1		1	0	1	4	1	0	1
5	0	1	1		0	1	1	5	1	0	1
	0	1	1		0	1	1		0	1	1
	1	1	1		1	1	1		0	1	1
	1	1	1		1	1	1		1	1	1
	1	1	1		1	1	1		1	1	1
$\mathbf{n}$	1	1	1	n	1	1	1	n	1	1	1
	0	0	0						0	0	0
	0	0	0						0	0	0
	0	0	0						0	0	0
	0	0	0						0	0	0
	0	0	0						0	0	0
	0	0	0					N	0	0	0
	0	0	0						0	0	0
	0	0	0						0	0	0
M	0	0	0						0	0	0
								Μ	0	0	0
								•			

It is convenient to express the conditional-on-z observation model concisely as:

$$y_i|z_i \sim \text{Bin}(K, pz_i)$$

Thus, if  $z_i = 0$  then the success probability of the binomial distribution is identically 0 whereas, if  $z_i = 1$ , then the success probability is p. This is useful in describing the model in the **BUGS** language, as shown below. Note the last line of the model specification here provides the expression for computing N from the data augmentation variables  $z_i$ .

```
p ~ dunif(0,1)
1691
1692
     psi~dunif(0,1)
1693
     # nind = number of individuals captured at least once
1694
         nz = number of uncaptured individuals added for PX-DA
1695
     for(i in 1:(nind+nz)) {
1696
         z[i]~dbern(psi)
1697
        mu[i] < -z[i] *p
1698
         y[i]~dbin(mu[i],K)
1699
1700
1701
     N<-sum(z[1:(nind+nz)])</pre>
1702
```

Specification of a more general model in terms of the individual encounter observations  $y_{ik}$  is not much more difficult than for the individual encounter frequencies. We define the observation model by a double loop and change the indexing of things accordingly, i.e.,

```
1707 for(i in 1:(nind+nz)) {
1708         z[i]~dbern(psi)
1709         for(k in 1:K){
1710             mu[i,k]<-z[i]*p
1711             y[i,k]~dbin(mu[i,k],1)
1712         }
1713     }</pre>
```

1703

1704

1706

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In this manner, it is straightforward to incorporate covariates on p (see discussion of this below and also chapt. 8 (REF XYZ) and consider other extensions.

# 4.2.3 Formal development of data augmentation

Use of DA for solving inference problems with unknown N can be justified as originating from the choice of uniform prior on N. The Unif(0, M) prior for N is innocuous in the sense that the posterior associated with this prior is equal to the likelihood for sufficiently large M. One way of inducing the Unif(0, M) prior on N

is by assuming the following hierarchical prior:

$$N \sim \text{Bin}(M, \psi)$$

$$\psi \sim \text{Unif}(0, 1)$$
(4.2.2)

which includes a new model parameter  $\psi$ . This parameter denotes the probability that an individual in the super-population of size M is a member of the population of N individuals exposed to sampling. The model assumptions, specifically the multinomial model (eq. XYZ) and eq. 4.2.2, may be combined to yield a reparameterization of the conventional model that is appropriate for the augmented data set of known size M:

$$(n_1, n_2, \dots, n_K) \sim \text{Multin}(M, \psi \pi(1), \psi \pi(2), \dots, \psi \pi(K))$$
 (4.2.3)

This arises by removing N from Eq. multinomial XYZ by integrating over the binomial prior distribution for N. Thus, the models we analyze under data augmentation arise formally by removing the parameter N from the ordinary model the model conditional on N - by integrating over a binomial prior distribution for N.

Note that the M-n unobserved individuals in the augmented data set have probability  $\psi\pi(0)+(1-\psi)$ , indicating that these unobserved individuals are a mixture of individuals that are sampling zeros ( $\psi\pi_0$ , and belong to the population of size N) and others that are "structural zeros" (occurring in the augmented data set with probability  $1-\psi$ ). In Eq. 4.2.3 N has been eliminated as a formal parameter of the model by marginalization (integration) and replaced with the new parameter  $\psi$ , which we will call the "data augmentation parameter." However, the full likelihood containing both N and  $\psi$  can be analyzed (see Royle et al., 2007).

# 4.2.4 Remarks on Data Augmentation

Data augmentation may seem like a strange and mysterious black-box, and likely it is unfamiliar to most people even those with extensive experience with capture-recapture models. However, it really is a formal reparameterization of capture-recapture models in which N is removed from the ordinary (conditional-on-N) model by integration. In the case of Model M0, data augmentation produces the zero-inflated binomial which is distinct from the original observation model, but only in the sense that it embodies, explicitly, the  $\mathrm{Unif}(0,M)$  prior for N. Choice of M might be cause for some concern related to potential sensitivity to choice of M. The guiding principle is that it should be chosen large enough so that the posterior for N is not truncated, but no larger because large values entail more computational burden. It seems likely that the properties of the Markov chains should be affected by M and so some optimality might exist (Gopalaswamy, 2012), as in occupancy models (Mackenzie and Royle, 2005). Formal analysis of this is required.

We emphasize the motivation for data augmentation being that it produces a data set of fixed size, so that the parameter dimension in any capture-recapture model is also fixed. As a result, MCMC is a relatively simple proposition using standard Gibbs Sampling. Consider the simplest context - analyzing Model M0 using the occupancy model. In this case, DA converts Model M0 to a basic occupancy model and the parameters p and  $\psi$  have known full-conditional distributions (in fact, beta distributions) that can be sampled from directly. Furthermore, the data augmentation variables - the latent data augmentation variables z, can be sampled from Bernoulli full conditionals. MCMC is not too much more difficult for complicated models - sometimes the hyperparameters need to be sampled using a Metropolis-Hastings step, but nothing more sophisticated than that is required.

There are other approaches to analyzing models with unknown N, using reversible jump MCMC (RJMCMC) or other so-called "trans-dimensional" (TD) algorithms<sup>2</sup> (Durbin and Elston, 2012; King, missing; Schofield and Barker, missing). What distinguishes DA from RJMCMC and related TD methods is that DA is used to create a distinctly new model that is unconditional on N and we (usually) analyze the unconditional model. The various TD/RJMCMC approaches seek to analyze the conditional-on-N model in which the dimensional of the parameter space is a variable function of N. TD/RJMCMC approaches might appear to have the advantage that one can model N explicitly or consider alternative priors for N. However, despite that N is removed as an explicit parameter in DA, it is possible to develop hierarchical models that involve structure on N (Converse and Royle, 2010; Royle et al., 2011a) which we consider in chapt. XYZ.

#### 4.2.5 Example: Black Bear Study on Fort Drum

To illustrate the analysis of Model M0 using data augmentation, we use a data set collected at Fort Drum Military Installation in upstate New York by the Department of Defense, Cornell University and colleagues. These data have been analyzed in various forms by Gardner (2009); Gardner et al. (2010), and Wegan (missing). The specific data used here are encounter histories on 47 individuals obtained from an array of 38 baited "hair snares" (Fig. 4.1) during June and July 2006. Barbed wire traps were baited and checked for hair samples each week for eight weeks, thus we have K=8 sample intervals. The data are provided on the Web Supplement and the analysis can be set up and run as follows. Here, the data were augmented with M-n=128 (M=175) all-zero encounter histories.

```
## Consider adding comments to your code.
## Good idea. This will be done in final draft
trapmat<-read.csv("FDtrapmat.csv")
bearArray<-source("FDbeararray.R")$value
nind<-dim(bearArray)[1]
```

<sup>&</sup>lt;sup>2</sup>Look these citeations up in Royle-Dorazio EURING paper

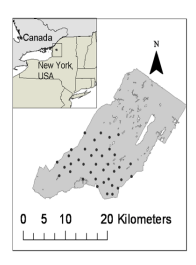


Figure 4.1. Fort Drum study area and hair snare locations.

```
K<-dim(bearArray)[3]</pre>
1794
     ntraps<-dim(bearArray)[2]
1795
1796
    M = 175
1797
    nz<-M-nind
1798
     Xaug <- array(0, dim=c(M,ntraps,K))</pre>
1800
     Xaug[1:nind,,]<-bearArray</pre>
1801
    y<- apply(Xaug,c(1,3),sum)
1802
    y[y>1]<-1
1803
    ytot<-apply(y,1,sum)</pre>
                               # total encounters out of K
1804
        Note that the raw data, y, is an M \times K array of individual encounter events
1805
     (i.e., y_{ik} = 1 if individual i was encountered in any trap and 0 otherwise). For
1806
     i = 48, ..., 175, y_{ik} = 0 as these are augmented observations. For Model M0 it is
1807
    sufficient to reduce the data to individual encounter frequencies which we have
1808
     labeled ytot above. The BUGS model file along with commands to fit the model
1809
     are as follows:
1810
     set.seed(2013)
                                      # to obtain the same results each time
1811
     data0<-list(y=y,M=M,K=K)
1812
     params0<-list('psi','p','N')</pre>
     zst=c(rep(1,nind),rbinom(M-nind, 1, .5))
1814
     inits = function() {list(z=zst, psi=runif(1), p=runif(1)) }
1815
1816
    cat("
1817
```

```
model {
1818
1819
    psi~dunif(0, 1)
1820
    p~dunif(0,1)
1821
1822
    for (i in 1:M){
1823
        z[i]~dbern(psi)
1824
        for(k in 1:K){
1825
           tmp[i,k] < -p*z[i]
1826
           y[i,k]~dbin(tmp[i,k],1)
1827
            }
1828
            }
1829
    N<-sum(z[1:M])
1830
    }
1831
     ",file="modelMO.txt")
1832
1833
    fit0 = bugs(data0, inits, params0, model.file="modelM0.txt",
1834
            n.chains=3, n.iter=2000, n.burnin=1000, n.thin=1,
1835
            debug=TRUE,working.directory=getwd())
1836
        The posterior summary statistics from this analysis are as follows:
1837
    > print(fit0,digits=2)
1838
    Inference for Bugs model at "modelMO.txt", fit using WinBUGS,
1839
      3 chains, each with 2000 iterations (first 1000 discarded)
1840
      n.sims = 3000 iterations saved
1841
                                2.5%
                                                 50%
                                                               97.5% Rhat n.eff
                 mean
                          sd
                                         25%
                                                         75%
1842
                 0.29
                        0.04
                                0.22
                                        0.26
                                                0.29
                                                        0.31
                                                                0.36
                                                                         1
                                                                            3000
1843
    psi
                 0.30
                        0.03
                                0.25
                                        0.28
                                                0.30
                                                        0.32
                                                                0.35
                                                                         1
                                                                            3000
1844
    р
                49.94
                       1.99
                              47.00
                                      48.00
                                              50.00
                                                      51.00
                                                               54.00
                                                                            3000
    N
1845
    deviance 489.05 11.28 471.00 480.45 488.80 495.40 513.70
                                                                            3000
1846
1847
    [.. some output deleted ...]
1848
```

**WinBUGS** did well in choosing an MCMC algorithm for this model – we have  $\hat{R}=1$  for each parameter, and an effective sample size of 3000, equal to the total number of posterior samples. We see that the posterior mean of N under this model is 49.94 and a 95% posterior interval is (48,54). We revisit these data later in the context of more complex models.

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In order to obtain an estimate of density, D, we need an area to associate with the estimate of N, and commonly used procedures to conjure up such an area include buffering the trap array by the home range radius, often estimated by the mean maximum distance moved (MMDM)<sup>3</sup>, 1/2 MMDM (Dice, 1938) or directly from telemetry data (REF XXX NEED REF HERE XXXXXX). Typically, the trap array is defined by the convex hull around the trap locations, and this is what we

<sup>&</sup>lt;sup>3</sup>really MMDM? How can this be an estimate of the home range radius? Reference for this?

applied a buffer to. We computed the buffer by using an estimate of the mean female 1860 home range radius (2.19 km) estimated from telemetry studies (Bales et al., 2005) instead of using an estimate based on our relatively more sparse recapture data<sup>4</sup>. 1862 For the Fort Drum study, the convex hull has area 157.135  $km^2$ , and the buffered convex hull has area  $277.011 \ km^2$ . To create this we used functions contained in the 1864 R package rgeos and created a utility function behave which is in our R package 1865 scrbook. The commands are as follows: 1866

```
library("rgeos")
1867
1868
    bcharea<-function(buff,traplocs){
1869
    p1<-Polygon(rbind(traplocs,traplocs[1,]))</pre>
1870
    p2<-Polygons(list(p1=p1), ID=1)
1871
    p3<-SpatialPolygons(list(p2=p2))
1872
    p1ch<-gConvexHull(p3)
1873
      bp1<-(gBuffer(p1ch, width=buff))</pre>
1874
     plot(bp1, col='gray')
1875
     plot(p1ch, border='black', lwd=2, add=TRUE)
1876
     gArea(bp1)
1877
1878
1879
    bcharea(2.19, traplocs=trapmat)
    The resulting buffered convex hull is shown in Fig. 4.2.
1881
        To conjure up a density estimate under model M_0, we compute the appropriate
1882
    posterior summary of N and the prescribed area (277.011 km^2):
1883
    > summary(fit0$sims.list$N/277.011)
1884
        Min. 1st Qu.
                        Median
                                    Mean 3rd Qu.
                                                       Max.
1885
      0.1697 0.1733 0.1805
                                 0.1803 0.1841
                                                     0.2130
1886
1887
      quantile(fit0$sims.list$N/277.011,c(0.025,0.975))
1888
          2.5%
                     97.5%
1889
    0.1696684 0.1949381
1890
    which yields a density estimate of about 0.18 ind/km<sup>2</sup>, and a 95% Bayesian confi-
```

dence interval of (0.170, 0.195). The obvious limitation of this estimate and, indeed, of the whole process, is that our choice of "area" is completely subjective - which area should we use? MMDM? One-half MMDM? Estimated from telemetry data? And, furthermore, how certain

are we of this area? Can we quantify our uncertainty about this quantity? More important, what exactly is the meaning of this area and, in this context, how do

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<sup>1897</sup> we gauge bias and/or variance of "estimators" of it? (i.e., what is it estimating?).

<sup>&</sup>lt;sup>4</sup>BETH: Why?

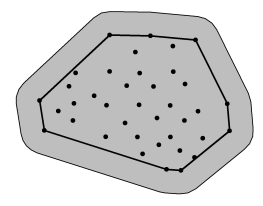


Figure 4.2. buffered convex hull of the bear hair snare array

#### 4.3 TEMPORALLY VARYING AND BEHAVIORAL EFFECTS

The purpose of this chapter is mainly to emphasize the central importance of the binomial model in capture-recapture and so we have considered models for individual encounter frequencies - the number of times individuals are captured out of K samples. Sometimes it is not acceptable to aggregate the encounter data for each individual – such as when encounter probability varies over time among samples. A type of time-varying response that seems relevant in most capture-recapture studies is "effort" such as amount of search time, number of observers, or trap effort or when p depends on date (Kéry et al., 2010; Gardner et al., 2010). A common situation is that in which there exists a "behavioral response" to trapping (even if the animal is not physically trapped).

Behavioral response is an important concept in carnivore studies because individuals might learn to come to baited traps or avoid traps due to trauma related to being encountered. There are a number of ways to parameterize a behavioral response to encounter. The distinction between persistent and ephemeral was made by Yang and Chao (2005) who considered a general behavioral response model of the form:

$$logit(p_{ik}) = \alpha_0 + \alpha_1 * y_{i,k-1} + \alpha_2 x_{ik}$$

where  $x_{ik}$  is a covariate indicator variable of previous capture (i.e.,  $x_{ik} = 1$  if

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captured in any previous period). Therefore, encounter probability changes depending on whether an individual was captured in the immediate previous period (ephemeral behavioral response) or in any previous period (persistent behavioral response). The former probably models a behavioral response due to individuals moving around their territory relatively slowly over time and the latter probably accommodates trap happiness due to baiting or shyness due to trauma. In spatial capture-recapture models it makes sense to consider a local behavioral response that is trap-specific (Royle et al., 2011c) - that is, the encounter probability is modified for individual traps depending on previous capture in specific traps.

Models with temporal effects are easy to describe in the **BUGS** language and analyze and we provide a number of examples in chapt. 8. XXXXX ?? XXXXXX

# 4.4 MODELS WITH INDIVIDUAL HETEROGENEITY

Here we consider models with individual-specific encounter probability parameters, say  $p_i$ , which we model according to some probability distribution,  $q(\theta)$ . We denote this basic model assumption as  $p_i \sim g(\theta)$ . This type of model is similar in concept to extending a GLM to a GLMM but in the capture-recapture context N is unknown. The basic class of models is often referred to as "Model  $M_h$ " but really this is a broad class of models, each being distinguished by the specific distribution assumed for  $p_i$ . There are many different varieties of Model  $M_h$  including parametric and various putatively non-parametric approaches (Burnham and Overton, 1978; Norris III and Pollock, 1996; Pledger, 2000). One important practical matter is that estimates of N can be extremely sensitive to the choice of heterogeneity model (Fienberg et al., 1999; Dorazio and Royle, 2003; Link, 2003). Indeed, Link (2003) showed that in some cases it's possible to find models that yield precisely the same expected data, yet produce wildly different estimates of N. In that sense, N for most practical purposes is not identifiable across classes of mixture models, and this should be understood before fitting any such model. One solution to this problem is to seek to model explicit factors that contribute to heterogeneity, e.g., using individual covariate models (See 4.5 below). Indeed, spatial capturerecapture models seek to do just that, by modeling heterogeneity due to the spatial organization of individuals in relation to traps or other encounter mechanism. For additional background and applications of Model  $M_h$  see Royle and Dorazio (2008, chapt. 6) and Kery and Schaub (2011, chapt. 6).

Model  $M_h$  has important historical relevance to spatial capture-recapture situations (Karanth, 1995) because investigators recognized that the juxtaposition of individuals with the array of trap locations should yield heterogeneity in encounter probability, and thus it became common to use some version of Model  $M_h$  in spatial trapping arrays to estimate N. While this doesn't resolve the problem of not knowing the area relevant to N, it does yield an estimator that accommodates the heterogeneity in p induced by the spatial aspect of capture-recapture studies.

To see how this juxtaposition induces heterogeneity, we have to understand

the relevance of movement in capture-recapture models. Imagine a quadrat that can be uniformly searched by a crew of biologists for some species of reptile (see Royle and Young (2008)). Figure 4.3 shows a sample quadrat searched repeatedly over a period of time. Further, suppose that species exhibits some sense of spatial fidelity in the form of a home range or territory, and individuals move about their home range (home range centroids are given by the blue dots) in some kind of random fashion. It is natural to think about it in terms of a movement process and sometimes that movement process can be modeled explicitly using hierarchical models (Royle and Young, 2008; Royle et al., 2011b). Heuristically, we imagine that each individual in the vicinity of the study area is liable to experience variable exposure to encounter due to the overlap of its home range with the sampled area - essentially the long-run proportion of times the individual is within the sample plot boundaries, say  $\phi$ . We might model the exposure of an individual to capture by supposing that  $z_i = 1$  if individual i is available to be captured (i.e., within the survey plot) during any sample, and 0 otherwise. Then,  $\Pr(z_i = 1) = \phi$ . In the context of spatial studies, it is natural that  $\phi$  should depend on where an individual lives, i.e., it should be individual-specific  $\phi_i$  (Chandler et al., 2011). This system describes, precisely, that of "random temporary emigration" (Kendall, 1997) where  $\phi_i$  is the individual-specific probability of being "available" for capture.

Conceptually, SCR models aim to deal with this problem of variable exposure to sampling due to movement in the proximity of the trapping array explicitly and formally with auxiliary spatial information. If individuals are detected with probability  $p_0$ , conditional on  $z_i = 1$ , then the marginal probability of detection of individual i is

$$p_i = p_0 \phi_i$$

so we see clearly that individual heterogeneity in encounter probability is induced as a result of the juxtaposition of individuals (i.e., their home ranges) with the sample apparatus and the movement of individuals about their home range.

We will work with a specific type of Model  $M_h$  here, that in which we extend the basic binomial observation model of Model  $M_0$  so that

$$logit(p_i) = \mu + \eta_i$$

where 1985

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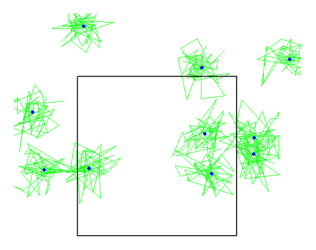
1988

$$\eta_i \sim \text{Normal}(0, \sigma_p^2)$$

We could as well write 1986

$$logit(p_i) \sim Normal(\mu, \sigma_p^2)$$

This "logit-normal mixture" was analyzed by Coull and Agresti (1999) and else-1987 where. It is a natural extension of the basic model with constant p, as a mixed GLMM, and similar models occur throughout statistics. It is also natural to con-1989 sider a beta prior distribution for  $p_i$  (Dorazio and Royle, 2003) and so-called "finite-1990 mixture" models are also popular (Norris III and Pollock, 1996; Pledger, 2000). 1991



**Figure 4.3.** A quadrat searched for lizards and the locations of each lizard over some period of time.

#### 4.4.1 Analysis of Model Mh

1992

2005

2006

2007

If N is known, it is worth taking note of the essential simplicity of Model  $M_h$  as 1993 a binomial GLMM. This is a type of model that is widely applied in just about 1994 every scientific discipline and using standard methods of inference based either on 1995 integrated likelihood (Laird and Ware, 1982; Berger et al., 1999) which we discuss in chapt. 9 or standard Bayesian methods. However, because N is not known, 1997 inference is somewhat more challenging. We address that here using Bayesian anal-1998 ysis based on data augmentation (DA). Although we use data augmentation in the 1999 context of Bayesian methods here, we note that heterogeneity models formulated 2000 under DA are easily analyzed by conventional likelihood methods as zero-inflated 2001 binomial mixtures (Royle, 2006) and more traditional analysis of model  $M_h$  based 2002 on integrated likelihood, without using data augmentation, has been considered by 2003 Coull and Agresti (1999), Dorazio and Royle (2003), and others. 2004

As with model  $M_0$ , we have the Bernoulli model for the zero-inflation variables:  $z_i \sim \text{Bern}(\psi)$  and the model of the observations expressed conditional on the latent variables  $z_i$ . For  $z_i = 1$ , we have a binomial model with individual-specific  $p_i$ :

$$y_i|z_i=1 \sim \text{Bin}(K,p_i)$$

and otherwise  $y_i|z_i=0 \sim \delta(0)$ . Further, we prescribe a distribution for  $p_i$ . Here we assume

$$logit(p_i) \sim Normal(\mu, \sigma^2)$$

The basic **BUGS** description for this model, assuming a Unif(0,1) prior for  $p_0 = \log_{10}^{-1} (\mu)$ , is given as follows:

```
2012
    model{
2013
    p0 ~ dunif(0,1)
                              # prior distributions
2014
    mup < - log(p0/(1-p0))
2015
     taup~dgamma(.1,.1)
2016
    psi~dunif(0,1)
2017
2018
     for(i in 1:(nind+nz)){
2019
       z[i]~dbern(psi)
                              # zero inflation variables
2020
       lp[i] ~ dnorm(mup,taup) # individual effect
2021
       logit(p[i])<-lp[i]</pre>
2022
       mu[i]<-z[i]*p[i]
2023
       y[i]~dbin(mu[i],J) #
                                observation model
2024
2025
    N<-sum(z[1:(nind+nz)]) # N is a derived parameter
2027
    }
2028
```

# 4.4.2 Analysis of the Fort Drum data

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The logit-normal heterogeneity model was fitted to the bear data from the Fort Drum study, and we used data augmentation to produce a data set of M = 500 individuals. We ran the model using **JAGS** with the instructions given as follows<sup>5</sup>.

```
[... get data as before ....]
2033
2034
     set.seed(2013)
2035
2036
     cat("
2037
     model{
2038
     p0 ~ dunif(0,1)
                               # prior distributions
2039
     mup < - log(p0/(1-p0))
2040
     taup~dgamma(.1,.1)
2041
     sigmap<-sqrt(1/taup)</pre>
2042
     psi~dunif(0,1)
2043
2044
     for(i in 1:(nind+nz)){
       z[i]~dbern(psi)
                               # zero inflation variables
2046
       lp[i] ~ dnorm(mup,taup) # individual effect
2047
       logit(p[i])<-lp[i]</pre>
2048
       mu[i]<-z[i]*p[i]
2049
       y[i]~dbin(mu[i],K)
                              #
                                  observation model
2050
2051
     N<-sum(z[1:(nind+nz)])</pre>
2053
2054
     ",file="modelMh.txt")
2055
2056
     library("rjags")
2057
     jm<- jags.model("modelMh.txt", data=data1, inits=inits, n.chains=4,</pre>
2058
                         n.adapt=1000)
2059
     jout <- coda.samples(jm, params1, n.iter=50000, thin=1)
2060
```

CHANGE THIS TO RUN SIGMA DUNIF(0,5) PRIOR INSTEAD OF TAUY ANDY IS WORKING THIS SECTION RIGHT NOW. KEY ISSUE IS THAT BEAR DATA HAVE VERY LONG RIGHT TAIL. PSSIBLY NOT EVEN IDENTIFIABLE FOR LOGIT-NORMAL MODEL. NEED TO RUN WINBUGS AND JAGS FOR M=500 AND RUN A 200K RUN AND THEN MAYBE A 500K RUN TO SEE HOW THINGS LOOK. THEN RUN MY R CODE BELOW FOR 5 MILLION ITERS OR SOME bs like that. I'm not sure if this is a teaching moment (Link 2003) or if we need a different example here!

This produces the posterior distribution for N shown in Fig. 4.4. Posterior summaries of parameters are given as follows:

<sup>&</sup>lt;sup>5</sup>For WinBUGS, should provide starts for lp and sigma or sometimes WinBUGS breaks

```
> summary(jout)
2072
    Iterations = 1001:201000
2073
    Thinning interval = 1
2074
    Number of chains = 4
2075
    Sample size per chain = 2e+05
2076
2077
     1. Empirical mean and standard deviation for each variable,
2078
        plus standard error of the mean:
2079
2080
                                 Naive SE Time-series SE
                              SD
                  Mean
2081
    N
            108.63259 52.53176 5.873e-02
                                                    2.077726
    рO
               0.08615
                        0.05919 6.618e-05
                                                    0.001950
2083
               0.21841
                         0.10615 1.187e-04
                                                    0.004141
    psi
2084
              1.94096
                        0.51014 5.703e-04
                                                    0.018244
    sigmap
2085
2086
    2. Quantiles for each variable:
2087
2088
                 2.5%
                                      50%
                                                75%
                                                        97.5%
                            25%
2089
    N
            59.00000 77.00000 93.00000 121.0000 261.0000
2090
             0.00418
                       0.03852
                                  0.07657
                                             0.1240
                                                       0.2210
    p0
2091
             0.11230
                        0.15373
                                  0.18920
                                             0.2457
                                                       0.5241
2092
    sigmap
             1.11906
                       1.57643
                                  1.87752
                                             2.2386
                                                       3.1166
2093
```

We used M=500 for this analysis and we note that while the posterior mass of N is concentrated away from this upper bound (Fig. 4.4), the posterior has an extremely long right tail, with some posterior values at the upper bound N=500. Maybe or maybe not sufficient data augmentation. The model runs effectively in **WinBUGS** but sometimes with apparently inefficient mixing for reasons that may be related to bad starting values. In some cases this was resolved if we supplied starting values for the  $logit(p_i)$  parameters and  $\tau$ .

to do: insert final results. longer run. more data augmentation. compare with winbugs.

The posterior mode compares well with the MLE which we obtained using the **R** code contained in Panel 6.1 of Royle and Dorazio (2008). The MLE of  $log(n_0)$ , the logarithm of the number of uncaptured individuals, is  $log(n_0) = 3.86$  and therefore the MLE is  $\hat{N} = exp(3.86) + 47 = 94.47$  consistent with the apparent mode in Fig. 4.4. <sup>6</sup> To convert this to density we use the buffered area as computed above  $(255.3 \ km^2)^7$  and perform the required summary analysis on the posterior samples of N, which results in about 0.37 individuals/ $km^2$ . The reader should carry out this analysis to confirm the estimates, and also obtain the 95% confidence interval.

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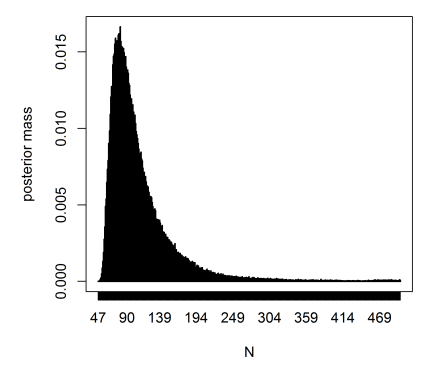
2107

2108

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 $<sup>^6</sup>$ We note that the result is inconsistent with Gardner et al. (2009) who reported an MLE of 104.1 ( $density = 0.437 inds/km^2$ ) although we do not know the reason for this at the present time.

 $<sup>^7</sup>$ WRONG #



**Figure 4.4.** Posterior of N for Fort Drum bear study data under the logit-normal version of model  $M_h$ . From WinBUGS output. 200k samples.

# 4.4.3 Building your own MCMC algorithm

For fun, we construct our own MCMC algorithm using a Metropolized Gibbs sampler for Model  $M_h$ . In chapter 10 we devise MCMC algorithms for spatial capture-recapture models and the basic conceptual and technical considerations are entirely analogous to Model  $M_h$ .

To begin, we first collect all of our model components which are as follows:  $[y_i|p_i,z_i], [p_i|\mu_p,\sigma_p], \text{ and } [z_i|\psi] \text{ for } each \ i=1,2,\ldots,M \text{ and then prior distributions } [\mu_p], [\sigma_p] \text{ and } [\psi].$  The joint posterior distribution of all unknown quantities in the model is proportional to the joint distribution of all elements  $y_i, p_i, z_i$  and also the prior distributions of the prior parameters:

$$\left\{\prod_{i=1}^{M}[y_i|p_i,z_i][p_i|\mu_p,\sigma_p][z_i|\psi]\right\}[\mu_p,\sigma_p,\psi]$$

For prior distributions, we assume that  $\mu_p, \sigma_p, \psi$  are mutually independent and for  $\mu_p$  and  $\sigma_p$  we use improper uniform priors, and  $\psi \sim \text{Unif}(0,1)$ . Note that the likelihood contribution for each individual, when conditioned on  $p_i$  and  $z_i$ , does not depend on  $\psi$ ,  $\mu_p$ , or  $\sigma_p$ . As such, the full-conditionals for the structural parameters  $\psi$  only depends on the collection of data augmentation variables  $z_i$ , and that for  $\mu_p$  and  $\sigma_p$  will only depends on the collection of latent variables  $p_i$ ; i = 1, 2, ..., M. The full conditionals for all the unknowns are as follows:

(1) For  $p_i$ :

$$[p_i|y_i, \mu_p, \sigma_p, z_i = 1] \propto [y_i|p_i][p_i|\mu_p, \sigma_p^2] \text{ if } z_i = 1$$
$$[p_i|\mu_p, \sigma_p] \text{ if } z_i = 0$$

2129 **(2)** for 
$$z_i$$
:

$$z_i|\cdot \propto [y_i|z_i*p_i]\mathrm{Bern}(z_i|\psi)$$

2130 (3) For 
$$\mu_p$$
:

$$[\mu_p|\cdot] \sim \prod_i [p_i|\cdot] * \text{const}$$

2131 (4) For 
$$\sigma_p$$
:

$$[\sigma_p|\cdot] \sim \prod_i [p_i|\cdot] * \text{const}$$

(5) For 
$$\psi$$
:

$$\psi|\cdot \sim \text{Beta}(1 + \sum z_i, 1 + M - \sum z_i)$$

What we've done here is identify each of the full conditional distributions in sufficient detail to toss them into our Metropolis-Hastings algorithm. With the exception of  $\psi$  which has a convenient analytic solution – it is a beta distribution which we can easily sample directly. In truth, we could also sample  $\mu_p$  and  $\sigma_p^2$  directly with

certain choices of prior distributions. For example, if  $\mu_p \sim \text{Normal}(0, 1000)$  then the full conditional for  $\mu_p$  is also normal, etc.. We implement an MCMC algorithm for this model in the following block of  $\mathbf{R}$  code. The basic structure is: initialize the parameters and create any required output or intermediate "holders", and then begin the main MCMC loop which, in this case, generates 100000 samples.

```
2142
    ## obtain the bear data by executing the previous data grabbing
2143
    ## function
2144
2145
    temp<-getdata()
    M<-temp$M
2147
    K<-temp$K
2148
    ytot<-temp$ytot
2149
2151
    ###
2152
    ### MCMC algorithm for Model Mh
2153
    out <- matrix (NA, nrow=100000, ncol=4)
2155
    dimnames(out)<-list(NULL,c("mu","sigma","psi","N"))</pre>
    lp<- rnorm(M,-1,1)</pre>
2157
    p<-expit(lp)
    mu < - -1
2159
    p0<-exp(mu)/(1+exp(mu))
2160
    sigma<- 1
2161
    psi<- .5
2162
    z<-rbinom(M,1,psi)</pre>
    z[ytot>0]<-1
2164
2165
    for(i in 1:100000){
2166
2167
    ### update the logit(p) parameters
2168
    lpc<- rnorm(M,lp,1) # 0.5 is a tuning parameter</pre>
    pc<-expit(lpc)</pre>
2170
    lik.curr<-log(dbinom(ytot,K,z*p)*dnorm(lp,mu,sigma))</pre>
    lik.cand<-log(dbinom(ytot,K,z*pc)*dnorm(lpc,mu,sigma))</pre>
2172
    kp<- runif(M) < exp(lik.cand-lik.curr)</pre>
    p[kp] <-pc[kp]
2174
    lp[kp]<-lpc[kp]
2176
    p0c<- rnorm(1,p0,.05)
    if(p0c>0 & p0c<1){
```

```
muc < -log(p0c/(1-p0c))
2179
     lik.curr<-sum(dnorm(lp,mu,sigma,log=TRUE))</pre>
     lik.cand<-sum(dnorm(lp,muc,sigma,log=TRUE))</pre>
2181
     if(runif(1) < exp(lik.cand-lik.curr)) {</pre>
      mu<-muc
2183
      p0<-p0c
2185
     }
2186
2187
     sigmac<-rnorm(1,sigma,.5)</pre>
2188
     if(sigmac>0){
2189
     lik.curr<-sum(dnorm(lp,mu,sigma,log=TRUE))</pre>
2190
     lik.cand<-sum(dnorm(lp,mu,sigmac,log=TRUE))</pre>
2191
     if(runif(1) < exp(lik.cand-lik.curr))</pre>
2192
      sigma<-sigmac
2193
2194
2195
     ### update the z[i] variables
2196
     zc<- ifelse(z==1,0,1) # candidate is 0 if current = 1, etc..</pre>
     lik.curr<- dbinom(ytot,K,z*p)*dbinom(z,1,psi)</pre>
2198
     lik.cand<- dbinom(ytot,K,zc*p)*dbinom(zc,1,psi)</pre>
    kp<- runif(M) < (lik.cand/lik.curr)</pre>
2200
     z[kp] \leftarrow zc[kp]
2202
    psi \leftarrow rbeta(1, sum(z) + 1, M - sum(z) + 1)
2203
     out[i,]<- c(mu,sigma,psi,sum(z))</pre>
2205
     }
2206
```

**Remarks**: (1) for parameters with bounded support, i.e.,  $\sigma_p$  and  $p_0$ , we are using a random walk candidate generator but rejecting draws outside of the parameter space. (2) We mostly use Metropolis-Hastings except for the data augmentation parameter  $\psi$  which we sample directly from its full-conditional distribution which is a beta distribution. (3) Even the latent data augmentation variables  $z_i$  are updated using Metropolis-Hastings although they too can be updated directly from their full-conditional.

#### 4.4.4 Exercises related to model Mh

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- (1) Enclose the MCMC algorithm in an R function and provide arguments for some of the parameters of the function that a user might wish to modify.
- 2217 (2) Execute the function and compare the results to those generated from Win-2218 BUGS in the previous section

(3) Note that the prior distribution for the "mean" parameter is given on  $p_0 = exp(\mu)/(1 + exp(\mu))$ . Reformulate the algorithm with a flat prior on  $\mu$  and see what happens. Contemplate this.

(4) Using Bayes rule, figure out the full conditional for  $z_i$  so that you don't have to use MH for that one. It might be more efficient. Is it?

# 4.5 INDIVIDUAL COVARIATE MODELS: TOWARD SPATIAL CAPTURE-RECAPTURE

A standard situation in capture-recapture models is when an individual covariate is measured, and this covariate is thought to influence encounter probability. As with other closed population models, we begin with the basic binomial observation model:

$$y_i \sim \text{Bin}(K, p_i)$$

and we assume also a model for encounter probability according to:

$$logit(p_i) = \alpha_0 + \alpha_1 x_i$$

Classical examples of covariates influencing detection probability are type of animal (juvenile/adult or male/female), a continuous covariate such as body mass (Royle and Dorazio, 2008, chapt. 6), or a discrete covariate such as group or cluster size. For example, in models of aerial survey data, it is natural to model detection probabilities as a function of the observation-level individual covariate, "group size" (Royle, 2008, 2009; Langtimm, 2010).

Such "individual covariate models" are similar in structure to Model  $M_h$ , except that the individual effects are observed for the n individuals that appear in the sample. These models are important here because spatial capture-recapture models are precisely a form of individual covariate model, an idea that we will develop here and elsewhere. Specifically, they are such models, but where the individual covariate is a partially observed latent variable similar. That is, unlike Model Mh, we do have some direct information about the latent variable, which comes from the spatial locations/distribution of individual recaptures. More on that later.

Traditionally, estimation of N in individual covariate models is achieved using methods based on ideas of unequal probability sampling (i.e., Horwitz-Thompson estimation), see Huggins (1989) and Alho (1990). An estimator of N is

$$\hat{N} = \sum_{i} \frac{1}{\tilde{p}_{i}}$$

where  $\tilde{p}_i$  is the probability that individual i appeared in the sample. That is,  $\tilde{p}_i = \Pr(y_i > 0)$ . In practice,  $\tilde{p}_i$  is estimated from the conditional-likelihood formed by the encounter histories. Namely,

$$\Pr(y_i|y_i>0) = \Pr(y_i)/\Pr(y_i>0)$$

where we substitute

$$Pr(y_i > 0) = (1 - (1 - p_i)^K)$$

2250 with

$$logit(p_i) = \alpha_0 + \alpha_1 x_i$$

Here we take a formal model-based approach to Bayesian analysis of such models using data augmentation (Royle, 2009). Classical likelihood analysis of the so-called "full likelihood" is covered in some detail by Borchers et al. (2002). For Bayesian analysis of individual covariate models, because the individual covariate is unobserved for the N-n uncaptured individuals, we require a model to describe variation among individuals, essentially allowing the sample to be extrapolated to the population. For our present purposes, we consider a continuous covariate and we assume that it has a normal distribution:

$$x_i \sim \text{Normal}(\mu, \sigma^2)$$

Data augmentation can be applied directly to this class of models. In particular, reformulation of the model under DA yields a basic zero-inflated binomial model of the form:

$$z_i \sim \operatorname{Bern}(\psi)$$
  
 $y_i|z_i = 1 \sim \operatorname{Bin}(K, p_i)$   
 $y_i|z_i = 0 \sim \delta(0)$ 

In addition, we assume that  $p_i$  is functionally related to a covariate  $x_i$ , e.g., by the logit model given above, and we assume a distribution for  $x_i$  appropriate for the context.

Fully spatial capture-recapture models essentially use this formulation with a latent covariate that is directly related to the individual detection probability (see next Section). As with the previous models, implementation is trivial in the BUGS language. The BUGS specification is very similar to that for model  $M_h$ , but we require the distribution of the covariate to be specified, along with priors for the parameters of that distribution.

# 4.5.1 Example: Location of capture as a covariate.

If we had a regular grid of traps over some closed geographic system then we imagine that the average location of capture would be a decent estimate (heuristically) of an individual's home range center. Intuitively some measure of typical distance from home range center to traps for an individual should be a decent covariate to explain heterogeneity in encounter probability, i.e., individuals with more exposure to traps should have higher encounter probabilities and vice versa. A version of

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this idea was put forth by Boulanger and McLellan (2001) (see also Ivan (2012)), but using the Huggins-Alho estimator and with covariate "distance to edge" of the trapping array. A limitation of this basic approach is that it does not provide a solution to the problem that the trap area is fundamentally ill-defined, nor does it readily accommodate the inherent and heterogeneous variation in this measured covariate. Here, we provide an example of this type of heuristically motivated approach using the fully model-based individual covariate model described above analyzed by data augmentation. We take a slightly different approach than that adopted by Boulanger and McLellan (2001). By analyzing the full likelihood and placing a prior distribution on the individual covariate, we resolve the problem of having an ill-defined area over which the population size is distributed. After you read later chapters of this book, it will be apparent that SCR models represent a formalization of this heuristic procedure.

For our purposes here, we define  $x_i = ||s_i - x_0||$  where  $s_i$  is the average encounter location of individual i and  $x_0$  is the centroid of the trap array. Conceptually, individuals in the middle of the array should have higher probability of encounter and, as  $x_i$  increases,  $p_i$  should therefore decrease. We note that we have defined  $s_i$  in terms of a sample quantity - the observed mean - which is ad hoc but maybe satisfactory under the circumstances. That said, for an expansive, dense trapping grid then we might expect the sample mean encounter location to be a good estimate of home range center but, clearly this is biased for individuals that live around the edge (or off) the trapping array. Regardless, it should be good enough for our present purposes of demonstrating this heuristically appealing application of an individual covariate model. A key point is that  $s_i$  is missing for each individual that is not encountered and thus so is  $x_i$ . Thus, it is a latent variable, or random effect, and we need therefore to specify a probability distribution for it. As a measurement of distance we know it must be positive-valued. Suppose further than we imagine no individual could have a home range radius larger than  $D_{max}$ . As such, we think a reasonable distribution for this individual covariate is

$$x_i \sim uniform(0, D_{max})$$

where  $D_{max}$  is a specified constant. In practice, people have used distance from edge of the trap array but that is less easy to define and compute. 2308

# Fort Drum Bear Study

We have to do a little bit of data processing to fit this individual covariate model to the Fort Drum data. To compute the average location of capture for each individual and the distance from the centroid of the trap array, we execute the following R instructions:

```
avg.s<-matrix(NA,nrow=nind,ncol=2)
2314
    for(i in 1:nind){
    tmp<-NULL
```

```
for(j in 1:T){
2317
    aa<-bearArray[i,,j]
2318
    if(sum(aa)>0){
2319
       aa<- trapmat[aa>0,]
       tmp<-rbind(tmp,aa)</pre>
2321
    }
2322
    }
2323
     avg.s[i,]<-c(mean(tmp[,1]),mean(tmp[,2]))
2324
    }
2325
    Cx<-mean(trapmat[,1])</pre>
2326
    Cy<-mean(trapmat[,2])
2327
    avg.s<-rbind(avg.s,matrix(NA,nrow=nz,ncol=2))</pre>
2328
    xcent < - sqrt((avg.s[,1]-Cx)^2 + (avg.s[,2]-Cy)^2)
2329
        To define the maximum distance (maxD) from the centroid, we use that of the
2330
    farthest trap, and so maxD is computed as follows:
2331
    minx<- min(trapmat[,1]-Cx)</pre>
2332
    maxx<-max(trapmat[,1]-Cx)</pre>
2333
    miny<- min(trapmat[,2]-Cy)
2334
    maxy<- max(trapmat[,2]-Cy)</pre>
2335
    # most extreme point determines maxD
2336
    ul<- c(minx,maxy)</pre>
2337
    \max D < - \operatorname{sqrt}( (ul[1]-0)^2 + (ul[2]-0)^2)
2338
        For the bear data the maxD was about 11.5 km. As such, the model described
2339
    above will produce an estimate of the population size of bears within 11.5 units of
2340
    the trap centroid<sup>8</sup>. The BUGS model specification and R commands to package
2341
    the data and fit the model are as follows:
2342
    cat("
2343
    model{
2344
    p0 ~ dunif(0,1)
                                # prior distributions
    mup < -log(p0/(1-p0))
2346
    psi~dunif(0,1)
     beta~dnorm(0,.01)
2348
2349
    for(i in 1:(nind+nz)){
2350
       xcent[i]~dunif(0,maxD)
2351
       z[i]~dbern(psi)
                                # DA variables
2352
       lp[i] <- mup + beta*xcent[i] # individual effect</pre>
2353
       logit(p[i])<-lp[i]</pre>
2354
```

<sup>&</sup>lt;sup>8</sup>To be convincing this might need a little bit of hand-holding

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```
mu[i]<-z[i]*p[i]
2355
      y[i]~dbin(mu[i],K)
                            #
                               observation model
2356
2357
    N<-sum(z[1:(nind+nz)])
2359
    ",file="modelMcov.txt")
    data2<-list(y=ytot,nz=nz,nind=nind,K=T,xcent=xcent,maxD=11.5)
2361
    params2<-list('p0','psi','N','beta')</pre>
2362
    inits = function() {list(z=z, psi=psi, p0=runif(1),beta=rnorm(1)) }
2363
    fit2 = bugs(data2, inits, params2, model.file="modelMcov.txt",working.directory=getwd(),
2364
            debug=T, n.chains=3, n.iter=4000, n.burnin=1000, n.thin=4)
2365
```

Posterior summaries are given in Table ?? XYZ, and the posterior distribution of N is given in Figure XYZ. It might be perplexing that the estimated N is much lower than obtained by model Mh but there is a good explanation for this, discussed subsequently. That issue notwithstanding, it is worth pondering how this model could be an improvement (conceptually or technically) over some other model/estimator including M0 and Mh considered previously. Well, for one, we have accounted formally for heterogeneity due to spatial location of individuals relative to exposure to the trap array, characterized by the centroid of the array. Moreover, we have done so using a model that is based on an explicit mechanism, as opposed to a phenomenological one such as Model Mh. Moreover, importantly, using our new model, the estimated N applies to an explicit area which is defined by our prescribed value of maxD. That is, this area is a fixed component of the model and the parameter N therefore has explicit spatial context, as the number of individuals with home range centers less than maxD from the centroid of the trap array. As such, the implied "effective trap area" for any maxD is that of a circle with radius maxD.

```
%% Not sure whether this should be a table or verbatim print-out
2382
    \begin{table}
2383
    \tabular{cccccccc}
    Node statistics
    node mean sd MC error 2.5% median 97.5% start sample
    N 58.89 5.483 0.2199 50.0 58.0 71.0 251 2250
    beta -0.246 0.06087 0.003892 -0.3592 -0.2457 -0.126 251 2250
    deviance 459.4 13.29 0.4496 435.7 458.4 487.8 251 2250
    p0 0.5409 0.06817 0.004052 0.4072 0.544 0.6678 251 2250
2390
    psi 0.1706 0.02572 7.759E-4 0.1247 0.1692 0.2242 251 2250
    \end{tabular}
2392
    \caption{..... xyz ......}
    \end{table}
2394
```

<sup>&</sup>lt;sup>9</sup>This is a bad use of this term. We have never defined ETA or ESA. What is it, exactly?

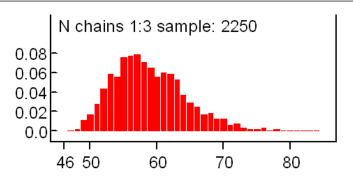


Figure 4.5. Needs a caption

5 \label{tab.maxD}

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We'll remake this figure in R. For now, insert it as is.

# 4.5.2 Extension of the Model

One important issue in understanding the meaning of estimates produced under the individual covariate model is that the uniform distribution on maxD implies that density is not constant over space. In particular, this model implies that it decreases as we move away from the centroid of the trap array. This is one reason we have a lower estimate of density than that obtained previously and also why, if we were to increase maxD, we would see density continue to decrease:  $x[i] \sim$ Uniform(0, maxD) implies constant N in each distance band from the centroid but obviously the area of each distance band is increasing. The reader can verify this as a homework exercise. Obviously, the use of an individual covariate model is not restricted to use of this specific distribution for the individual covariate. Clearly, it is a bad choice and, therefore, we should think about whether we can choose a better distribution for maxD - one that doesn't imply a decreasing density as distance from the centroid increases. Conceptually, what we want to do is impose a prior on distance from the centroid, x, such that density is proportional to the amount of area in each successive distance band as you move farther away from the centroid. In fact, there is theory that exists which tells us what the correct distribution of x is  $2x/maxD^2$ . This can be derived by noting that F(x) = Pr(X < x) =pi \* x \* x/pi \* maxD \* maxD. Then, f(x) = dF/dx = 2 \* x/(maxD \* maxD). This might be called a triangular distribution, I think, which makes sense because the incremental area in each additional distance band increases linearly with radius (i.e., distance from centroid). It is sometimes comforting to verify things empirically:

> u<-runif(10000,-1,1)

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It would be useful if we could describe this distribution in \*BUGS but there is not a built-in way to do this. One possibility is to use a discrete version of the pdf. We might also be able to use what is referred to in WinBUGS jargon as the "zeros trick" (see Advanced BUGS tricks) although we haven't pursued this approach. Instead, we consider using a discrete version and break Dmax into L distance classes of width  $\delta$ , with probabilities proportional to 2\*x. In particular, if the cut-points are  $xg[1] = 0, xg[2], \ldots, xg[L+1] = Dmax$  and the interval midpoints are  $xm[i] = xg[i+1] - \delta$ . Then, the interval probabilities are p[i] = 2\*xm[i]\*delta/(Dmax\*Dmax), which we can compute once and then send them to WinBUGS as data.

The R script is as follows. In the model description the variable x (observed home range center) has been rounded so that the discrete version of the f(x) can be used as described previously. The new variable labeled **xround** is actually then the integer category label in units of delta from 0. Thus, to convert back to distance in the expression for lp[i], **xround[i]** has to be multiplied by  $\delta$ .

```
delta<-.2
2440
    xround<-xcent%/%delta + 1
    Dgrid <- seq(delta, maxD, delta)
2442
    xprobs<- delta*(2*Dgrid/(maxD*maxD))</pre>
    xprobs<-xprobs/sum(xprobs)
2444
2445
    cat("
2446
    model{
2447
    p0 ~ dunif(0,1)
                              # prior distributions
    mup < - log(p0/(1-p0))
    psi~dunif(0,1)
    beta~dnorm(0,.01)
2451
    for(i in 1:(nind+nz)){
2453
      xround[i]~dcat(xprobs[])
2454
      z[i]~dbern(psi)
                                                # zero inflation variables
2455
      lp[i] <- mup + beta*xround[i]*delta # individual effect</pre>
      logit(p[i])<-lp[i]</pre>
2457
      mu[i]<-z[i]*p[i]
      y[i]~dbin(mu[i],K) # observation model
2459
2460
```

**Table 4.4.** Table: Analysis of Fort Drum bear hair snare data using the individual covariate model, for different values of Dmax, the upper limit of the uniform distribution of 'distance from centroid of the trap array'

maxD mn SD [1,] 12 0.230 0.038 [2,] 15 0.244 0.041 [3,] 17 0.249 0.044 [4,] 18 0.249 0.043 [5,] 19 0.250 0.043 [6,] 20 0.250 0.044

```
2462 N<-sum(z[1:(nind+nz)])
2463 }
2464 ",file="modelMcov.txt")

To fit the model we do this - keeping in mind that the data objects required
2465 below have been defined in previous analyses of this chapter:

2467 data2<-list(y=ytot,nz=nz,nind=nind,K=T,xround=xround,xprobs=xprobs,delta=delta)
2468 params2<-list('p0','psi','N','beta')
2469 inits = function() {list(z=z, psi=psi, p0=runif(1),beta=rnorm(1) ) }
2470 fit = bugs(data2, inits, params2, model.file="modelMcov.txt",working.directory=getwd(),
2471 debug=FALSE, n.chains=3, n.iter=11000, n.burnin=1000, n.thin=2)</pre>
```

This is a useful model because it induces a clear definition of area in which the population of N individuals reside. Under this model, that area is defined by specification of maxD. We can apply the model for different values of maxD and observe that the estimated N varies with maxD. Fortunately, we see empirically, that while N seems highly sensitive to the prescribed value of maxD, density seems to be invariant to maxD as long as it is chosen to be sufficiently large. We fit the model for maxD = 12 (points in close proximity to the trap arra) to 20 for and the results are given in Table ??.

We see that the posterior mean and SD of density (individuals per square km) appear insensitive to choice of  $\max D$  once we get a slight ways away from the maximum observed value of about 11.5. The estimated density of 0.250 per km² is actually quite a bit lower than we reported using model Mh (0.37, see section XYZ above) for which sample area is not an explicit feature of the model. On the other hand it is higher than that reported from Model M0 using the buffered area (0.195). There is no basis really for comparing or contrasting these various estimates and it would be a useful philosophical exercise for the reader to discuss this matter. In particular, application of model M0 and Mh are distinctly not spatially explicit models – the area within which the population not resides is not defined under either model. There is therefore no reason at all to think that the estimates produced under either model, using a buffered area, are justifiable based on any theory. In fact, we would get exactly the same estimate of N no matter what we declare the area to be. On the other hand, the individual covariate model explicitly describes

 $<sup>^{10}</sup>$ We need to look back at Chapter 1 and make sure we quit calling this "sample area" - it really isn't that at al, but rather the area within which N resides.

a distribution for "distance from centroid" that is a reasonable and standard null model - it posits, in the absence of direct information, that individual home range centers are randomly distributed in space and that probability of detection depends on the distance between home range center and the centroid of the trap array. Under this definition of the system, we see that density is invariant to the choice of sample area which seems like a desirable feature. The individual covariate model is not ideal, however, because it does not make full use of the spatial information in the data set, i.e., the trap locations and the locations of each individual encounter.

# 4.5.3 Invariance of density to maxD

Under the model above, and also under models that we consider in later chapters, a general property of the estimators is that while N increases with the prescribed trap area (equivalent to maxD in this case), we expect that density estimators should be invariant to this area. In the model used above, we note that  $Area(maxD) = \pi * maxD*maxD$  and  $E[N(maxD)] = \lambda * A(maxD)$  and thus  $E[Density(maxD)] = \lambda$  which is constant. This should be interpreted as the prior density. Absent data, then realizations under the model will have density  $\lambda$  regardless of what maxD is prescribed to be. As we verified empirically above, the posterior density is also invariant if maxD as long as the implied area (implied by maxD) is large enough so that the data no longer provide information about density (i.e., "far away"), then our estimator of density should become insensitive.

#### 4.5.4 Toward Fully Spatial Capture-recapture Models

We developed this model for the average observed location and equated it to home range center  $s_i$ . Intuitively, taking the average encounter location as an estimate of home range center makes sense but more so when the trapping grid is dense and expansive relative to typical home range sizes. However, our approach also ignored the variable precision with which each s[i] is estimated and also, as noted previously, estimates of s[i] around the "edge" (however we define that) are biased because the observations are truncated (we can only observe locations within the trap array). In the next Chapter we provide a further extension of this individual covariate model that definitively resolves the ad hoc nature of the individual covariate approach we took here. In that model we build a model in which s[i] are regarded as latent variables and the observation locations (i.e., trap specific encounters) are linked to those latent variables with an explicit model. We note that the model fitted previously could be adapted easily to deal with  $s_i$  as a latent variable, simply by adding a prior distribution for  $s_i$ . The reader should contemplate how to do this in WinBUGS.

# 4.6 DISTANCE SAMPLING: A PRIMATIVE SPATIAL CAPTURE-RECAPTURE MODEL

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Distance sampling is one of the most popular methods for estimating animal abundance. One of the great benefits of distance sampling is that it provides explicit 2531 estimates of density. The distance sampling model is a special case of a closed 2532 population model with a covariate. The covariate in this case,  $x_i$ , is the distance 2533 between an individual's location "u" and the observation location or transect. In fact, the model underlying distance sampling is precisely the same model as that 2535 which applies to the individual-covariate models, except that observations are made at only K=1 sampling occasion. In a sense, distance sampling is a spatial capture-2537 recapture model, but without the "recapture." This first and most basic spatial 2538 capture-recapture model has been used routinely for decades and, formally, it is a 2539 spatially-explicit model in the sense that it describes, explicitly, the spatial organi-2540 zation of individual locations (although this is not always stated explicitly) and, as a result, somewhat general models of how individuals are distributed in space can 2542 be specified (Royle, 2004b; Johnson, 2010; Sillett, 2011). As before, the distance 2543 sampling model, under data augmentation, includes a set of M zero-inflation vari-2544 ables  $z_i$  and the binomial model expressed conditional on z (binomial for z=1, and fixed zeros for z=0). In distance sampling we pay for having only a single 2546 sample (i.e., K = 1) by requiring constraints on the model of detection probability. 2547 A standard model is 2548

$$\log(p_i) = b * x_i^2$$

for b < 0, where  $x_i$  denotes the distance at which the *i*th individual is detected relative to some reference location where perfect detectability (p = 1) is assumed. This function corresponds to the "half-normal" detection function (i.e., with  $b = 1/\sigma^2$ ). If K > 1 then the intercept alpha is identifiable and such models are usually called "capture-recapture distance sampling" (Borchers, missing) and others XYZ?????).

As with previous examples, we require a distribution for the individual covariate  $x_i$ . The customary choice is

$$x_i \sim \text{Uniform}(0, B)$$

wherein B>0 is a known constant, being the upper limit of data recording by the observer (i.e., the point count radius, or transect half-width). In practice, this is sometimes asserted to be infinity, but in such cases the distance data are usually truncated. Specification of this distance sampling model in the BUGS language is shown in Panel 4.1. Royle and Dorazio (2008), p. xyz) provide a distance sampling example analyzed by DA using the famous Impala data.

As with the individual covariate model in the previous section, the distance sampling model can be equivalently specified by putting a prior distribution on individual *location* instead of distance between individual and observation point

```
b~dunif(0,10)
psi~dunif(0,1)
for(i in 1:(nind+nz)){
                       # DA Variables
   z[i]~dbern(psi)
   x[i]~dunif(0,B)
                       # B=strip width
   p[i]<-exp(logp[i])</pre>
                         # DETECTION MODEL
   logp[i]<-
               -((x[i]*x[i])*b)
   mu[i]<-z[i]*p[i]
   y[i]~dbern(mu[i])
                       # OBSERVATION MODEL
N \le sum(z[1:(nind+nz)])
D<- N/striparea # area of transects
```

Panel 4.1: Distance sampling model in WinBUGS, using a "half-normal" detection function.

(or transect). Thus we can write the general distance sampling model as

$$logit(p[i]) = alpha + beta * ||u[i] - x0||$$

567 Along with

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2577

$$\mathbf{u}_i \sim Uniform(\mathcal{S})$$

where  $x_0$  is a fixed point (or line) and u[i] is the individual's location which is observable for n individuals. In practice it is easier to record distance instead of location. Basic math can be used to argue that if individuals have a uniform distribution in space, then the distribution of Euclidean distance is also uniform. In particular, if a transect of length L is used and x is distance to the transect then  $F(x) = Pr(X \le x) = L * x/L * B = x/B$  and f(x) = dF/dx = (1/B). For measurements of radial distance, see the previous section.

In the context of our general characterization of SCR models (chapter 1.XYZ), we suggested that every SCR model can be described, conceptually, by a hierarchical model of the form:

Distance sampling ignores s, and treats u as observed data<sup>11</sup>. Thus, we are left with

In contrast, as we will see in the next chapters, basic SCR models (chapter 4) ignore u and condition on s, which is not observed:

<sup>&</sup>lt;sup>11</sup>Formally we could also say that  $[u] = \int [y|s][s]ds$ 

Since [u] and [s] are both assumed to be uniformly distributed, these are structurally equivalent models! The main differences have to do with interpretation of model components and whether or not the latent variables are observable (in distance sampling they are).

So why bother with SCR models when distance sampling yields density estimates and accounts for spatial heterogeneity in detection? For one, imagine try to collect distance sampling data on tigers! Clearly, distance sampling requires that one can collect large quantities of distance data, which is not always possible. For tigers, it is much easier, efficient, and safer to employ camera traps or tracking plates and then apply SCR models. Furthermore, as we will see in Ch XYZ, SCR models can use distance data to estimate all the parameters of our enchilada, allowing us to study distribution, movement, and density. Thus, SCR models are much more flexible than distance sampling models, and can accommodate data from virtually all animal survey designs.

# 4.6.1 Example: Muntjac deer survey from Nagarahole, India

Here we fit distance sampling models to distance sampling data on the muntjac deer (Muntiakus muntjak) collected in the year 2004 from Nagarahole National Park in southern India (Kumar, missing)(Kumar et al. unpublished data). The muntjac is a solitary species and distance measurements were made on 57 groups that were largely singletons with XYZ pairs of individuals. Commands for reading in and organizing the data for WinBUGS, followed by writing the model to a text file. Note that the total sampled area of the transects is fed in as "striparea" which is 708 (km of transect) multiplied by the strip width (B=150 = 0.15 km) multiplied by 2.

```
library("R2WinBUGS")
2606
     data <- read.csv("Muntjac.csv")
2607
     nind<-nrow(data)</pre>
2608
     y<-rep(1,nind)
2609
     nz<-400
2610
     y < -c(y, rep(0, nz))
2611
    x < -data[,3]
2612
     x<-c(x,rep(NA,nz))
2613
     data<-list(y=y,x=x,nz=nz,nind=nind,B=150,striparea=708*.15*2)
2615
2616
     cat("
2617
    model{
2618
     b~dunif(0,10)
2619
     psi~dunif(0,1)
2620
2621
```

```
for(i in 1:(nind+nz)){
2622
       z[i]~dbern(psi)
                             # DA Variables
2623
       x[i]~dunif(0,B)
                             # B=strip width
2624
                               # DETECTION MODEL
       p[i] <-exp(logp[i])</pre>
       logp[i]<-</pre>
                     -((x[i]*x[i])*b)
2626
       \#logp[i] \leftarrow -b*log(x[i]+1)
       mu[i]<-z[i]*p[i]
       y[i]~dbern(mu[i])
                            # OBSERVATION MODEL
2629
     }
2630
    N<-sum(z[1:(nind+nz)])</pre>
2631
    D<- N/striparea # area of transects
2632
    }
2633
    ",file="dsamp.txt")
2634
       Next, we provide inits, indicate which parameters to monitor, and then pass
2635
    those things to WinBUGS:
2636
    params<-list('b','N','D','psi')</pre>
2637
    inits = function() {list(z=z, psi=runif(1), b=runif(1,0,.02) )}
2638
    fit = bugs(data, inits, params, model.file="dsamp.txt",
    working.directory=getwd(),debug=T, n.chains=3, n.iter=4000, n.burnin=1000, n.thin=2)
2640
    Posterior summaries are provided in the following table. Estimated density is pretty
2641
    low, 1.1 individuals per sq. km. 12
     node mean sd MC error 2.5% median 97.5% start sample
2643
    D 1.096 0.1694 0.009122 0.8098 1.078 1.474 501 4500
    N 232.8 35.99 1.938 172.0 229.0 313.0 501 4500
2645
    b 5.678E-4 1.05E-4 4.129E-6 3.867E-4 5.616E-4 7.949E-4 501 4500
    deviance 681.2 16.72 0.7536 650.8 680.6 716.6 501 4500
    psi 0.5099 0.08238 0.004442 0.3681 0.5033 0.6918 501 4500
```

# 4.7 SUMMARY AND OUTLOOK

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Traditional closed population capture-recapture models are closely related to binomial generalized linear models. Indeed, the only real distinction is that in capture-recapture models, the population size parameter N (corresponding also to the size of a hypothetical "complete" data set) is unknown. This requires special consideration in the analysis of capture-recapture models. The classical approach to inference recognizes that the observations don't have a standard binomial distribution but, rather, a truncated binomial (from which which the so-called "conditional")

 $<sup>^{12}</sup>$  much lower than Samba's : Observers walked about 708 km from 39 transects in Nagarahole and the muntjac density is about 3 per sq km.. I need to get to the bottom of this.

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likelihood" derives) since we only have encounter frequency data on observed individuals. If instead we analyze the models using data augmentation, the observations can be modeled using a zero-inflated binomial distribution. In short, when we deal with the unknown-N problem using data augmentation then we are left with zero-inflated GLM and GLMMs instead of ordinary GLM or GLMMs. The analysis of such zero-inflated models is practically convenient, especially using the various Bayesian analysis packages that use the BUGS language.

Spatial capture-recapture models that we will consider in the rest of the chapters of this book are closely related to what have been called individual covariate models. Heuristically, spatial capture-recapture models arise by defining individual covariates based on observed locations of individuals – we can think of using some function of mean encounter location as an individual covariate. We did this in a novel way, by using distance to the centroid of the trapping array as a covariate. We analyzed the "full likelihood" using data augmentation, and placed a prior distribution on the individual covariate which was derived from an assumption that individual locations are, a priori, uniformly distributed in space. This assumption provides for invariance of the density estimator to the choice of population size area (induced by maximum distance from the centroid of the). The model addressed some important problems in the use of closed population models: it allows for heterogeneity in encounter probability due to the spatial context of the problem and it also provides a direct estimate of density because area is a feature of the model (via the prior on the individual covariate). The model is still not completely general because the model does not make use of the fully spatial encounter histories, which provide direct information about the locations and density of individuals. A specific individual covariate model that is in widespread use is classical "distance sampling." The model underlying distance sampling is precisely a special kind of SCR model - but one without replicate samples. Understanding distance sampling and individual covariate models more broadly provides a solid basis for understanding and analyzing spatial capture-recapture models.

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2687	FULLY SPATIAL CAPTURE-RECAPTURE
2688	MODELS

# FULLY SPATIAL CAPTURE-RECAPTURE MODELS

In previous sections we discussed some classes of models that could be viewed as primitive spatial capture-recapture models. We looked at a basic distance sampling model and we also considered a classical individual covariate modeling approach in which we defined a covariate to be the distance from (estimated) home range center to the center of the trap array. These were spatial in the sense that they included some characterization of where individuals live but, on the other hand, only a primitive or no characterization of trap location. That said, very little distinguishes these two models from spatial capture-recapture models that we consider in this chapter which fully recognize the spatial attribution of both individual animals and the locations of encounter devices.

Fully spatial capture-recapture models must accommodate the spatial organization of individuals and the encounter devices because the encounter process occurs at the level of individual traps. Failure to consider the trap-specific collection of data is the key deficiency with classical ad-hoc approaches which aggregate encounter information to the resolution of the entire trap array. We have seen previously some problems that this induces - imbalance in trap-level effort over time is problematic, and not being able to deal with trap-specific behavioral responses. Here, we resolve that by developing what is basically an individual covariate model but operating at the level of traps. That is, we develop our first fully spatial capture-recapture model which turns out to be precisely the model considered in section 3.XXX but instead of defining the individual covariate to be distance to centroid of the array we define J individual covariates - the distance to each trap. And, instead of using estimates of individual locations  $\mathbf{s}$ , we consider a fully hierarchical model in which we regard  $\mathbf{s}$  as a latent variable and impose a prior distribution on

it. We can think of having J independent capture-recapture studies generating one data set for each trap, and applying the individual covariate model with random activity centers, and that is all the basic SCR model is.

In the following sections of this chapter we investigate the basic spatial capturerecapture model and address some important considerations related to its analysis in **WinBUGS**. We also demonstrate how to summarize posterior output for the purposes of producing density maps or spatial predictions of density.

# 6.1 SAMPLING DESIGN AND DATA STRUCTURE

In our development here, we will assume a standard sampling design in which an array of J traps is operated for K time periods (say, nights) producing encounters of n individuals. Because sampling occurs by traps and also over time, the most general data structure yields encounter histories for each individual that are temporally and spatially indexed. Thus a typical data set will include an encounter history matrix for each individual. For the most basic model, there are no time-varying covariates that influence encounter, there are no explicit individual-specific covariates, and there are no covariates that influence density we will develop models in this chapter for encounter data that are aggregated over the temporal replicates. For example, suppose we observe 6 individuals in sampling at 4 traps over 3 nights of sampling then a plausible data set is the  $6\times 4$  matrix of encounters, out of 3, of the form:

2	736		trap1	trap2	trap3	trap4
2	737	[1,]	1	0	0	0
2	738	[2,]	0	2	0	0
2	739	[3,]	0	0	0	1
2	740	[4,]	0	1	0	0
2	741	[5,]	0	0	1	1
2	742	[6,]	1	0	1	0

We develop models in this chapter for devices such as "hair snares" or other DNA sampling methods (Kéry et al., 2010; Gardner et al., 2010) and related types of sampling problems so that we can suppose that "traps" may capture any number of individuals and an individual may be captured in any number of traps during each occasion but individuals can be encountered at most 1 time in a trap during any occasion. Thus, this is a "multi-catch" type of sampling (?, p. xyz). The statistical assumptions are that individual encounters within and among traps are independent. These basic (but admittedly at this point somewhat imprecise) assumptions define the basic spatial capture-recapture model, which we will refer to as "SCR0" henceforth¹ so that we may use that model as a point of reference

<sup>&</sup>lt;sup>1</sup>RC: It would be nice to have a running series of figures to display the various types of models. Each figure could have the same set of traps, use the same symbols, etc... It's probably worth showin g example data (and latent variables) in a table too

without having to provide a long-winded enumeration of assumptions and sampling design each time we do. We will make things more precise as we develop a formal statistical definition of the model shortly.

While the model is mostly directly relevant for hair snares and other DNA sampling methods for which multiple detections of an individual are not distinguishable, we will also make use of the model for data that arise from camera-trapping studies. In practice, with camera trapping, individuals might be photographed several times in a night but we will typically distill such data into a single binary encounter event for reasons discussed later in chapter 6.

#### 6.2 THE BINOMIAL OBSERVATION MODEL

We assume that the individual and trap-specific encounters,  $y_{ij}$ , are mutually independent outcomes of a binomial random variable:

$$y_{ij} \sim \text{Bin}(K, p_{ij}) \tag{6.2.1}$$

This is the basic model underlying "logistic regression" (chapter 2) as well as standard closed population models (chapter 3). The key element of the model is that the encounter probability  $p_{ij}$  is indexed by (i.e., depends on) both individual and trap. In a sense, then, we can think of each trap as producing individual level encounter history data of the classical variety - an nind x nrep matrix of 0's and 1's (this is the "encountered at most 1 time" assumption).

As we did in section XXX.YYY, we will make explicit the notion that  $p_{ij}$  is defined conditional on "where" individual i lives. Naturally, we think about defining an individual home range and then relating  $p_{ij}$  explicitly to the centroid of the individuals home range, or its center of activity (Efford, 2004; Borchers and Efford, 2008; Royle and Young, 2008). Therefore, define  $\mathbf{s}_i$ , a two-dimensional spatial coordinate, to be the activity center for individual i. Then, the basic SCR model postulates that encounter probability,  $p_{ij}$ , is related by a decreasing function to distance between trap j, having location  $\mathbf{x}_j$ , and  $\mathbf{s}_i$ . Naturally, if we think of modeling binomial counts using logistic regression, we might specify the model according to:

$$logit(p_{ij}) = \alpha_0 + \theta * ||s[i] - x[j]||$$
(6.2.2)

where, here,  $||\mathbf{s}_i - \mathbf{x}_j||$  is the distance between  $\mathbf{s}_i$  and  $\mathbf{x}_j$ . We sometimes write  $||\mathbf{s}_i - \mathbf{x}_j|| = dist(\mathbf{s}_i, \mathbf{x}_j) = d_{ij}$ . Alternatively, if we think about distance sampling then we might use the "half-normal" model of the form:

$$p_{ij} = p_0 * \exp(-\theta * ||\mathbf{s}_i - \mathbf{x}_j||^2)$$

Or any of a large number of standard detection models that are commonly used (we consider more in chapter XYZ). The half-normal model implies

$$\log(p_{ij}) = \log(p_0) - \theta * ||\mathbf{s}_i - \mathbf{x}_j||^2$$

$$(6.2.3)$$

Whatever model encounter probability we choose, we should always keep in mind that the model is described conditional on  $\mathbf{s}_i$ , which is an unobserved random variable. Thus, to be precise about this, we should write the observation model as

$$y_{ij}|\mathbf{s}_i \sim \text{Bin}(K, p(\mathbf{s}_{ij}; \theta))$$

Note that we probably expect that the parameter  $\theta$  in Eq. 6.2.2 or 6.2.3 should be negative, so that the probability of encounter decreases with distance between the trap and individual home range center. The joint likelihood for the data, conditional on the collection of individual activity centers, can therefore be expressed as

$$\mathcal{L}(\theta|\{\mathbf{y}_i,\mathbf{s}_i\}_{i=1}^N) = \prod_i \prod_j \text{Bin}(y_{ij}|p_{ij}(\theta))$$

Which, if we switch the indices on the product operators, this shows the SCR likelihood (conditional on  $\mathbf{s}$ ) to be the product of J independent capture-recapture likelihoods - one for each trap. However, the data have a "repeated measures" type of structure, with each of the j likelihood contributions for each individual being grouped by individual. Thus, we cannot analyze the model meaningfully by J trapspecific models. In classical repeated measures types of models, we accommodate the group structure of the data using random effects (random individual or group level variables). For SCR models we take the same basic approach, which we develop subsequently.

# 6.2.1 Distance as a latent variable

If we knew precisely every  $\mathbf{s}_i$  in the population (and how many, N), then the model specified by eqs. 6.2.1 and 6.2.2 or 6.2.3 is just an ordinary logistic regression type of a model which we learned how to fit using **WinBUGS** previously (chapt. 2), with a covariate  $d_{ij}$ . However, the activity centers are unobservable even in the best possible circumstances. In that case,  $d_{ij}$  is an unobserved variable, analogous to classical "random effects" models. We need to therefore extend the model to accommodate these random variables with an additional model component. A standard, and perhaps not unreasonable, assumption is the so-called "uniformity assumption" which is to say that the  $\mathbf{s}_i$  are uniformly distributed over space (the obvious next question "which space?" is addressed below). This uniformity assumption amounts to a uniform prior distribution on  $\mathbf{s}_i$ , i.e., the pdf of  $\mathbf{s}_i$  is constant, which we may express

$$Pr(\mathbf{s}_i)proptoconst$$
 (6.2.4)

To summarize the preceeding model developing, a basic SCR model is defined by 3 essential components:

- (1) Observation model:  $y_{ij}|\mathbf{s}_i \sim \text{Bin}(K, p_{ij})$
- (2) Encounter probability:  $logit(p_{ij}) = \alpha_0 + \theta * ||\mathbf{s}_i \mathbf{x}_j||$

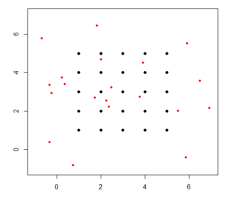


Figure 6.1. Realization of a binomial point process

# (3) Point process model: $\Pr[\mathbf{s}_i] \propto const$

Therefore, the SCR model is little more than an ordinary capture-recapture model for closed populations. It is such a model, but augmented with a set of "individual effects",  $\mathbf{s}_i$ , which relate some sense of individual location to encounter probability. As it turns out, assumption (3) is usually not precise enough to fit a model in practice for reasons we discuss in the following section. We will give another way to represent this prior distribution that is more concrete, but it depends on specifying the "state-space" of the random variable  $\mathbf{s}_i$ . The term "state-space" is a technical way of saying "possible outcomes".

# 6.3 THE BINOMIAL POINT-PROCESS MODEL

The collection of individual activity centers  $\mathbf{s}_1,\ldots,\mathbf{s}_N$  represent a realization of a binomial point process (Illian, 2008a, p. xyz). The binomial point process (BPP) is analogous to a Poisson point process in the sense that it represents a "random scatter" of points in space - except that the total number of points is fixed, whereas, in a Poisson point process it is random (having a Poisson distribution). As an example, we show in Fig. 6.1 locations of 20 individual activity centers (black dots) in relation to a grid of 25 traps. For a Poisson point process the number of such points in the prescribed state-space would be random whereas often we will simulate fixed numbers of points, e.g., for evaluating the performance of procedures such as how well does our estimator perform of N=50?

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It is natural to consider a binomial point process in the context of capture-recapture models because it preserves N in the model and thus preserves the linkage directly with closed population models. In fact, under the binomial point process model then Model M0 and other closed models are simple limiting cases of SCR models. In addition, use of the BPP model allows us to use data augmentation for Bayesian analysis of the models as in chapter 3, thus yielding a methodologically coherent approach to analyzing the different classes of models. Despite this, making explicit assumptions about N, such as Poisson, is convenient in some cases (see chapt. XYZ).

One consequence of having fixed N, in the BPP model, is that the model is not strictly a model of "complete spatial randomness". This is because if one forms counts  $n(A_1), \ldots, n(A_k)$  in any set of disjoint regions say  $A_1, \ldots, A_k$ , then these counts are not independent. In fact, they have a multinomial distribution (see Illian, 2008a, p. XYZ). Thus, the BPP model introduces a slight bit of dependence in the distribution of points. However, in most situations this will have no practical effect on any inference or analysis and, as a practical matter, we will usually regard the BPP model as one of spatial independence among individual activity centers because each activity center is distributed independently of each other activity center. Despite this implicit independence we see in Fig. 6.1 that realizations of randomly distributed points will typically exhibit distinct non-uniformity. Thus, independent, uniformly distributed points will almost never appear regularly, uniformly or systematically distributed. For this reason, the basic binomial (or Poisson) point process models are enormously useful in practical settings. More relevant for SCR models is that we actually have a little bit of data for some individuals and thus the resulting posterior point pattern can deviate strongly from uniformity (we should note this elsewhere too). The uniformity hypothesis is only a prior distribution which is directly affected by the quantity and quality of observations.

# 6.3.1 Definition of home range center

Some will be offended by our use of the concept of "home range center" and thus will have difficulty in believing that the resulting model is really useful for anything. Indeed, the idea of a home range or activity center is a vague concept anyway, a purely phenomenological construct. Despite this, it doesn't really matter whether or not a home range makes sense for a particular species - individuals of any species inhabit some region of space and we can define the "home range center" to be the center of the space that individual was occupying (or using) during the period in which traps were active. Thinking about it in that way, it could even be observable (almost) as the centroid of a very large number of radio fixes over the course of a survey period or a season. Thus, this practical version of a home range center is a well-defined construct regardless of whether one thinks the home range concept is meaningful, even if individuals are not particularly territorial. This is why we usually use the term "activity center" or maybe even "centroid of space usage"

and we recognize that this construct is a transient thing which applies only to a well-defined period of study.

# 6.3.2 The state-space of the point process

Shortly we will focus on Bayesian analysis of this model with N known so that we can directly apply what we learned in chapter 2 to this situation. To do this, we note that the individual effects  $\mathbf{s}_i, \ldots, \mathbf{s}_N$  are unknown quantities and we will need to be able to simulate each  $\mathbf{s}_i$  in the population from the posterior distribution. It should be self-vident that we cannot simulate the  $\mathbf{s}_i$  unless we describe precisely the region over which those  $\mathbf{s}_i's$  are uniformly distributed. This is the quantity referred to above as the state-space, denoted henceforth by S, which is a region or a set of points comprising the potential values of  $\mathbf{s}_i$ . Thus, an equivalent explicit statement of the "uniformity assumption" is

$$\mathbf{s}_i \sim \mathrm{Unif}(\mathcal{S})$$

# Prescribing the state-space

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Evidently, we need to define the state-space,  $\mathcal{S}$ . How can we possibly do this objectively? Prescribing any particular  $\mathcal{S}$  seems like the equivalent of specifying a "buffer" which we criticized previously as being ad hoc. How is it that choosing a state-space is *not* ad hoc? As a practical matter, it turns out that estimates of density are insensitive to choice of the state-space. As we observed in chapter 11, it is true that N increases with  $\mathcal{S}$ , but only at the same rate as  $\mathcal{S}$  under the prior assumption of constant density. As a result, we say that density is invariant to  $\mathcal{S}$  as long as  $\mathcal{S}$  is sufficiently large. Thus, while choice of  $\mathcal{S}$  is (or can be) essentially arbitrary, once  $\mathcal{S}$  is chosen, it defines the population being exposed to sampling, which scales appropriately with the size of the state-space.

For our simulated system developed previously in this chapter, we defined the state space to be a square within which our traps were centered perfectly. For many practical situations this might be an acceptable approach to defining the state-space. We provide an example of this in section 6.7 below in which the trap array is irregular and also situated within a realistic landscape that is distinctly irregular. In general, it is most practical to define the state-space as a regular polygon (e.g., rectangle) containing the trap array without differentiating unsuitable habitat. Although defining the state-space to be a regular polygon has computational advantages (e.g., we can implement this more efficiently in WinBUGS and cannot for irregular polygons), a regular polygon induces an apparent problem of admitting into the state-space regions that are distinctly non-habitat (e.g., oceans, large lakes, ice fields, etc.). It is difficult to describe complex sets in mathematical terms that can be admitted to this spatial model. As an alternative, we can provide a representation of the state-space as a discrete set of points (section 6.9) that will allow specific points to be deleted or not depending on whether they represent habitat, or we can define the state-space as an intersection of polygons, and analysis of

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models with state-space defined in that way can be analyzed easily using MCMC (see section XYZ in chapt. 6). In what follows below we provide an analysis of the camera data defining the state-space to be a regular continuous polygon (a rectangle).

# 6.3.3 Invariance and the State-space as a model assumption

We will assert for all models we consider in this book that density is invariant to the size and extent of S, if S is sufficiently large. In fact, this only holds as long as our model relating  $p_{ij}$  to  $\mathbf{s}_i$  is a decreasing function of distance. We can prove this thinking about a 1-d case where E[y] for the "last cell" (i.e., for d > Bfor B large enough) is 0. So it always contributes nothing to the likelihood, i.e., E[n(lastcell)] = 0. [sketch out a proof of this], in regular situations in which the detection function decays monotonically with distance and prior density is constant. Sometimes our estimate of density can be influenced if we make S too small but this might be sensible if  $\mathcal{S}$  is naturally well-defined. As we discussed in chapter 1, choice of S is part of the model and thus it makes sense that estimates of density might be sensitive to its definition in problems where it is natural to restrict  $\mathcal{S}$ . One could imagine however that in specific cases where you're studying a small population with well-defined habitat preferences that a problem could arise because changing the state-space around based on differing opinions and GIS layers really changes the estimate of total population size. But this is a real biological problem and a natural consequence of the spatial formalization of capture-recapture models - a feature, not a bug or some statistical artifact - and it should be resolved with better information and research, and not some arbitrary statistical artifact. For situations where there is not a natural choice of  $\mathcal{S}$ , we should default to choosing S to be very large in order to achieve invariance or otherwise evaluate sensitivity of density estimates by trying a couple of different values of  $\mathcal{S}$ . This is a standard "sensitivity to prior" argument that Bayesians always have to be conscious of. We demonstrate this in our analysis of section 6.7 below. Note that  $area(\mathcal{S})$  affects data augmentation. If you increase  $area(\mathcal{S})$  then there are more individuals to account for and therefore the size of the augmented data set M must increase.

We have been told that one can carry-out non-Bayesian analyses of SCR models without having to specify the state-space of the point process or perhaps while only specifying it imprecisely. This assertion is incorrect. We assume people are thinking this because they don't have to specify it explicitly because someone else has done it for them in a package that does integrated likelihood. Even to do integrated likelihood (see chapter 9) we have to integrate the conditional-on-s likelihood over some 2-dimensional space. It might work that the integration can be done from  $-\infty$  to  $+\infty$  but that is a mathematical artifact of specific detection functions, and an implicit definition of a state-space that doesn't make biological sense, even though it may in fact be innocuous;

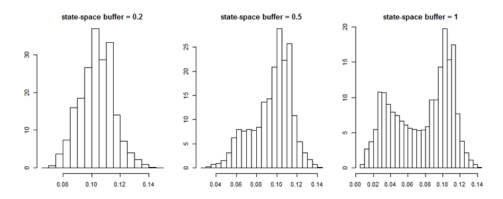


Figure 6.2. Needs a caption

#### 6.3.4 Connection to Model Mh

SCR models are closely related to heterogeneity models. In SCR models, heterogeneity in encounter probability is induced by both the effect of distance in the model for detection probability and also from specification of the state-space. Clearly then the state-space is explicitly part of the model. To understand this, we have a random effect with some prior distribution:

$$s \sim uniform(S)$$

And  $p(\mathbf{s}) = p(y=1|\mathbf{s})$  is some function of  $\mathbf{s}$ . Therefore, for any specific g(p) and  $\mathcal{S}$  we can work out what the implied heterogeneity model is for example, the mean, variance or other moments of the population distribution of p can be evaluated by integrating  $p(\mathbf{s})$  over the state-space of  $\mathbf{s}$ . Obviously the choice of  $p(\mathbf{s})$  and the choice of  $\mathcal{S}$  interact to determine the effective heterogeneity in p. We show an illustration in Fig. 6.2 below which shows a histogram of  $p(\mathbf{s})$  for a hypothetical population of 100000 individuals on a state-space enclosing our  $p(\mathbf{s})$  trap array above, under the logistic model for distance. Recode is provided in the Repackage scrbook to produce this analysis for the logistic and half-normal models. The histogram shows the encounter probability under buffers of 0.2, 0.5 and 1.0. We see the mass shifts to the left as the buffer increases, implying more individuals in the population but with lower encounter probability as their home range centers increase in distance from the trap array.

Another way to understand this is by representing S as a set of discrete points on a grid. In the coarsest possible case where S is a single arbitrary point, then every individual has exactly the same p. As we increase the number of points in S then more distinct values of p are possible. As such, when S is characterized by discrete points then SCR models are precisely a type of finite-mixture model (Norris III and

Pollock, 1996; Pledger, 2000), except where we have some information about which group an individual belong (i.e., where their activity center is), as a result of their captures in traps.

This context suggests the problem raised by Link (2003). He showed that in most practical situations N may not be identifiable across classes of mixture distributions which in the context of SCR models is the pair  $(g, \mathcal{S})$ . The difference, however, is that we do obtain some direct information about  $\mathbf{s}$  in SCR models and therefore N is identifiable across models characterized by  $(g, \mathcal{S})$ .

# 6.3.5 Connection to Distance Sampling

It is worth emphasizing that the basic SCR model is a binomial encounter model in which distance is a covariate. As such, it is striking similarity to a classical distance sampling model. Both have distance as a covariate but in classical distance sampling problems the focus is on the distance between the observer and the animal at an instant in time, not the distance between a trap and an animal's home range center. Thus in distance sampling, "distance" is observed for those individuals that appear in the sample. Conversely, in SCR problems, it is only imperfectly observed (we have partial information in the form of trap observations). Clearly, it is preferable to observe distance if possible, but as we will discuss in chapter XYZ, distance sampling requires field methods that are often not practical in many situations, e.g. when surveying tigers. Furthermore, SCR models allow us to relax many of the assumption made in classical distance sampling, and SCR models allow for estimates of quantities other than density, such as home range size.

#### 6.4 SIMULATING SCR DATA

It is always useful to simulate data because it allows you to understand the system that you're modeling and also calibrate your understanding with the parameter values of the model. That is, you can simulate data using different parameter values until you obtain data that "looks right" based on your knowledge of the specific situation that you're interested in. Here we provide a simple script to illustrate how to simulate spatial encounter history data. In this exercise we simulate data for 100 individuals and a 25 trap array laid out in a  $5\times 5$  grid of unit spacing. The specific encounter model is the half-normal model given above and we used this code to simulate data used in subsequent analyses. The 100 activity centers were simulated on a state-space defined by a  $8\times 8$  square within which the trap array was centered (thus the trap array is buffered by 2 units). Therefore, the density of individuals in this system is fixed at 100/64.

```
set.seed(2013)

representations with unit spacing traplocs traplocs cbind(sort(rep(1:5,5)),rep(1:5,5))
```

```
Dmat <- e2dist(traplocs, traplocs) # in cases where speed doesn't matter, it might be
3019
                                        # clearer to just show the slow for-loop.
3020
                                        # Plus, people will want to copy/paste this stuff
3021
    ntraps<-nrow(traplocs)</pre>
3022
3023
    # define state-space of point process. (i.e., where animals live).
3024
    # "delta" just adds a fixed buffer to the outer extent of the traps.
3025
3026
    X1<-min(traplocs[,1] - delta)</pre>
3027
    Xu<-max(traplocs[,1] + delta)</pre>
3028
    Yl<-min(traplocs[,2] - delta)
3029
    Yu<-max(traplocs[,2] + delta)
3030
3031
    N<-100
              # population size
3032
    K<- 20
               # number nights of effort
3033
3034
    sx<-runif(N,X1,Xu)</pre>
                             # simulate activity centers
3035
    sy<-runif(N,Y1,Yu)
3036
    S<-cbind(sx,sy)
3037
    D<- e2dist(S,traplocs) # distance of each individual from each trap
3038
3039
    alpha0<- -2.5
                         # define parameters of encounter probability
3040
    sigma<- 0.5
                         #
3041
    theta<- 1/(2*sigma*sigma)
3042
    probcap<- expit(-2.5)*exp( - theta*D*D)</pre>
                                                    # probability of encounter
    # now generate the encounters of every individual in every trap
    Y<-matrix(NA,nrow=N,ncol=ntraps)
3045
    for(i in 1:nrow(Y)){
3046
        Y[i,]<-rbinom(ntraps,K,probcap[i,])
3047
    }
3048
        Subsequently we will generate data using this code packaged in an R function
3049
    called simSCRO.fn which takes a number of arguments including discard0 which,
3050
    if TRUE, will return only the encounter histories for captured individuals. A second
3051
    argument is array3d which, if TRUE, returns the 3-d encounter history array instead
3052
    of the aggregated nind ×ntraps encounter frequencies (see below). Finally we
3053
    provide a random number seed, sd which we always set to 2013 in our analyses.
3054
    Thus we obtain a data set as above using the following command
3055
    data<-simSCRO.fn(discardO=TRUE, array3d=FALSE, sd=2013)
3056
    The R object data is a list, so let's take a look at what's in the list and then harvest
3057
    some of its elements for further analysis below.
3058
    > names(data)
3059
    [1] "Y"
                       "traplocs" "xlim"
                                                                "N"
                                                                             "alpha0"
                                                  "ylim"
                                                                                           "beta"
```

"K"

[8] "sigma"

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```
<sub>1062</sub> > Y<-data$Y
<sub>1063</sub> > traplocs<-data$traplocs
```

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#### 6.4.1 Formatting and manipulating real data sets

Conventional capture-recapture data are easily stored and manipulated as a 2dimensional array, an nind × nperiod matrix, which is maximally informative for any conventional capture-recapture model, but not for spatial capture-recapture models. For SCR models we must preserve the spatial information in the encounter history information. We will routinely analyze data from 3 standard formats:

- 3070 (1) The basic 2-dimensional data format, which is an nind × ntraps encounter frequency matrix such as that simulated previously;
  - (2) The maximally informative 3-dimensional array which we establish here the convention that it has dimensions nind × nperiods × ntraps and
- 3074 (3) We use a compact format the "SCR flat format" which we describe below in section 6.7.

To simulate data in the most informative format - the "3-d array" - we can use the **R** commands given previously but replace the last 4 lines with the following:

```
3078 Y<-array(NA,dim=c(N,K,ntraps))
3079 for(i in 1:nrow(Y)){
3080 for(j in 1:ntraps){
3081 Y[i,1:K,j]<-rbinom(K,1,probcap[i,j])
3082 }
3083 }</pre>
```

We see that a collection of K binary encounter events are generated for each individual and for each trap. The probabilities have those Bernoulli trials are computed based on the distance from each individuals home range center and the trap (see calculation above), and those are housed in the matrix probcap. Our data simulator function simSRCO.fn will return the full 3-d array if array3d=TRUE is specified in the function call. To recover the 2-d matrix from the 3-d array, and subset the 3-d array to individuals that were captured, we do this:

```
Y2d<- apply(Y,c(1,3),sum) # sum over the ''replicates'' dimension (2nd margin of the array)
ncaps<-apply(Y2d,1,sum) # compute how many times each individual was captured
Y<-Y[ncaps>0,,] # keep those individuals that were captured
```

#### 6.5 FITTING AN SCR MODEL IN BUGS

Clearly if we somehow knew the value of N then we could fit this model directly because, in that case, it is a special kind of logistic regression model - one with a random effect, but that enters into the model in a peculiar fashion - and also with a distribution (uniform) which we don't usually think of as standard for random

effects models. So our aim here is to analyze the known-N problem, using our simulated data, as an incremental step in our progress toward fitting more generally useful models.

To begin, we use our simulator to grab a data set and then harvest the elements of the resulting object for further analysis.

```
data<-simSCRO.fn(discard0=FALSE,sd=2013)
3103
    y<-data$Y
3104
    traplocs<-data$traplocs
3105
    nind<-nrow(y)</pre>
3106
    X<-data$traplocs
3107
    J<-nrow(X)
3108
    y<-rbind(y,matrix(0,nrow=(100-nrow(y)),ncol=J ) )</pre>
3109
    X1<-data$xlim[1]
3110
    Yl<-data$ylim[1]
3111
    Xu<-data$xlim[2]
3112
    Yu<-data$ylim[2]
```

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Note that we specify  ${\tt discard0} = {\tt FALSE}$  so that we have a "complete" data set, i.e., one with the all-zero encounter histories corresponding to uncaptured individuals. Now, within an  ${\tt R}$  session, we can create the  ${\tt BUGS}$  model file and fit the model using the following commands. This model describes the half-normal detection model but it would be trivial to modify that to various others including the logistic described above. One consequence of using the half-normal is that we have to constrain the encounter probability to be in [0,1] which we do here by defining alpha0 to be the logit of the intercept parameter p0. Note that the distance covariate is computed within the  ${\tt BUGS}$  model specification given the matrix of trap locations,  ${\tt X}$ , which is provided to  ${\tt WinBUGS}$  as data.

```
cat("
3124
     model {
3125
     alpha0~dnorm(0,.1)
3126
     logit(p0)<- alpha0
3127
     theta~dnorm(0,.1)
3128
     for(i in 1:N){
      s[i,1]~dunif(X1,Xu)
3130
      s[i,2]~dunif(Y1,Yu)
3131
     for(j in 1:J){
3132
     d[i,j] \leftarrow pow(pow(s[i,1]-X[j,1],2) + pow(s[i,2]-X[j,2],2),0.5)
3133
     y[i,j] ~ dbin(p[i,j],K)
3134
    p[i,j] \leftarrow p0*exp(-theta*d[i,j]*d[i,j])
3135
3137
     }
3138
3139
     ",file = "SCROa.txt")
```

3141

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Next we do a number of organizational activities including bundling the data for **WinBUGS**, defining some initial values, the parameters to monitor and some basic MCMC settings. We choose initial values for the activity centers **s** by generating uniform random numbers in the state-space but, for the observed individuals, we replace those values by each individual's mean trap coordinate for all encounters

```
sst<-cbind(runif(nind,X1,Xu),runif(nind,Y1,Yu)) # starting values for s
     for(i in 1:nind){
     if(sum(y[i,])==0) next
     sst[i,1]<- mean( X[y[i,]>0,1] )
     sst[i,2]<- mean( X[y[i,]>0,2] )
3150
3151
3152
    data <- list (y=y,X=X,K=K,N=nind,J=J,X1=X1,Y1=Y1,Xu=Xu,Yu=Yu)
3153
     inits <- function(){</pre>
      list (alpha0=rnorm(1,-4,.4),theta=runif(1,1,2),s=sst)
3155
3156
3157
    library("R2WinBUGS")
3158
    parameters <- c("alpha0","theta")</pre>
3159
    nthin<-1
    nc<-3
3162
    nb<-1000
3163
     out <- bugs (data, inits, parameters, "SCROa.txt", n.thin=nthin,
3164
    n.chains=nc, n.burnin=nb,n.iter=ni,debug=TRUE,working.dir=getwd())
3165
3166
```

There is little to say about the preceding basic operations other than to suggest that the interested reader explore the output and additional analyses by running the script provided in the  ${\bf R}$  package scrbook. We ran 1000 burn-in and 1000 after burn-in, 3 chains, to obtain 3000 posterior samples. Because we know N for this particular data set we only have 2 parameters of the detection model to summarize (alpha0 and theta). When the object out is produced we print a summary of the results as follows:

```
> print(out,digits=3)
3173
    Inference for Bugs model at "SCROa.txt", fit using WinBUGS,
     3 chains, each with 2000 iterations (first 1000 discarded)
3175
     n.sims = 3000 iterations saved
3176
                          sd
                                 2.5%
                                          25%
                                                  50%
                                                          75%
                                                                97.5% Rhat n.eff
                 mean
3177
               -2.496
                       0.224
                              -2.954 -2.648
                                               -2.48
                                                       -2.340
                                                               -2.091 1.013
                                                                               190
    alpha0
3178
                                1.638
                                        2.145
                                                 2.44
                                                        2.721
                                                                 3.303 1.005
                2.442 0.419
                                                                               530
    theta
3179
    deviance 292.803 21.155 255.597 277.500 291.90 306.000 339.302 1.006
                                                                               380
    For each parameter, n.eff is a crude measure of effective sample size,
3182
    and Rhat is the potential scale reduction factor (at convergence, Rhat=1).
3183
3184
    DIC info (using the rule, pD = Dbar-Dhat)
```

```
pD = -138.8 and DIC = 154.0 DIC is an estimate of expected predictive error (lower deviance is better).
```

We know the data were generated with alpha0 = -2.5 and theta = -2. The estimates look reasonably close to those data-generating values and we probably feel pretty good about the performance of the Bayesian analysis and MCMC algorithm that WinBUGS cooked-up based on our sample size of 1 data set. It is worth noting that the Rhat statistics indicate reasonable convergence but, as a practical matter, we might choose to run the MCMC algorithm for additional time to bring these closer to 1.0 and to increase the effective posterior sample size (n.eff). Other summary output includes "deviance" and related things including the deviance information criterion (DIC). We discuss these things in chapter XXXX.

#### 6.6 UNKNOWN N

In all real applications N is unknown and that fact is kind of an important feature of the capture-recapture problem! We handled this important issue in chapter 3 using the method of data augmentation which we apply here to achieve a realistic analysis of Model SCR0. As with the basic closed population models considered previously, we formulate the problem here by augmenting our observed data set with a number of "all zero" encounter histories - what we referred to in Chapter 3 as potential individuals. If n is the number of observed individuals, then let M-n be the number of potential individuals in the data set. For the basic  $y_{ij}$  data structure (individuals x traps encounter frequencies) we simply add additional rows of "all 0" observations to that data set. This is because such "individuals" are unobserved, and therefore necessarily have  $y_{ij} = 0$  for all j. A data set, say with 4 traps and 6 individuals, augmented with 4 pseudo-individuals therefore might look like this:

3210		trap1	trap2	trap3	trap4
3211	[1,]	1	0	0	0
3212	[2,]	0	2	0	0
3213	[3,]	0	0	0	1
3214	[4,]	0	1	0	0
3215	[5,]	0	0	1	1
3216	[6,]	1	0	1	0
3217	[7,]	0	0	0	0
3218	[8,]	0	0	0	0
3219	[9,]	0	0	0	0
3220	[10,]	0	0	0	0

We typically have more than 4 traps and, if we're fortunate, many more individuals in our data set.

For the augmented data, we introduce a set of binary latent variables (the data augmentation variables),  $z_i$ , and the model is extended to describe  $Pr(z_i = 1)$  which is, in the context of this problem, the probability that an individual in the

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3252

augmented data set is a member of the population that was sampled. In other words, if  $z_i = 1$  for one of the "all zero" encounter histories, this is implied to be a sampling zero whereas observations for which  $z_i = 0$  are "structural zeros" under the model.

How big does the augmented data set have to be? We discussed this issue in chapt. 3 where we noted that the size of the data set is equivalent to the upper limit of a uniform prior distribution on N. Practically speaking, it should be sufficiently large so that the posterior distribution for N is not truncated. On the other hand, if it is too large then unnecessary calculations are being done. An approach to choosing M by trial-and-error is indicated. You can take a ballpark estimate of the probability that an individual is captured (at all during the study), obtain N as n/pcap, and then set M=2\*N, as a first guess. Do a short MCMC run and then consider whether you need to do something different. See chapt. 10 for an example of this. Kery and Schaub (2011, ch. 6) provide an assessment of choosing M in closed population models.

Analysis by data augmentation removes N as an explicit parameter of the model. Instead, N is a derived parameter, computed by  $N = \sum_{i=1}^{M} z_i$ . Similarly, density, D, is also a derived parameter computed as  $D = N/area(\mathcal{S})$ . For our simulator, we're using an  $8 \times 8$  state-space and thus we will compute D as D = N/64.

#### 6.6.1 Analysis using data augmentation in WinBUGS

As before we begin by obtaining a data set using our simSCRO.fn routine and then harvesting the required data objects from the resulting data list. Note that we use the discardO=TRUE option this time so that we get a "real" data set with no all-zero encounter histories. After harvesting the data we produce the WinBUGS model specification which now includes M encounter histories including the augmented potential individuals, the data augmentation parameters  $z_i$ , and the data augmentation parameter  $\psi$ .

```
data<-simSCRO.fn(discard0=TRUE,sd=2013)
3253
     y<-data$Y
3254
     traplocs<-data$traplocs
3255
     nind<-nrow(y)</pre>
     X<-data$traplocs
3257
     J<-nrow(X)
3258
     X1<-data$xlim[1]
3259
     Y1<-data$ylim[1]
3260
     Xu<-data$xlim[2]</pre>
3261
     Yu<-data$ylim[2]
3262
     cat("
3264
     model {
3265
     alpha0~dnorm(0,.1)
     logit(p0)<- alpha0
```

111

```
theta~dnorm(0,.1)
3268
    psi~dunif(0,1)
3269
3270
    for(i in 1:M){
3271
     z[i] ~ dbern(psi)
3272
     s[i,1]~dunif(Xl,Xu)
3273
     s[i,2]~dunif(Y1,Yu)
3274
    for(j in 1:J){
3275
    d[i,j] \leftarrow pow(pow(s[i,1]-X[j,1],2) + pow(s[i,2]-X[j,2],2),0.5)
3276
    y[i,j] ~ dbin(p[i,j],K)
3277
    p[i,j] < z[i] *p0*exp(-theta*d[i,j]*d[i,j])
3278
3280
    N < -sum(z[])
3281
    D<-N/64
3282
    }
3283
     ",file = "SCROa.txt")
3284
        To prepare our data we have to augment the data matrix y with M-n all-
3285
    zero encounter histories, we have to create starting values for the variables z_i and
3286
     also the activity centers \mathbf{s}_i of which, for each, we require M values. Otherwise the
    remainder of the code for bundling the data, creating initial values and executing
     WinBUGS looks much the same as before except with more or differently named
3289
    arguments.
3290
    ## Data augmentation stuff
    y<-rbind(y,matrix(0,nrow=M-nind,ncol=ncol(y)))
    z<-c(rep(1,nind),rep(0,M-nind))
```

```
3291
3292
     sst<-cbind(runif(M,X1,Xu),runif(M,Y1,Yu)) # starting values for s
3296
    for(i in 1:nind){
3297
    if(sum(y[i,])==0) next
3298
    sst[i,1]<- mean( X[y[i,]>0,1] )
3299
    sst[i,2]<- mean( X[y[i,]>0,2] )
3300
3301
    }
    data \leftarrow list (y=y,X=X,K=K,M=M,J=J,X1=X1,Y1=Y1,Xu=Xu,Yu=Yu)
3302
    inits <- function(){</pre>
3303
      list (alpha0=rnorm(1,-4,.4), theta=runif(1,1,2), s=sst, z=z)
3304
    }
3305
3306
    library("R2WinBUGS")
    parameters <- c("alpha0","theta","N")
3309
    nthin<-1
    nc<-3
3310
    nb<-1000
3311
    ni<-2000
3312
```

3315

3316

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3349

3354

out <- bugs (data, inits, parameters, "SCROa.txt", n.thin=nthin,n.chains=nc,
n.burnin=nb,n.iter=ni,debug=TRUE,working.dir=getwd())

**Remarks**: (1) Note the differences in this new **WinBUGS** model with that appearing in the known-N version. (2) Also the input data has changed - the augmented data set has more rows of all-zeros. Previously we knew that N=100 but in this analysis we pretend not to know N, but think that N=200 is a good upper-bound; (3) Population size N(S) is a derived parameter, being computed by summing up all of the data augmentation variables  $z_i$  (as we've done previously); (4) Density,  $D \equiv D(S)$ , is also a derived parameter. Summarizing the output from **WinBUGS** produces:

```
> print(out1,digits=2)
3323
    Inference for Bugs model at "SCROa.txt", fit using WinBUGS,
      3 chains, each with 2000 iterations (first 1000 discarded)
      n.sims = 3000 iterations saved
3326
                mean
                         sd
                              2.5%
                                       25%
                                               50%
                                                      75%
                                                           97.5% Rhat n.eff
3327
               -2.57
                      0.23
                             -3.04
                                     -2.72
                                            -2.56
                                                    -2.41
                                                            -2.15 1.01
                                                                         320
    alpha0
3328
    theta
                2.46
                      0.42
                              1.63
                                      2.16
                                             2.46
                                                     2.73
                                                             3.33 1.02
                                                                         120
3329
    N
              113.62 15.73
                             86.00 102.00 113.00 124.00 147.00 1.01
                                                                         260
3330
3331
    D
                1.78 0.25
                              1.34
                                      1.59
                                             1.77
                                                     1.94
                                                            2.30 1.01
                                                                         260
    deviance 302.60 23.67 261.19 285.47 301.50 317.90 354.91 1.00
3332
3333
    For each parameter, n.eff is a crude measure of effective sample size,
3334
    and Rhat is the potential scale reduction factor (at convergence, Rhat=1).
3335
3336
    DIC info (using the rule, pD = var(deviance)/2)
3337
    pD = 279.9 and DIC = 582.5
    DIC is an estimate of expected predictive error (lower deviance is better).
3339
```

The column labeled "MC error" is the Monte Carlo error - the error inherent in the attempt to compute these posterior summaries by MCMC. It is desirable to run the Markov chain algorithm long enough so as to reduce the MC error to a tolerable level. What constitutes tolerable is up to the investigator. Certainly less than 1% is called for. As a general rule, Rhat gets closer to 1 and MC error decreases toward 0 as the number of iterations increases. We see that the estimated parameters ( $\alpha_0$  and  $\theta$ ) are comparable to the previous results obtained for the known-N case, and also not too different from the data-generating values. The posterior of N overlaps the data-generating value substantially with a mean of 113.62. To obtain these results we fitted the true data-generating model, that based on the half-normal detection model, to a single simulated data set. For fun and excitement we fit the wrong model - that with the logistic-linear detection model - to the same data set. This is easily achieved by modifying the WinBUGS model specification above, although we provide the R script in the R package scrbook. Those results are given below. We see that the estimate of N, the main parameter of interest, is very similar to that obtained under the correct model, convergence is worse (as measured by Rhat) which probably doesn't have anything to do with the model being wrong, and the posterior deviance and DIC favor the correct model. We consider the use of DIC for carrying-out model selection in chapter 12.

```
> print(out2,digits=2)
3359
    Inference for Bugs model at "SCROa.txt", fit using WinBUGS,
3360
     3 chains, each with 2000 iterations (first 1000 discarded)
3361
     n.sims = 3000 iterations saved
3362
                mean
                         sd
                              2.5%
                                       25%
                                              50%
                                                      75%
                                                           97.5% Rhat n.eff
3363
                                            -1.58
                                                           -1.071.05
3364
    alpha0
               -1.59
                      0.27
                             -2.16
                                    -1.77
                                                   -1.42
                                                                          60
                      0.43
                              2.92
                                      3.48
                                             3.79
                                                     4.05
                                                            4.66 1.04
                                                                           70
                3.77
3365
    N
              122.57 18.67
                             90.00 109.00 122.00 135.00 163.00 1.00
                                                                        3000
3366
                1.92
                     0.29
                              1.41
                                      1.70
                                             1.91
                                                     2.11
                                                            2.55 1.00
                                                                        3000
3367
    deviance 312.67 22.43 271.00 297.20 311.50 327.00 359.60 1.02
3368
3370
    For each parameter, n.eff is a crude measure of effective sample size,
    and Rhat is the potential scale reduction factor (at convergence, Rhat=1).
3371
3372
    DIC info (using the rule, pD = var(deviance)/2)
3373
    pD = 247.5 and DIC = 560.1
3374
    DIC is an estimate of expected predictive error (lower deviance is better).
3375
```

## 6.6.2 Use of other BUGS engines: JAGS

3376

There are two other popular BUGS engines in widespread use: OpenBUGS 3377 (Thomas et al., 2006) and **JAGS** (Plummer, 2003). Both of these are easily called 3378 from R. OpenBUGS can be used instead of WinBUGS by changing the package option in the bugs call to package=OpenBUGS. JAGS can be called using the 3380 function jags() in package R2JAGS which has nearly the same arguments as bugs(). 3381 We prefer to use the R library rjags (Plummer, 2009) which has a slightly different 3382 implementation that we demonstrate here as we reanalyze the simulated data set 3383 in the previous section (note: the same R commands are used to generate the data 3384 and package the data, inits and parameters to monitor). The function jags.model 3385 is used to initialize the model and run the MCMC algorithm for a period in which 3386 adaptive rejection (XXXX not sure XXXXX???) sampling is used. Then the 3387 Markov chains are updated using coda.samples() to obtain posterior samples for 3388 analysis, as follows: 3389

```
jm<- jags.model("SCROa.txt", data=data, inits=inits, n.chains=nc,
n.adapt=nb))
jm<- coda.samples(jm, parameters, n.iter=ni-nb, thin=nthin)</pre>
```

We find that JAGS seems to be 20-30% faster for the basic SCR model which the reader can evaluate using the script jags.winbugs.R in the R package scrbook.

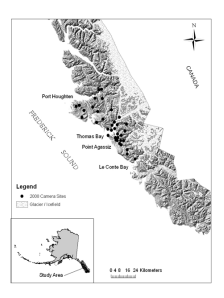


Figure 6.3. Wolverine camera trap locations from Magoun et al. (2011).

## 6.7 CASE STUDY: WOLVERINE CAMERA TRAPPING STUDY

We provide an analysis here of A. Magoun's wolverine data (Magoun et al., 2011; Royle et al., 2011c). The study took place in SE Alaska (Fig. 6.3) where 37 cameras were operational for variable periods of time (min = 5 days, max = 108 days, median = 45 days). A consequence of this is that the binomial sample size K (see Eq. 6.2.1) is variable for each camera. Thus, we must provide a matrix of sample sizes as data to BUGS and modify the model specification in sec. 6.6 accordingly. Our treatment of the data here is based on the analysis of Royle et al. (2011c).

To carry-out an analysis of these data, we require the matrix of trap coordinates and the encounter history data. We store data in an the "scr flat format" (see sec. 6.4.1 above), an efficient file format which is easily manipulated and also used as the input file format in our custom **R** script (ch. xxx) and **SPACECAP** (Gopalaswamy, 2012). To illustrate this format, the wolverine data are available as an encounter data **R** object named "wcaps" which has 3 columns and 115 rows, each representing a unique encounter event including the trap identity, the individual identity and the sample occasion index (sample). The first 10 rows of this matrix are as follows:

```
> wcaps
```

trapid individual sample

```
[1,]
                       1
                                       2
3414
                                       2
         [2,]
                       1
                                               128
3415
         [3.]
                       1
                                       2
                                               129
3416
                                      18
         [4,]
                       1
                                               130
3417
                       2
         [5,]
                                       3
                                               106
3418
                                      18
         [6,]
                       2
                                               104
3419
         [7,]
                       5
                                       5
                                                73
3420
         [8,]
                       5
                                       5
                                                89
3421
         [9,]
                       6
                                      18
                                              117
3422
        [10,]
                       6
                                      18
                                               118
3423
```

This "encounter data file" contains 1 row for each unique individual/trap encounter, and 3 variables (columns): trapid is an integer that runs from 1:ntraps, individual runs from 1:nind and sample runs from 1:nperiods. Often (as the case here) "sample" will correspond to daily sample intervals. The variable trapid will have to correspond to the row of a matrix containing the trap coordinates - a file named traplocs.csv available in the R package scrbook.

Note that these data do not represent a completely informative summary of the data. For example, if no individuals were captured in a certain trap or during a certain period, then this compact data format will have no record. Thus we will need to know ntraps and nperiods when reformatting this SCR data format into a 2-d encounter frequency matrix or 3-d array. In addition, the encounter data file does not provide information about which periods each trap was operated. This additional information is also necessary as the trap-specific sample sizes must be passed to **BUGS** as data. We provide this information in a 2nd data file - which we call the "trap deployment" file (described below).

The "encounter data file" wcaps.csv exists in the R package scrbook as a .csv file that people can read into R and do some basic summary statistics on. For our purposes we need to convert these data into the "individual x trap" array of binary encounter frequencies, although more general models might require an encounter-history formulation of the model which requires a full 3-d array. To obtain our nind x ntrap encounter frequency matrix, we do this the hard way by first converting the encounter data file into a 3-d array and then summarize to trap totals. We have a handy function SCR23darray.fn which takes the compact encounter data file with optional arguments ntraps and nperiods, and converts it to a 3-d array, and then we use the R function apply to summarize over the "sample" period dimension (by convention here, this is the 2nd dimension):

```
SCR23darray.fn <- function(caps,ntraps=NULL,nperiods=NULL){

nind<-max(caps[,2])

if(is.null(ntraps)) ntraps<-max(caps[,1])

if(is.null(nperiods)) nperiods<- max(caps[,3])

3454

3455 y<-array(0,c(nind,nperiods,ntraps))

tmp<-cbind(caps[,2],caps[,3],caps[,1])

y[tmp]<-1
```

```
3458     y
3459     }
3460
3461     # for the wolverine data do this:
3462
3463     Y3d <-SCR23darray.fn(wcaps,ntraps=37,nperiods=165)
3464     y <- apply(y3d,c(1,3),sum)</pre>
```

If ntraps and nperiods are not specified then they are assumed to be equal to the maximum value provided in the encounter data file. The 3-d array is necessary to fit certain types of models (e.g., behavioral response) and this is why we sometimes will require this maximally informative 3-d data format.

The other data file that we must have is the "trap deployment" file (henceforth "traps file") which provides the additional information not contained in the encounter data file. The traps file has  $\mathtt{nperiods} + 3$  columns. The first column is assumed to be a trap identifier, columns 2 and 3 are the easting and northing coordinates (assumed to be in a Euclidean coordinate system), and columns 4 to ( $\mathtt{nperiods} + 3$ ) are binary indicators of whether each trap was operational in each time period. The first 5 rows (out of 37) and 10 columns (out of 168) of the traps file for the wolverine data (" $\mathtt{wtraps.csv}$ " in the  $\mathbf{R}$  package  $\mathtt{scrbook}$  are:

```
Trap Easting Northing 1 2 3 4 5 6 7 <- column names
   1
       39040
                19216 0 0 0 0 0 0 0
   2
       41324
                19772 1 1 1 1 1 1 1
  3
       44957
                12985 0 0 0 0 0 0 0
   4
       41151
                23220 0 0 0 0 0 0 0
  5
       44240
                17198 0 0 0 0 0 0 0
```

This tells us that trap 2 was operated in periods 1-7 but the other traps were not operational during those periods. To extract the relevant information to fit the model in **WinBUGS** we do this:

```
traps<- read.csv("wtraps.csv")
traplocs<- traps[,2:3]
K<- apply(traps[,4:ncol(traps)],1,sum)</pre>
```

This results in a matrix traplocs which contains the coordinates of each trap and a vector K containing the number of days that each trap was operational. We now have all the information required to fit a basic SCR model in **WinBUGS**.

Summarizing these data files for the wolverine study, we see that 21 unique individuals were captured a total of 115 times. Most individuals were captured 1-6 times, with 4, 1, 4, 3, 1, and 2 individuals captured 1-6 times, respectively. In addition, 1 individual was captured each 8 and 14 times and 2 individuals each were captured 10 and 13 times. The number of unique traps that captured a particular individual ranged from 1-6, with 5, 10, 3, 1, 1, and 1 individual captured in each of 1-6 traps, respectively, for a total of 50 unique wolverine-trap encounters. These

numbers might be hard to get your mind around whereas some tabular summary is often more convenient. For that it seems natural to tabulate individuals by trap and total encounter frequencies. The spatial information in SCR data is based on multi-trap captures, and so, it is informative to understand how many unique traps each individual is captured in. At the same, it is useful to understand how many total captures we have of each individual because this is, in an intuitive sense, the effective sample size. So, we reproduce Table 1 from Royle et al. (2011c) which shows the trap and total encounter frequencies:

**Table 6.1.** Individual frequencies of capture for wolverines captured in camera traps in Southeast Alaska in 2008. Rows index unique trap frequencies and columns represent total number of captures (e.g., we captured 4 individuals 1 time, necessarily in only 1 trap; we captured 3 individuals 3 times but in 2 different traps)

								No.	of	captures
No. of traps	1	2	3	4	5	6	8	10	13	14
1	4	1	0	0	0	0	0	0	0	0
2	0	0	3	3	0	2	1	2	0	0
3	0	0	1	1	0	0	0	0	0	1
4	0	0	0	0	0	0	0	0	1	0
5	0	0	0	0	1	0	0	0	0	0
6	0	0	0	0	0	0	0	0	1	0

## 6.7.1 Fitting the model in WinBUGS

For illustrative purposes here we fit the simplest SCR model with the half-normal distance function although we revisit these data with more complex models in later chapters. The model is summarized by the following 3 components:

3511 (1)  $y_{ij}|\mathbf{s}_{i} \sim \text{Bin}(K, z_{i} \ p_{ij})$ 3512 (2)  $p_{ij} = p_{0} \exp(-\theta \ ||\mathbf{s}_{i} - x_{j}||^{2})$ 3513 (3)  $\mathbf{s}_{i} \sim \text{Unif}(\mathcal{S})$ 3514 (4)  $z_{i} \sim \text{Bern}(\psi)$ 

We assume customary flat priors on the structural (hyper-) parameters of the model,  $\alpha_0 = logit(p_0)$ ,  $\theta$  and  $\psi$ . It remains to define the state-space  $\mathcal{S}$ . For this, we nested the trap array (Fig. 6.3) in a a rectangular state-space extending 20 km beyond the traps in each cardinal direction. We also considered larger state-spaces up to 50 km to evaluate that choice. The buffer of the state space should be larger enough so that individuals beyond the state-space boundary are not likely to be encountered. Thus some knowledge of typical space usage patterns of the species is useful. The coordinate system was scaled so that a unit distance was equal to 10km, producing a rectangular state-space of dimension 9.88x10.5 units (area = 10374km\*km) within which the trap array was nested. As a general rule, we recommend scaling the state-space so that it is defined near the origin (x, y) = (0, 0). While the scaling of the

1.80 0.34

sigma

1.30

1.56

```
coordinate system is theoretically irrelevant, a poorly scaled coordinate system can
     produce Markov chains that mix poorly. We fitted this model in WinBUGS using
     data augmentation with M=300 potential individuals, using 3 Markov chains
3528
     each of 12000 total iterations, discarding the first 2000 as burn-in. [R commands
     for reading in the data and executing the analysis are as follows:
3530
    provide those commands here
     The output follows (note, we have a parameter "sigma" which we discuss shortly):
3532
     Buffer = 10 km
3533
     > print(out1$out,digits=2)
3534
     Inference for Bugs model at "modelfile.txt", fit using WinBUGS,
3535
      3 chains, each with 12000 iterations (first 2000 discarded)
3536
     n.sims = 30000 iterations saved
3537
                mean
                         sd
                               2.5%
                                       25%
                                               50%
                                                       75%
                                                            97.5% Rhat n.eff
                0.11
                       0.02
                              0.07
                                      0.10
                                              0.11
                                                     0.13
                                                             0.17
                                                                      1
                                                                         2400
3539
     psi
     sigma
                1.79
                       0.29
                              1.31
                                      1.58
                                              1.75
                                                     1.97
                                                             2.46
                                                                      1
                                                                          600
3540
    0g
                0.03
                       0.00
                              0.02
                                      0.03
                                              0.03
                                                     0.03
                                                             0.04
                                                                      1 13000
3541
    N
               33.02
                       4.99
                             25.00
                                     29.00
                                             32.00
                                                    36.00
                                                            44.00
                                                                         1600
3542
                                                                      1
    D
                4.93
                      0.75
                              3.73
                                      4.33
                                              4.78
                                                     5.38
                                                             6.57
                                                                         1600
3543
                                                                      1
                              0.08
                                                     0.20
                0.17 0.05
                                      0.13
                                              0.16
                                                             0.29
                                                                      1
                                                                          600
3544
    beta
     deviance 441.97 11.49 421.50 434.00 441.20 449.20 466.30
                                                                         6600
                                                                      1
3546
3547
     Buffer = 20 km
3548
     > print(out2$out,digits=2)
3549
     Inference for Bugs model at "modelfile.txt", fit using WinBUGS,
      3 chains, each with 12000 iterations (first 2000 discarded)
     n.sims = 30000 iterations saved
                mean
                         sd
                               2.5%
                                       25%
                                               50%
                                                       75%
                                                            97.5% Rhat n.eff
3553
     psi
                0.16
                       0.04
                              0.10
                                      0.13
                                              0.16
                                                      0.18
                                                             0.24
                                                                      1 4200
3554
                1.78
                       0.32
                              1.29
                                      1.55
                                              1.73
                                                     1.94
                                                             2.56
                                                                      1 20000
     sigma
3555
    p0
                0.03
                       0.00
                              0.02
                                      0.03
                                              0.03
                                                     0.03
                                                             0.04
                                                                         3000
3556
                                                                      1
               47.40
                      9.19
                             32.00
                                     41.00
                                                    53.00
                                                            68.00
                                                                         5900
                                             46.00
                                                                      1
3557
    N
                                                                      1 5900
                4.57
                      0.89
                              3.08
                                      3.95
                                                             6.55
    D
                                              4.43
                                                     5.11
                0.17 0.06
                              0.08
                                      0.13
                                              0.17
                                                     0.21
                                                             0.30
                                                                      1 20000
3559
     deviance 444.36 11.84 423.60 436.00 443.60 451.80 469.70
                                                                      1 1800
3560
3561
    Buffer = 25 \text{ km}
3562
    > print(out3$out,digits=2)
3563
     Inference for Bugs model at "modelfile.txt", fit using WinBUGS,
      3 chains, each with 12000 iterations (first 2000 discarded)
     n.sims = 30000 iterations saved
3566
                                       25%
                                               50%
                                                       75%
                                                            97.5% Rhat n.eff
3567
                mean
                         sd
                              2.5%
                0.19
                      0.04
                              0.11
                                      0.16
                                              0.19
                                                     0.22
                                                             0.29 1.00
                                                                          790
3568
    psi
```

1.75

1.98

2.59 1.01

400

```
0.03
    рO
                0.03 0.00
                               0.02
                                       0.03
                                                      0.03
                                                              0.04 1.00
3570
    N
                56.66 11.47
                              37.00
                                      48.00
                                             56.00
                                                     64.00
                                                             82.00 1.00
                                                                            570
3571
                 4.53
                      0.92
                               2.96
                                       3.84
                                              4.48
                                                      5.11
                                                              6.55 1.00
                                                                            570
    D
3572
                                              0.16
                                                      0.20
                 0.17 0.06
                               0.07
                                       0.13
                                                              0.30 1.01
                                                                            400
3573
    deviance 444.75 11.87 423.60 436.40 444.00 452.30 469.80 1.00 24000
3574
3575
    Buffer = 30 \text{ km}
3576
    > print(out4$out,digits=2)
3577
    Inference for Bugs model at "modelfile.txt", fit using WinBUGS,
3578
     3 chains, each with 12000 iterations (first 2000 discarded)
3579
3580
     n.sims = 30000 iterations saved
                                                50%
3581
                 mean
                         sd
                               2.5%
                                        25%
                                                        75% 97.5% Rhat n.eff
                 0.23
                       0.05
                               0.14
                                       0.19
                                               0.22
                                                      0.26
                                                              0.34 1.00
                                                                          1500
    psi
3582
                       0.34
                               1.29
                                               1.73
    sigma
                 1.79
                                       1.55
                                                      1.97
                                                              2.58 1.01
3583
    p0
                0.03 0.00
                               0.02
                                       0.03
                                               0.03
                                                      0.03
                                                              0.04 1.00 30000
3584
    N
                67.39 14.12
                              43.00
                                      57.00
                                             66.00
                                                     76.00
                                                             98.00 1.00
                                                                           1200
3585
    D
                 4.54 0.95
                               2.90
                                       3.84
                                               4.44
                                                      5.12
                                                              6.60 1.00
                                                                           1200
3586
                0.17 0.06
                               0.07
                                       0.13
                                              0.17
                                                      0.21
                                                              0.30 1.01
                                                                            560
3587
    deviance 444.58 11.83 423.60 436.40 443.80 452.20 469.90 1.00
                                                                           4700
3588
3589
    Buffer = 40 km (need to add this)
3590
3591
3592
3593
    Buffer = 45 \text{ km}
3594
    > print(out7$out,digits=2)
3595
    Inference for Bugs model at "modelfile.txt", fit using WinBUGS,
3596
     3 chains, each with 12000 iterations (first 2000 discarded)
3597
     n.sims = 30000 iterations saved
3598
                                                50%
                mean
                         sd
                               2.5%
                                        25%
                                                        75% 97.5% Rhat n.eff
3599
                      0.08
                 0.36
                               0.21
                                               0.35
                                                              0.53
                                                                       1 5000
    psi
                                       0.30
                                                       0.41
3600
                1.78 0.34
                               1.29
                                       1.55
                                               1.72
                                                       1.95
                                                              2.60
                                                                       1
                                                                            850
3601
    sigma
    p0
                 0.03
                       0.00
                               0.02
                                       0.03
                                               0.03
                                                       0.03
                                                              0.04
                                                                          3600
3602
              106.57 23.34
                              67.00
                                      90.00 104.00 121.00 157.00
                                                                           3400
3603
                 4.62
                      1.01
                               2.90
                                       3.90
                                               4.51
                                                      5.25
                                                              6.81
                                                                       1
                                                                           3400
3604
                 0.17 0.06
                               0.07
                                       0.13
                                               0.17
                                                       0.21
                                                              0.30
                                                                       1
                                                                            850
3605
3606
    deviance 444.80 11.84 423.60 436.40 444.10 452.30 470.00
                                                                       1 30000
3607
3608
    Buffer = 50 \text{ km}
    > print(out8$out,digits=2)
3609
    Inference for Bugs model at "modelfile.txt", fit using WinBUGS,
3610
     3 chains, each with 12000 iterations (first 2000 discarded)
3611
     n.sims = 30000 iterations saved
3612
                 mean
                          sd
                               2.5%
                                        25%
                                                50%
                                                        75%
                                                            97.5% Rhat n.eff
3613
3614
    psi
                 0.40
                       0.09
                               0.23
                                       0.33
                                               0.39
                                                       0.45
                                                              0.60 1.01 1300
    sigma
                 1.82
                      0.48
                               1.30
                                       1.56
                                               1.74
                                                      1.97
                                                              2.68 1.05
                                                                            200
3615
```

```
рO
                0.03
                       0.00
                               0.02
                                       0.03
                                              0.03
                                                      0.03
                                                              0.04 1.00
3616
    N
               118.47 26.81
                              71.00 100.00 117.00
                                                    135.00
                                                            176.00 1.01
                                                                          1200
3617
    D
                       1.02
                               2.71
                                       3.82
                                                                          1200
                4.52
                                              4.46
                                                      5.15
                                                              6.72 1.01
3618
                                                      0.21
                0.17
                       0.06
                               0.07
                                       0.13
                                              0.17
                                                              0.30 1.05
                                                                           200
    beta
    deviance 444.84 11.90 423.90 436.50 444.10 452.20 470.30 1.00
                                                                           500
```

We see that the estimated density is roughly consistent as we increase the state-space buffer from 20 to 50 km. We do note that the data augmentation parameter  $\psi$  (and, correspondingly, N) increase with the size of the state space in accordance with the deterministic relationship N=D\*A. However, density is constant more or less as we increase the size of the state-space beyond a certain point. For the 10 km state-space buffer, we see a noticeable effect on the posterior distribution of D. This is not a bug but rather a feature. As we noted above, the state-space is part of the model.

One thing we haven't talked about yet is that we can calibrate the desired size of the state-space by looking at the estimated home range radius of the species. For some models it is possible to convert the parameter  $\theta$  directly into the home range radius (section XXX XYZ). For the half-normal model we interpret the half-normal scale parameter  $\sigma$  which is related to  $\theta$  by  $\theta = 1/(2\sigma^2)$  as the radius of a bivariate normal movement model.

#### 6.7.2 Conclusion of Analysis

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Our point estimate of wolverine density from this study of approximately 4.5 individuals/ $1000~km^2$  and a 95% posterior interval is around [2.7, 6.3]. Density is estimated imprecisely which might not be surprising given the low sample size (n=21 individuals!). This seems to be a basic feature of carnivore studies although it should not (in our view) preclude the study of their populations nor attempts to estimate density or vital rates.

It is worth thinking about this model, and these estimates, computed under a rectangular state space roughly centered over the trapping array (Fig. 6.3). Does it make sense to define the state-space to include, for example, ocean? What are the possible consequences of this? What can we do about it? There's no reason at all that the state space has to be a regular polygon – we defined it as such here strictly for convenience and for ease of implementation in **WinBUGS** where it enables us to specify the prior for the activity centers as uniform priors for each coordinate. While it would be possible to define a more realistic state-space using some general polygon, it might take some effort to implement that in the **BUGS** language (see chapter XYZXYZ<sup>2</sup> for example of a simple case). Alternatively, we recommend using a discrete representation of the state-space – i.e., approximate  $\mathcal{S}$  by a grid of G points. We discuss this in the following section.

<sup>&</sup>lt;sup>2</sup>raccoon example or something?

## 6.8 CONSTRUCTING DENSITY MAPS

One of the most useful aspects of SCR models is that they are parameterized in 3654 terms of individual locations - i.e., where each individual lives - and, thus, we can 3655 compute many useful or interesting summaries of the activity centers. For example, 3656 we can make a spatial density plot by tallying up the number of activity centers 3657  $\mathbf{s}_i$  in boxes of arbitrary size and then producing a nice multi-color spatial plot of 3658 those which, we find, increases the acceptance probability of your manuscripts by a substantial amount. We discussed in chapter 2 the idea of estimating derived pa-3660 rameters from MCMC output. In SCR models, there are many derived parameters that are functions of the latent point locations  $(\mathbf{s}_1, \dots, \mathbf{s}_N)$ . In the present context, 3662 the number of individuals living in any well-defined polygon is a derived parameter. Specifically, let B(x) indicate a box centered at x then 3664

$$N(x) = \sum_{i} I(\mathbf{s}_{i} \in B(x))$$

is the population size of box B(x), and D(x) = N(x)/||B(x)|| is the local density. These are just "derived parameters" (see chapter 2) which are estimated from MCMC output using the appropriate Monte Carlo average. One thing to be careful about, in the context of models in which N is unknown, is that, for each MCMC iteration m, we only tabulate those activity centers which correspond to individuals in the sampled population. i.e., for which the data augmentation variable  $z_i = 1$ . In this case, we take all of the output for MCMC iterations  $m = 1, 2, \ldots, n$ iter and compute this summary:

$$N(x,m) = \sum_{z_{i,m}=1} I(s_{i,m} \in B(x))$$

Thus,  $N(x, 1), N(x, 2), \ldots$ , is the Markov chain for parameter N(x). In what follows we will provide a set of **R** commands for doing this calculations and making a basic image plot from the MCMC output.

Step 1: Define the center points of each box, B(x), or point at which local density will be estimated:

```
xg<-seq(X1,Xu,,50)
yg<-seq(Y1,Yu,,50)
```

Step 2: Extract the MCMC histories for the activity centers and the data augmentation variables. Note that these are each  $N \times \text{niter}$  matrices:

```
3682 Sxout<-out$sims.list$s[,,1]
3683 Syout<-out$sims.list$s[,,2]
2584 z<-out$sims.list$z
```

Step 3: We associate each coordinate with the proper box using the **R** command cut(). Note that we keep only the activity centers for which z=1 (i.e., individuals that belong to the population of size N):

```
Sxout<-cut(Sxout[z==1],breaks=xg,include.lowest=TRUE)
Syout<-cut(Syout[z==1],breaks=yg,include.lowest=TRUE)
```

Step 4: Use the table() command to tally up how many activity centers are in each B(x):

```
B692 Dn<-table(Sxout,Syout)
```

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3714

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Step 5: Use the image() command to display the resulting matrix.

```
image(xg,yg,Dn/nrow(z),col=terrain.colors(10))
```

Praise the Lord! This map is somewhat useful or at least it looks pretty and will facilitate the publication of your papers.

It is worth emphasizing here that density maps will not usually appear uniform despite that we have assumed that activity centers are uniformly distributed. This is because the observed encounters of individuals provide direct information about the location of the  $i=1,2,\ldots,n$  activity centers and thus their "estimated" locations will be affected by the observations. In a limiting sense, were we to sample space intensely enough, every individual would be captured a number of times and we would have considerable information about all N point locations. Consequently, the uniform prior would have almost no influence at all on the estimated density surface in this limiting situation. Thus, in practice, the influence of the uniformity assumption increases as the fraction of the population encountered decreases.

On the non-intuitiveness of image() – the R function image() might not be very intuitive to some – it plots M[1,1] in the lower left corner. If you want M[1,1] to be plotted "as you look at it" then M[1,1] should be in the upper left corner. We have a function rot() which does that. If you do image(rot(M)) then it puts it on the monitor as if it was a map you were looking at. You can always specify the x and y—labels explicitly as we did above.

**Spatial dot plots** – Now here is a cruder version based on the "spatial dot map" function spatial.plot. The useful functions in **R** are image() and image.scale() which is a function we grabbed off the web somewhere. Use of this function requires arguments of point locations and the resulting value to be displayed. The function is defined and applied as follows:

```
systial.plot<- function(x,y){
nc<-as.numeric(cut(y,20))
plot(x,pch=" ")
points(x,pch=20,col=topo.colors(20)[nc],cex=2)
image.scale(y,col=topo.colors(20))</pre>
```

```
3723 }
3724 # To execute the function do this:
3725 spatial.plot(cbind(xg,yg), Dn/nrow(z))
```

## 6.8.1 Example: Wolverine density map.

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We used the posterior output from the wolverine model fitted previous to compute a relatively coarse version of a density map, using a  $10 \times 10$  grid (Fig. 6.4) and using a  $30 \times 30$  grid (Fig. 6.5)<sup>3</sup>. In these figures density is expressed in units of individuals per  $1000 \ km^2$ , while the area of the pixels is about  $1037 \ km^2$  and  $115 \ km^2$ , respectively. That calculation is based on<sup>4</sup>:

```
3732 > total.area<- (Yu-Y1)*(Xu-X1)*1000
3733 > total.area/(10*10)
3734 [1] 1037.427
3735 > total.area/(30*30)
3736 [1] 115.2697
```

A couple of things are worth noting: First is that as we move away from "where the data live" - away from the trap array - we see that the density approaches the mean density. This is a property of the estimator as long as the "detection function" decreases sufficiently rapidly as a function of distance. Relatedly, it is also a property of statistical smoothers such as splines, kernel smoothers, and regression smoothers - predictions tend toward the global mean as the influence of data diminishes. Another way to think of it is that it is a consequence of the prior - which imposes uniformity, and as you get far away from the data, the predictions tend to the prior. The other thing to note about this map is that density is not 0 over water (although the coastline is not shown). This might be perplexing to some who are fairly certain that wolverines do not like water. However, there is nothing about the model that recognizes water from non-water and so the model predicts over water as if it were habitat similar to that within which the array is nested. But, all of this is ok as far as estimating density goes and, furthermore, we can compute valid estimates of N over any well-defined region which presumably wouldn't include water if we so choose.

#### 6.9 DISCRETE STATE-SPACE

The SCR model developed previously in this chapter assumes that individual activity centers are distributed uniformly over the prescribed state-space. Clearly this will not always be a reasonable assumption. In chapter 15 we talk about developing

<sup>&</sup>lt;sup>3</sup>Note: Not sure if we should use quantiles for color to make equal area slices. ??? Also should we use the same scale?

<sup>&</sup>lt;sup>4</sup>This is wrong and needs fixed. Move decimal one place over. i.e., 100 instead of 1000.

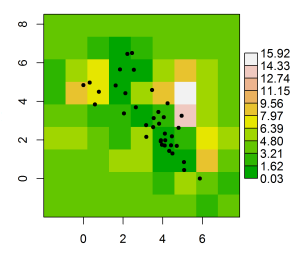


Figure 6.4. Needs a caption

models that allow explicitly for non-uniformity of the activity centers by modeling covariate effects on density. A simpler method of affecting the distribution of activity centers, which we address here, is to modify the shape of the state-space explicitly. For example, we might be able to classify the state-space into distinct blocks of habitat and non-habitat. In that case we can remove the non-habitat from the state-space and assume uniformity of the activity centers over the remaining portions judged to be suitable habitat. There are two ways to approach this: We can use a regular grid of points to represent the state-space, i.e., by the set of coordinates  $\mathbf{s}_1, \ldots, \mathbf{s}_G$ , and assign a equal probabilities to each possible value, or we can retain the continuous formulation of the state-space but use basic polygon operations to induce constraints on the state-space We focus here on the formulation of our basic SCR model in terms of a discrete state-space but later on (chapter 10 and also Appendix XYZ) we demonstrate the latter approach based on using polygon operations to define an irregular state-space.

Use of a discrete state-space can be computationally expensive in **WinBUGS**. That said, it isn't too difficult to do the MCMC calculations in  $\mathbf{R}$  which we discuss briefly in chapter 10. The  $\mathbf{R}$  package SPACECAP (Gopalaswamy et al., 2011) arose from the  $\mathbf{R}$  implementation developed for the application in Royle et al. (2009). As we will see in chapter 9, we must prescribe the state-space by a discrete mesh of points in order to do integrated likelihood and so if we are using a discrete

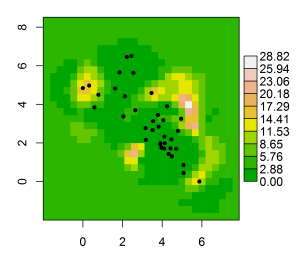


Figure 6.5. Needs a caption

state-space this can be accommodated directly in our code for obtaining MLEs.

While clipping out non-habitat seems like a good idea, its not obvious that we accomplish any biologically reasonable objective by doing so. We might prefer to do it when non-habitat represents a clear-cut restriction on the state-space such as a reserve boundary or a lake, ocean or river. It makes sense in those situations. Unfortunately, having the capability to do this also causes people to start defining "habitat" vs. "non-habitat" based on their understanding of the system whereas it can't be known whether the animal being studied has the same understanding. Moreover, differentiating of the landscape by habitat or habitat quality probably affects the geometry and morphology of home ranges much more than the plausible locations of activity centers. That is, a home range centroid could, in actual fact, occur in a walmart parking lot if there is pretty good habitat around walmart, so there is probably no sense to cut out the walmart lot and preclude it as the location for an activity center. It would generally be better to include some definition of habitat quality in the model for the detection probability (see chapter XYZ).

#### 6.9.1 Evaluation of Coarseness of Discrete Approximation

The coarseness of the state-space should not really have much of an effect on estimates if the grain is sufficiently fine relative to typical animal home range sizes.

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Why is this? We have two analogies that can help us understand this. First is the relationship to Model  $M_h$ . As noted in section 6.3.4 above, we can think about SCR models as a type of finite mixture (Norris III and Pollock, 1996; Pledger, 2000) where we are fortunate to be able to obtain direct information about which "group" individuals belong to (group being location of activity center). In the standard finite mixture models we typically find that only 1 or a very small number of groups (e.g., 2 or 3 at the most) can explain really high levels of heterogeneity and are adequate for most data sets of small to moderate sample sizes. We therefore expect a similar effect in SCR models when we discretize the state-space. We can also think about discretizing the state-space as being related to numerical integration where we find (see chapter 9) that we don't need a very fine grid of support points to evaluate the integral to a reasonable level of accuracy. We demonstrate this here by reanalyzing simulated data using a state-space defined by a different numbers of support points. We provide an R script called simSCROdiscrete.fn in the R package scrbook. We note that for this comparison we generated the actual activity centers as a continuous random variable and thus the discrete state-space is, strictly speaking, an approximation to truth. That said, we regard all state-space specifications as approximations to truth because they are all, strictly speaking, models of some unknown truth. Thus the use of any specific discrete state-space is not intrinsically more "wrong" than any specific continuous representation.

We used **JAGS** from the rjags function to obtain the results for  $6 \times 6$ ,  $9 \times 9$ ,  $12 \times 12$ ,  $15 \times 15$ ,  $20 \times 20$ ,  $25 \times 25$  and  $30 \times 30$  state-space grids. We used 2000 burn, 12000 total iters with 3 chains, therefore a total of 30000 posterior samples. For WinBUGS we used 3 chains of 5k total with 1k burnin means 12k total posterior samples. Summary results for these analyses are shown in Table XYZ<sup>5</sup>.

3819	Table	e XYZ.					
3820			Mean	SD	NaiveSE	Time-seriesSE	runtime
3821	6	N	109.7717	15.98959	0.0923160	0.377737	1239
3822	9	N	114.4621	16.72025	0.0965344	0.468659	1267
3823	12	N	115.4309	17.12403	0.098866	0.464830	1576
3824	15	N	114.7699	17.0242	0.0982894	0.425238	1638
3825	20	N	116.0370	17.10686	0.0987665	0.486867	1647
3826	25	N	116.3228	16.98323	0.0980527	0.465527	1661
3827	30	N	116.4252	17.4078	0.100504	0.533735	1806
3828	WinB	UGS					
3829			Mean	SD	NaiveSE	Time-seriesSE	runtime
3830	6	N	111.67	16.61			2274
3831	9	N	114.23	17.99			4300
3832	12	N	115.98	17.38			7100
3833	15	N	115.38	17.94			13010

<sup>&</sup>lt;sup>5</sup>Andy to finish later

3833 3834

```
Note: WinBUGS based on fewer samples too!

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To get SE and time-series SE do this:

You can use as.mcmc.list() to convert to a coda object. Then use summary.
```

The results in terms of the posterior summaries are, as we expect, very similar using **WinBUGS**. However, it was interesting to note that **WinBUGS** runtime is much worse (note the number of iterations is lower for **WinBUGS** yet the runtime is much longer) and, furthermore, it seems to scale with the size of the discrete state-space grid. While that was expected, it was unexpected that the runtime of **JAGS** would seem relatively consistent as we increase the grid size. We suspect that **WinBUGS** is evaluating the full-conditional for each activity center at all G possible values whereas it may be that **JAGS** is evaluating the full-conditional only at a subset of values or perhaps using previous calculations more effectively.

While this might suggest that one should always use **JAGS** for this analysis, we found in our analysis of the wolverine (next section) that **JAGS** could be extremely sensitive to starting values, producing MCMC algorithms that sometimes simply did not work.

#### 6.9.2 Analysis of the wolverine camera trapping data

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We reanalyzed the wolverine data using discrete state-space grids with points spaced by 2, 4 and 8 km (depicted in Fig. ??). These were constructed from the 40 km buffered state-space, and deleting the points over water (see Royle et al., 2011c). Our interest in doing this was to evaluate the relative influence of grid resolution on estimated density because the coarser grids will be more efficient from a computational stand-point and so we would prefer to use them, but perhaps not if there is a strong influence on estimated density.

**Note**: Results from WinBUGS are given below based on short runs that took a long long time. I am rerunning those. I will also show a density map for each analysis.

```
based on 2k burn 3k total and 3 chains = 3k total posterior samples.
    lots of MC error here.
3864
3865
3866
    For each parameter, n.eff is a crude measure of effective sample size,
3867
    and Rhat is the potential scale reduction factor (at convergence, Rhat=1).
3868
                        2.5%
                                25%
                                      50%
                                             75%
            mean
                    sd
                                                  97.5% Rhat n.eff
            0.28
                  0.06
                        0.17
                               0.24
                                     0.27
                                            0.32
                                                   0.41 1.01
                                                                230
    psi
    sigma
           0.64
                  0.05
                        0.55
                               0.60
                                     0.64
                                            0.67
                                                   0.73 1.02
                                                                 88
3871
           -3.00
                  0.16 -3.33 -3.11 -3.00 -2.90
                                                  -2.69 1.04
                                                                 52
3872
    p0
                  0.01 0.03 0.04 0.05
                                           0.05
                                                   0.06 1.04
                                                                 52
3873
           82.95 16.26 55.00 72.00 82.00 93.00 119.02 1.01
                                                                240
    N
3874
```

```
3875
    4 km
3876
    For each parameter, n.eff is a crude measure of effective sample size,
3877
    and Rhat is the potential scale reduction factor (at convergence, Rhat=1).
            mean
                    sd 2.5%
                                25%
                                      50%
                                             75% 97.5% Rhat n.eff
3879
            0.30
                  0.06 0.19
                              0.26
                                    0.29
                                           0.34
                                                   0.43 1.01
3880
    sigma 0.62
                  0.05 0.54 0.59
                                    0.62
                                            0.65
                                                   0.72 1.00
3881
    1am0 - 3.00
                  0.16 -3.33 -3.10 -2.99 -2.90
                                                  -2.67 1.01
3882
    рO
            0.05 0.01 0.03 0.04 0.05 0.05
                                                   0.06 1.01
                                                                390
3883
           88.78 16.76 60.00 77.00 87.00 99.00 125.00 1.01
    N
                                                                690
3884
3885
    For each parameter, n.eff is a crude measure of effective sample size,
3887
    and Rhat is the potential scale reduction factor (at convergence, Rhat=1).
3888
            mean
                    sd
                        2.5%
                                25%
                                      50%
                                             75% 97.5% Rhat n.eff
3889
            0.27
                  0.06
                        0.17
                               0.23
                                     0.27
                                            0.31
                                                   0.40 1.00
                                                               1500
3890
    psi
                  0.05 0.60
                               0.65
                                     0.68
                                           0.72
                                                   0.80 1.00
                                                               3000
    sigma 0.69
                  0.17 -3.41 -3.20 -3.07 -2.95
    1am0 - 3.07
                                                  -2.74 1.01
                                                                210
            0.04 0.01 0.03
                              0.04
                                    0.04 0.05
                                                   0.06 1.01
                                                                200
3893
           82.01 15.98 55.00 71.00 80.00 92.00 118.00 1.00
                                                               1300
3894
        We did the analysis in JAGS also. The results are shown below. Note: I am
3895
    going to run these again but for longer to finalize the results.
3896
    2km
3897
    Iterations = 7001:13000
3898
    Thinning interval = 1
3899
    Number of chains = 3
    Sample size per chain = 6000
                            SD Naive SE Time-series SE
3903
           86.28522 16.950626 1.263e-01
    N
                                               0.4878973
3904
            0.04807
                     0.007512 5.599e-05
                                               0.0002199
    lam0
3905
    рO
            0.04581
                     0.006820 5.083e-05
                                               0.0001996
3906
            0.28904 0.062117 4.630e-04
                                               0.0017481
3907
    psi
    sigma 0.62769 0.043596 3.249e-04
                                               0.0018724
3909
    4km
3910
                            SD Naive SE Time-series SE
               Mean
3911
           85.53139 16.998966 1.267e-01
                                               0.5181297
    N
3912
            0.04636
                     0.007542 5.621e-05
                                               0.0002382
3913
    lam0
            0.04425
                     0.006867 5.118e-05
                                               0.0002172
    p0
3914
    psi
            0.28650
                     0.061922 4.615e-04
                                               0.0018276
           0.64281
                     0.048321 3.602e-04
                                               0.0022911
3916
3917
    8km
3918
                            SD Naive SE Time-series SE
               Mean
3919
```

```
83.97039 16.508146 1.230e-01
    N
                                                 0.4548782
3920
    lam0
            0.04519
                      0.006919 5.157e-05
                                                 0.0001738
3921
    p0
                      0.006319 4.710e-05
                                                 0.0001589
            0.04319
3922
            0.28146
                      0.060653 4.521e-04
                                                 0.0016555
    psi
                      0.040989 3.055e-04
            0.66956
                                                 0.0015070
3924
```

#### 6.9.3 SCR models as multi-state models

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While we invoke a discrete state-space artificially, by gridding the underlying continuous state-space, sometimes the state-space is more naturally discrete. Consider a situation in which discrete patches of habitat are searched using some method and it might be convenient (or occur inadvertently) to associate samples to the patch level instead of recording observation locations. In this case we might use a model  $\mathbf{s}_i \sim dcat(probs[])$  where probs[] are the probabilities that an individual inhabits a particular patch. We consider such a case study in chapter XXPoissonXXX from Mollet et al. (2012) who obtained a population size estimate of a large grouse species known as the capracaillie. Forest patches were searched for scat which was identified to individual by DNA analysis. Even when space is not naturally discrete, measurements are often made at a fairly coarse grain (e.g., meters or tens of meters along a stream), or associated with spatial quadrats for scat searches and therefore the state-space may be effectively discrete in many situations.

This discrete formulation of SCR models suggests that SCR models are related to ordinary multi-state models (Kery and Schaub, 2011, ch. 9) which are also parameterized in terms of a discrete state variable which is often defined as a spatially-indexed state related either to location of capture or breeding location. While many multi-state models exist in which the state variable is not related to space, multi-state models have been extremely useful in development models of movements among geographic states and indeed this type of problem motivated their early developments by Arnason (1973, 1974) and Hestbeck (1991). We pursue this connection a little bit more in chapter XXX XYZ.

## 6.10 SUMMARY AND OUTLOOK

A point we tried to emphasize in this chapter is that the basic SCR model is not much more than an ordinary capture-recapture model for closed populations – it is simply that model but augmented with a set of "individual effects",  $\mathbf{s}_i$ , which relate encounter probability to some sense of individual location. SCR models are therefore a type of individual covariate model (as introduced in chapter 11 – but with imperfect information about the individual covariate. In other words, they are GLMM type models when N is known or, when N is unknown, they are zero-inflated GLMMs (see Royle (2006)). Another class of capture-recapture models that SCR models are closely related to is so-called "Model  $M_h$ ." The effect of introducing a spatial location for individuals is that it induces heterogeneity in

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detection probability, as in Model  $M_h$ . However, unlike Model  $M_h$ , we obtain some information about the individual effect which is completely latent in Model  $M_h$ . If the state-space of the random effect s is discrete then the SCR model resembles more closely the finite-mixture class of heterogeneity models (Norris III and Pollock, 1996) which parameterizes heterogeneity by assuming that individuals belong to discrete classes or groups (e.g., high, medium, low). In the context of SCR models we obtain some information about the "group membership" in the locations where individuals are captured. Given the direct relationship of SCR models with so many standard classes of models, we find that they are really quite easy to analyze using standard MCMC methods encased in black boxes such as **WinBUGS** or **JAGS** and possibly other packages. They are also easy to analyze using classical likelihood methods, which we address in chapter 9.

Formal consideration of the collection of individual locations  $(\mathbf{s}_1, \dots, \mathbf{s}_N)$  in the model is fundamental to all of the models considered in this book. In statistical terminology, we think of the collection of points  $\{\mathbf{s}_i\}$  as a realization of a point process and part of the promise, and ongoing challenge, of SCR models is to develop models that reflect interesting biological processes, for example interactions among points or temporal dynamics in point locations. Here we considered the simplest possible point process model - the points are independent and uniformly ("randomly") distributed over space. Despite the simplicity of this assumption, it should suffice in many applications of SCR models although we do address generalizations of this model in later chapters. Moreover, even though the prior distribution on the point locations is uniform, the realized pattern may deviate markedly from uniformity as the observed encounter data provide information to impart deviations from uniformity. Thus, the estimated density map will typically appear distinctly non-uniform. As a general rule, information in the data will govern estimates of individual point locations so even fairly complex patterns of non-independence or non-uniformity will appear in the data. That is, we find in applications of the basic SCR model that this simple a priori model can effectively reflect or adapt to complex realizations of the underlying point process. For example, if individuals are highly territorial then the data should indicate this in the form of individuals not being encountered in the same trap - the resulting posterior distribution of point locations should therefore reflect non-independence. Obviously the complexity of posterior estimates of the point pattern will depend on the quantity of data, both number of individuals and captures per individual. Because the point process is such an integral component of SCR models, the state-space of the point process plays an important role in developing SCR models. As we tried to emphasize in this chapter, the choice of the stat-espace is part of the model. It can have an influence on parameter estimates and other inferences such as model selection (see chapter 12). We emphasize however that this is not an arbitrary decision like "buffering" because the model induces an explicit interpretation of parameters and statistical effect on estimators.

We showed how to conduct inference about the underlying point process includ-

ing calculation of density maps from posterior output. We can do other things we normally do with spatial point processes such as compute "K-functions" and test for "complete spatial randomness" (CSR) which we develop in chapter 12. Modifying and applying point process methods to SCR problems seems to us to be a fruitful area of research.

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An obvious question that might be floating around in your mind is why should we ever go through all of this trouble when we could just use MARK or CAP-**TURE** to get an estimate of N and apply 1/2 MMDM methods? The main reason is that these conventional methods are predicated on models that represent explicit misspecifications of both the observation and ecological process - they are wrong! Not just wrong, because of course all models are wrong, but they're not even plausible models! Thus while we might be able to show adequate fit or whatever, we think as a conceptual and philosophical model one should not be using models that are not even plausible data-generating models – even if the plausible ones don't fit! Perhaps more charitably, these ordinary non-spatial models are models of the wrong system. They do not account for trap identity. They don't account for spatial organization or "clustering" of individual encounters in space. And, "density" is not a parameter of those models because density has no meaning absent an explicit representation of space. If we do define space explicitly, e.g., as a buffered minimum convex hull, then the normal models  $(M_0, M_h, \text{ etc.})$  assume that individual capture-probability is not related to space, no matter how we define the buffer. Conversely, the SCR model is a model for trap-specific encounter data how individuals are organized in space and interact with traps. SCR models provide a coherent framework for inference about density or population size and also, because of the formality of their derivation, can be extended and generalized to a large variety of different situations, as we demonstrate in subsequent chapters.

In the next few chapters we continue to work with this basic SCR design and model but consider some important extensions of the basic model. For example, we consider extensions to include covariates that vary by individual, trap, or over time (chapter 13), spatial covariates on density (chapter 15), open populations (chapter 16), model assessment and selection (chapter 12) and other topics. We also consider technical details of Bayesian (chapter 10) and maximum likelihood (chapter 9) estimation so that the interested reader can develop or extend their own methods to suit their needs.

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1027	OTHER OBSERVATION MODELS

4039 MAXIMUM LIKELIHOOD ESTIMATION

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## LIKELIHOOD ANALYSIS OF SCR MODELS

In this book we mainly focus on Bayesian analysis of spatial capture-recapture models. And, in the previous chapters we learned how to fit some basic spatial capture-recapture models using a Bayesian formulation of the models analyzed in BUGS engines including **WinBUGS** and **JAGS**. Despite our focus on Bayesian analysis, it is instructive to develop the basic conceptual and methodological ideas behind classical analysis based on likelihood methods and frequentist inference. In fact, simple SCR models can be analyzed fairly easily using such methods. This has been the approach taken by Borchers and Efford (2008); Dawson and Efford (2009) and related papers.

This chapter provides some conceptual and technical footing for likelihood-based analysis of spatial capture-recapture models. We recognized earlier (chapt. 4) that SCR models are versions of binomial (or other) GLMs, but with random effects i.e., GLMMs. These models are routinely analyzed by likelihood methods. In particular, likelihood analysis is based on the integrated likelihood in which the random effects are removed by integration from the likelihood. In SCR models, the random effect, s, i.e., the 2-dimensional coordinate, is a bivariate random effect. In this chapter, we show that it is straightforward to compute the maximum likelihood estimates (MLE) for SCR models by integrated likelihood. We develop the MLE framework using R, and we also provide a basic introduction to an R package secr (Efford, 2011) which is based on the stand-alone package **DENSITY** (Efford et al., 2004). To set the context we analyze the SCR model here when N is known because, in that case, it is precisely a GLMM and does not pose any difficulty at all. We generalize the model to allow for unknown N using both conventional ideas based on the "joint likelihood" (e.g., Borchers et al., 2002) and also using a formulation based on data augmentation. We consider likelihood analysis of SCR models in the context of the wolverine camera trapping study (Magoun et al., 2011) we analyzed

in previous chapters to compare/contrast the results.

## 9.1 LIKELIHOOD ANALYSIS

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We noted in chapter 4 that, with N known, the basic SCR model is a type of binomial regression with a random effect. For such models we can easily obtain maximum likelihood estimators of model parameters based on integrated likelihood. The integrated likelihood is based on the marginal distribution of the data y in which the random effects are removed by integration. Conceptually, our model is a specification of the conditional-on-s model  $[y|s,\theta]$  and we have a "prior distribution" for s, say [s], and the marginal distribution of the data y is

$$[y|\theta] = \int_{\mathbf{s}} [y|\mathbf{s}, \theta][\mathbf{s}] d\mathbf{s}.$$

When viewed as a function of  $\theta$  for pursposes of estimation, the marginal distribution  $[y|\theta]$  is often referred to as the *integrated likelihood*.

It is worth analyzing the simplest SCR model with known-N in order to understand the underlying mechanics and basic concepts. These are directly relevant to the manner in which many capture-recapture models are classically analyzed, such as model Mh, and individual covariate models (see chapt. 6 from Royle and Dorazio (2008)). To develop integrated likelihood for SCR models, we first identify the conditional likelihood.

The observation model for each encounter observation  $y_{ij}$ , specified conditional on  $\mathbf{s}_i$ , is

$$y_{ij}|\mathbf{s}_i \sim \text{Bin}(K, p_{\theta}(\mathbf{x}_i, \mathbf{s}_i))$$
 (9.1.1)

where we have indicated the dependence of  $p_{ij}$  on  $\mathbf{s}$  and parameters  $\theta$  explicitly. For the random effect we have  $\mathbf{s}_i \sim \text{Unif}(\mathcal{S})$ . The joint distribution of the data for individual i is the product of J such terms (i.e., contributions from each of J traps).

$$[\mathbf{y}_i|\mathbf{s}_i,\theta] = \prod_j \operatorname{Bin}(K,p_{\theta}(\mathbf{x}_j,\mathbf{s}_i))$$

We note that this assumes that encounter of individual i in each trap is independent of encounter in every other trap, conditional on  $\mathbf{s}_i$ , this is the fundamental property of SCR0 or "multi-catch" traps.

The so-called "marginal likelihood" is computed by removing  $\mathbf{s}_i$ , by integration, from the conditional-on-s likelihood. That is, we compute:

$$[y|\theta] = \int_{\mathcal{S}} \mathcal{L}(\theta|\mathbf{y}_i|\mathbf{s}_i)g(\mathbf{s}_i)d\mathbf{s}_i$$

here  $g(s) = 1/||\mathcal{S}||$ .

The joint likelihood for all N individuals, assuming independence of encounters among individuals, is the product of N such terms:

$$\prod_{i} f(y[i,j])$$

We emphasize that two independence assumptions are explicit in this development: independence of trap-specific encounters within individuals and also independence among individuals. In particular, this would only be valid when individuals are not physically restrained or removed upon capture, and when traps do not "fill up".

The key operation for computing the likelihood is solving a 2-dimensional integration problem. There are some general purpose  $\mathbf{R}$  packages that implement a number of multi-dimensional integration routines including adapt (Genz et al., 2007) and R2cuba (Hahn et al., 2011). In practice, we won't rely on these extraneous  $\mathbf{R}$  packages but instead will use perhaps less efficient methods in which we replace the integral with a summation over an equal area mesh of points on the state-space  $\mathcal{S}$  and explicitly evaluate the integrand at each point. We invoke the rectangular rule for integration here<sup>1</sup> in which we evaluate the integrand on a regular grid of points of equal area and compute the average of the integrand over that grid of points. Let  $u=1,2,\ldots,nG$  index a grid of nG points,  $\mathbf{s}_u$ , where the area of grid cell u is A. In this case, the integrand, i.e., the marginal probability of  $y_{ij}$ , is approximated by

$$f(y_{ij}|\theta) = \frac{1}{nG} \sum_{u=1}^{nG} f(\mathbf{y}_i|\mathbf{s}_u)$$
(9.1.2)

This is a specific case of the general expression that could be used for approximating the integral for any arbitrary bivariate distribution g(u). The general case is

$$[y] = \frac{A}{nG} \sum_{u} [y|\mathbf{s}_{u}][\mathbf{s}_{u}]$$

In the present context note that  $[\mathbf{s}] = (1/A)$  and thus the grid-cell area cancels in the above expression to yield eq. 9.1.2.

Not surprisingly this the same answer we get if S were inherently discrete, having nG unique values with equal probabilities 1/nG, and we apply the Law of Total Probability directly to compute the marginal probability  $[y|\theta]$ .

## 9.1.1 Implementation (simulated data)

Here we will illustrate how to carryout this integration and optimization based on the integrated likelihood using simulated data (i.e., following that from Chapter 4). Using simSCRO.fn we simulate data for 100 individuals and a 25 trap array layed

<sup>1</sup>e.g., http://en.wikipedia.org/wiki/Rectangle\_method

out in a  $5 \times 5$  grid of unit spacing. The specific encounter model is the half-normal model. The 100 activity centers were simulated on a state-space defined by a  $8 \times 8$  square within which the trap array was centered (thus the trap array is buffered by 2 units). Therefore, the density of individuals in this system is fixed at 100/64.

In the following set of R commands we generate the data and then harvest the required data objects:

```
data<-simSCRO.fn(discardO=FALSE,sd=2013)
4134
    y<-data$Y
4135
    traplocs<-data$traplocs
4136
    nind<-nrow(y)</pre>
    X<-data$traplocs
4138
    J<-nrow(X)
    K<-data$K
4140
    X1<-data$xlim[1]
    Y1<-data$ylim[1]
4142
    Xu<-data$xlim[2]
    Yu<-data$ylim[2]
4144
    Now we need to define the integration grid, say G, which we do with the following
4145
    set of R commands (here, delta is the grid spacing):
4146
    delta<- .2
4147
    xg<-seq(X1+delta/2, Xu-delta/2, by=delta)
    yg<-seq(Yl+delta/2, Yu-delta/2, by=delta)
4149
                                  # assumes xg and yg same dimension here
    npix<-length(xg)
    area<- (Xu-X1)*(Yu-Y1)/((npix)*(npix)) # dont need area for anything
4151
    G<-cbind(rep(xg,npix),sort(rep(yg,npix)))
4152
    nG<-nrow(G)
4153
4154
```

In this case, the integration grid is set up as a grid with spacing  $\delta = 0.2$  which produces a  $40 \times 40$  grid of points for evaluating the integrand if the state-space buffer is set at 2.

We next create an  ${\bf R}$  function that defines the likelihood as a function of the data objects y and X which were created above but, in general, you would read these files into  ${\bf R}$ , e.g., from a .csv file. In addition to these data objects, we need to have defined the various quantities associated with the integration grid  ${\bf G}$  and nG. However, instead of worrying about making all of these objects and keeping track of them we just put that code above into the likelihood function and pass  $\delta$  as an additional (optional) argument and a few other things that we need such as the boundary of the state-space over which the integration (summation) is being done. Here is one reasonably useful variation of a function for estimation based on the integrated likelihood:

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```
intlik1<-function(parm,y=y,delta=.2,X=traplocs,ssbuffer=2){</pre>
4168
4169
     X1<-min(X[,1]) - ssbuffer</pre>
4170
     Xu < -max(X[,1]) + ssbuffer
4171
     Yu < -max(X[,2]) + ssbuffer
4172
     Yl<-min(X[,2]) - ssbuffer
4173
4174
     xg<-seq(X1+delta/2, Xu-delta/2,,length=npix)
4175
     yg<-seq(Y1+delta/2, Yu-delta/2,,length=npix)
4176
     npix<-length(xg)
4177
     G<-cbind(rep(xg,npix),sort(rep(yg,npix)))
4179
     nG<-nrow(G)
4180
     D<- e2dist(X,G)
4181
4182
     alpha0<-parm[1]
4183
     alpha1<-parm[2]
4184
     probcap<- plogis(alpha0)*exp(-alpha1*D*D)</pre>
4185
     Pm<-matrix(NA,nrow=nrow(probcap),ncol=ncol(probcap))</pre>
4186
                             # all zero encounter histories
4187
     n0 < -sum(apply(y,1,sum) == 0)
4188
                             # encounter histories with at least 1 detection
4189
     ymat<-y[apply(y,1,sum)>0,]
4190
     ymat<-rbind(ymat,rep(0,ncol(ymat)))</pre>
4191
4192
     lik.marg<-rep(NA,nrow(ymat))</pre>
     for(i in 1:nrow(ymat)){
4193
     Pm[1:length(Pm)] <- (dbinom(rep(ymat[i,],nG),K,probcap[1:length(Pm)],log=TRUE))</pre>
4194
     lik.cond<- exp(colSums(Pm))</pre>
4195
     lik.marg[i]<- sum( lik.cond*(1/nG))</pre>
4196
     }
4197
     nv<-c(rep(1,length(lik.marg)-1),n0)</pre>
4198
     -1*( sum(nv*log(lik.marg)) )
4199
     }
4200
```

The function accepts as input the encounter history matrix, y, the trap locations, X, and the state-space buffer. This allows us to vary the state-space buffer and easily evaluate the sensitivity of the MLE to the size of the state-space. Note that we have a peculiar handling of the encounter history matrix y. In particular, we remove the all-zero encounter histories from the matrix and tack-on a single all-zero encounter history as the last row which then gets weighted by the number of such encounter histories (n0). This is a bit long-winded and strictly unnecessary when N is known, but we did it this way because the extension to the unknown-N case is now transparent (as we demonstrate in the following section). The matrix Pm holds the log-likelihood contributions of each encounter frequency for each possible state-space location of the individual. The log contributions are summed up and the result exponentiated on the next line, producing lik.cond, the conditional-on-

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s likelihood (Eq. 9.1.1 above). The marginal likelihood ( $\mathtt{lik.marg}$ ) sums up the conditional elements weighted by  $\Pr(\mathbf{s})$  (formula XXX above). Finally, this function assumes that K, the number of replicates, is constant for each trap. Further, it assumes that the state-space is a square. As an exercise, consider resolving these two issues by generalizing the code.

Here is the **R** command for maximizing the likelihood and saving the results into an object called frog. The output is a list of the following structure and these specific estimates are produced using the simulated data set:

```
# should take 15-30 seconds
4221
4222
    > starting.values <- c(-2, 2)
4223
     > frog<-nlm(intlik1, starting.values, y=y, delta=.1, X=traplocs, ssbuffer=2, hessian=TRUE)
4224
4225
     $minimum
4227
     [1] 297.1896
4228
4229
     $estimate
4230
     [1] -2.504824
                      2.373343
4231
4232
     $gradient
     [1] -2.069654e-05 1.968754e-05
4234
4235
     $hessian
4236
                 [,1]
                             [,2]
4237
     [1,] 48.67898 -19.25750
4238
     [2,] -19.25750 13.34114
4240
     $code
4241
     [1] 1
4242
4243
     $iterations
4244
4245
     [1] 11
```

Details about this output can be found on the help page for nlm. We note briefly that frog\$minimum is the negative log-likelihood value at the MLEs, which are stored in the frog\$estimate component of the list. The hessian is the observed Fisher information matrix, which can be inverted to obtain the variance-covariance matrix using the commands:

## > solve(frog\$hessian)

It is worth drawing attention to the fact that the estimates are different than the Bayesian estimates reported in the previous chapter (section XYZ)!!! How can that be?! There are several reasons for this. First Bayesian inference is based on

the posterior distribution and it is not generally the case that the MLE should correspond to any particular value of the posterior distribution. If the prior distributions in a Bayesian analysis are uniform, then the mode of the posterior is the MLE, but note that Bayesians almost always report posterior means and so there will typically be a discrepancy there. Secondly, we have implemented an approximation to the integral here and there might be a slight bit of error induced by that. We will evaluate that shortly. Third, the Bayesian analysis by MCMC is subject to some amount of Monte Carlo error which the analyst should always be aware of in practical situations. All of these different explanations are likely responsible for some of the discrepancy. Accounting for these, as a practical matter, we see general consistency between the two estimates.

To compute the integrated likelihood we used a discrete representation of the state-space so that the integral could be approximated as a summation over possible values of s with each value being weighted by its probability of occurring, which is 1/nG under the assumption that s is uniform on the state-space S. In chapter 4 we used a discrete state-space in developing a Bayesian analysis of the model in order to be able to modify the state-space in a flexible manner. Bayesian analysis requires simulation of the point process conditional on the observations, and this can be a difficult task when the state-space is continuous but has irregular geometry. Conversely, if the state-space is a regular polygon then Bayesian analysis by MCMC is possibly more efficient with a continuous state-space. We emphasize that the state-space is a part of the model. In some cases there wont be a natural choice of state space beyond "some large rectangle containing the trap grid" and, in such cases, for regular detection functions the estimate of density is invariant to the size of the state-space (i.e., the buffer) as long as it is sufficiently large. However if there are good reasons to restrict the state-space, it will tend to have an influence on the likelihood and hence AIC and so forth. As an illustration, lets do that by changing the state space here.....Use my polygon clipping stuff .....

In summary, we note that, for the basic SCR model, integrated likelihood is a really easy calculation when N is known. Even for N unknown it is not too difficult, and we will do that shortly. However, if you can solve the known-N problem then you should be able to do a real analysis, for example by considering different values of N and computing the results for each value and then making a plot of the log-likelihood or AIC and choosing the value of N that produces the best log likelihood or AIC. As a homework problem we suggest that the reader take the code given above and try to estimate N without modifying the code by just repeatedly calling that code for different values of N and trying to deduce the best value. Nevertheless, we will formalize the unknown-N problem shortly. We note that the software package **DENSITY** (Efford et al., 2004) implements certain types of SCR models using integrated likelihood methods. **DENSITY** has been made into an **R** package called **secr** (Efford, 2011) and we provide an analysis of some data using **secr** shortly along with a discussion of its capabilities.

## 9.2 MLE WHEN N IS UNKNOWN

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Here we build on the previous introduction to integrated likelihood but we consider now the case in which N is unknown. We will see that adapting the analysis based on the N-known model is really straightforward for the more general problem. The main distinction is that we dont observe the all-zero encounter history so we have to make sure we compute the probability for that encounter history which we do by tacking a row of zeros onto the encounter history matrix. In addition, we include the number of such all-zero encounter histories as an unknown parameter of the model. Call that unknown quantity  $n_0$ . In addition, we have to be sure to include a combinatorial term to account for the fact that of the n observed individuals there are  $\binom{N}{n}$  ways to realize a sample of size n. The combinatorial term involves the unknown  $n_0$  and thus it must be included in the likelihood.

## DETAILS NEEDED HERE

To summarize, when N is unknown, the n observed encounter histories have a multinomial distribution with probabilities pi(i) and sample size  $N^2$ . The last cell the "zero cell" is computed by carrying out the integral in expression XYZ above for the all-zero encounter history and we have to account for the fact that there are  $n_0 = N - n$  such encounter histories.

To analyze a specific case, well read in our fake data set (simulated using the parameters given above). To set some things up in our workspace we do this:

```
data<-simSCRO.fn(discard0=TRUE,sd=2013)
4316
    y<-data$Y
4317
    nind<-nrow(y)</pre>
     X<-data$traplocs
4319
     J<-nrow(X)
4320
     K<-data$K
4321
    X1<-data$xlim[1]
     Yl<-data$vlim[1]
4323
    Xu<-data$xlim[2]</pre>
     Yu<-data$ylim[2]
4325
```

Recall that these data were generated with N=100, on an  $8\times 8$  unit state-space representing the trap locations (**X**) buffered by 2 units. As before, the likelihood is defined in the **R** workspace as an **R** function which takes an argument being the unknown parameters of the model and additional arguments as prescribed. In particular, as before, we provide the encounter history matrix **y**, the trap locations traplocs, the spacing of the integration grid ( $\delta$ ) and the state-space buffer. Here is the new likelihood function:

intlik2<-function(parm,y=y,delta=.3,X=traplocs,ssbuffer=2){</pre>

 $<sup>^2</sup>$  Maybe you could show an alternative simulation script to generate data using the rmultinom function. This would make it a little more clear for people

```
4334
    Xl<-min(X[,1]) -ssbuffer</pre>
4335
    Xu<-max(X[,1])+ ssbuffer</pre>
4336
    Yu < -max(X[,2]) + ssbuffer
    Yl<-min(X[,2])- ssbuffer
4338
4339
    #delta<- (Xu-X1)/npix</pre>
4340
    xg<-seq(X1+delta/2, Xu-delta/2, delta)
4341
    yg<-seq(Yl+delta/2, Yu-delta/2, delta)
4342
    npix.x<-length(xg)
4343
    npix.y<-length(yg)
4344
    area<- (Xu-X1)*(Yu-Y1)/((npix.x)*(npix.y))
4345
    G<-cbind(rep(xg,npix.y),sort(rep(yg,npix.x)))</pre>
4346
    nG<-nrow(G)
4347
    D<- e2dist(X,G)
4348
4349
    alpha0<-parm[1]
4350
    alpha1<-parm[2]
4351
    n0 < -exp(parm[3])
    probcap<- plogis(alpha0)*exp(-alpha1*D*D)</pre>
4353
    Pm<-matrix(NA,nrow=nrow(probcap),ncol=ncol(probcap))</pre>
    ymat<-rbind(y,rep(0,ncol(y)))</pre>
4355
4356
    lik.marg<-rep(NA,nrow(ymat))</pre>
4357
    for(i in 1:nrow(ymat)){
4358
    Pm[1:length(Pm)]<- (dbinom(rep(ymat[i,],nG),K,probcap[1:length(Pm)],log=TRUE))
4359
    lik.cond<- exp(colSums(Pm))</pre>
4360
    lik.marg[i]<- sum( lik.cond*(1/nG) )</pre>
4361
    }
4362
    nv<-c(rep(1,length(lik.marg)-1),n0)</pre>
4363
    part1<- lgamma(nrow(y)+n0+1) - lgamma(n0+1)</pre>
4364
    part2<- sum(nv*log(lik.marg))</pre>
4365
      -1*(part1+ part2)
4366
    }
4367
        To execute this function for the data that we created with simSCRO.fn, we
4368
    execute the following command (saving the result in our friend frog). This re-
4369
    sults in the usual output, including the parameter estimates, the gradient, and the
4370
    numerical Hessian which is useful for obtaining asymptotic standard errors (see
4371
    below):
4372
    > frog<-nlm(intlik2,c(-2.5,2,log(4)),hessian=TRUE,y=y,X=X,delta=.2,ssbuffer=2)</pre>
4373
    There were 50 or more warnings (use warnings() to see the first 50)
4375
```

```
4376
    > frog
4377
    $minimum
4378
    [1] 113.5004
4380
    $estimate
    [1] -2.538334
                      2.466515
                                  4.232810
4382
4383
    [. Additional output deleted .]
4384
    While this produces some R warnings, these happen to be harmless in this case,
    and we will see from the nlm output that the algorithm performed satisfactory in
4386
    minimizing the objective function. The estimate of population size for the state-
    space (using the default state-space buffer) is
4388
    > nrow(y) + exp(4.2328)
    [1] 110.9099
4390
    Which differs from the data-generating value (N = 100) as we might expect. We
    usually will present an estimate of uncertainty assocated with this MLE which we
    can obtain by inverting the Hessian. Note that Var(\hat{N}) = n + Var(\hat{n}_0). Since we
    have parameterized the model in terms of log(n_0) we use a delta approximation to
4394
    obtain the variance on the scale of n_0 as follows:
    > (exp(4.2328)^2)*solve(frog$hessian)[3,3]
4396
    [1] 260.2033
    > sqrt(260)
    [1] 16.12452
```

#### 9.2.1 Exercises

- 4401 1. Run the analysis with different state-space buffers and comment on the result.
- 2. Conduct a brief simulation study using this code by simulating 100 data sets and obtain the MLEs for each data set. Do things seem to be working as you expect?
- 3. Further extensions: It should be straightforward to generalize the integrated likelihood function to accommodate many different situations. For examples, if we want to include more covariates in the model we can just add stuff to the object probcap, and add the relevant parameters to the argument that gets passed to the main function. For the simulated data, make up a covariate by generating a Bernoulli covariate ("trap type" perhaps baited or not baited) randomly and try to modify the likelihood to accommodate that.

4. We would probably be interested in devising the integrated likelihood for the full
3-d encounter history array so that we could include temporally varying covariates.
This is not difficult but naturally will slow down the execution substantially. The
interested reader should try to expand the capabilities of this basic **R** function.

## 9.2.2 Integrated Likelihood using the model under data augmentation

Note that this likelihood analysis is based on the standard likelihood in which N(or  $n_0$ ) is an explicit parameter. This is usually called the "joint likelihood" or "unconditional likelihood". We could also express the joint likelihood using data augmentation, replacing the parameter N with  $\psi$  (e.g., Royle and Dorazio, 2008, sec. xyz). We don't go into detail here, but we do note that the likelihood under data augmentation is a zero-inflated binomial mixture precisely as an occupancy type model (Royle, 2006). The interested reader could adapt the material from Royle and Dorazio (2008) with the R code given above for the likelihood and implement the likelihood analysis based on the model under data augmentation. While we can carryout likelihood analysis of models under data augmentation, we primarily advocate data augmentation for Bayesian analysis. 

#### 9.2.3 Extensions

There are other types of covariates of interest: behavioral response, sex-specificity of parameters and all of these things. Some of these can be added directly to the likelihood if the covariate is fixed and known for all individuals captured or not. This excludes most covariates but it does include behavioral response. Sex-specificity is more difficult since sex is not known for uncaptured individuals. Trap-specific covariates such as trap type or status, or time-specific covariates such as date, are relatively easy to deal with (we leave these as exercises). We apply these various models in Chapter XXXX. To analyze such models, we do Bayesian analysis of the joint likelihood facilitated by the use of data augmentation. For covariates that are not fixed and known for all individuals, it is hard to do MLE for these based on the joint likelihood as we have developed above. Instead what people normally do is use what is colloquially referred to as the "Huggins-Alho" type model which is one of the approaches taken in the software package secr (Efford, 2011, see sec. 9.5).

# 9.3 CLASSICAL MODEL SELECTION AND ASSESSMENT

In most analyses, one is interested in choosing from among various potential models. A good thing about classical analysis based on likelihood is we can do rote application of AIC without thinking about anything. With distance as a covariate (e.g., distance sampling) this is usually applied to some arbitrary selection of distance functions. We don't recommend this. Given there is hardly ever (if at all) a

rational science-based reason for choosing some particular distance function we believe that this standard approach will invariably lead to over-fitting. The fact that AIC is easy to compute does not mean that it should be abused in such fashions. Further discussion is made in chapters XYZ.

Goodness-of-fit In many analyses based on likelihood methods it is possible to cook-up fit statistics for which asymptotic distributions are knkown. In general, however, applied statisticians tend to adopt bootstraping based on heuristically appealing fit statistics. An omnibus global GoF statistic is not so obvious but we can apply bootstrapping principles to SCR models directly which we discussion in chapter XYZ. Bayesian goodness-of-fit is almost always addressed with Bayesian p-values or some other posterior predictive check (REF XXX). Thus the approach whether Bayesian or classical is the same. We identify a fit statistic, we do a bootstrap (classical) or a Bayesian p-value. Royle et al. (2011) decomposed the fit problem into separate evaluations of the CSR hypothesis and the encounter process model. We discuss all of this in Chapter XYZ.

# 9.4 LIKELIHOOD ANALYSIS OF THE WOLVERINE CAMERA TRAPPING DATA

Here we compute the MLEs for the wolverine data using an expanded version of the function we developed in the previous section. To accommodate that each trap might be operational a variable number of nights, we provided an additional argument to the likelihood function (allowing for a vector K), which requires also a modification to the construction of the likelihood. In addition, we had to accommodate that the state-space is a general rectangle, and we included a line in the code to compute the state-space area which we apply below for computing density. The more general function (intlik3) is given in the  $\bf R$  package. It has a general purpose wrapper named  $\bf scr$  which has other capabilities too.

The data were read into our R session and manipulated using the following commands. Note that we use the utility  ${\bf R}$  function SCR23darray.fn which we defined in chapt. 4.

```
4474 > wcaps<-source("wcaps.R")$value
4475 > wtraps<-source("wtraps.R")$value
4476 > K.wolv<-apply(wtraps[,4:ncol(wtraps)],1,sum)
4477 >
4478 > xx<-SCR23darray.fn(wcaps,ntraps=37,nperiods=165)
4479 > y.wolv<- apply(xx,c(1,3),sum)
4480 > traplocs.wolv<-wtraps[,2:3]
4481 > traplocs.wolv<-traplocs.wolv/10000
4482 >
4483 > frog<-nlm(intlik3,c(-1.5,1.2,log(4)),hessian=TRUE,y=y.wolv,K=K.wolv,X=traplocs.wolv,delta=.1,s
4484 There were 23 warnings (use warnings() to see them)</pre>
```

```
> frog
4485
4486
    $minimum
4487
    [1] 220.4355
4488
4489
    $estimate
    [1] -2.817570
                      1.255112 3.599040
4491
4492
    $gradient
4493
    [1] -6.274309e-06 2.146722e-05 -1.045566e-05
4494
    $hessian
4496
                  [,1]
                               [,2]
                                           [,3]
4497
           37.687931 -11.852236
                                      4.688911
4498
    [2,] -11.852236
                        30.846144 -9.199113
4499
             4.688911
                        -9.199113 13.050428
4500
4501
    $code
4502
    [1] 1
4503
4504
    $iterations
4505
    [1] 12
4506
    > exp(3.599)*sqrt(solve(frog$hessian)[3,3])
4508
    [1] 11.41059
4509
    >
4510
4511
```

4512

4513

4515

4517

We optained the MLEs for a state-space buffer of 2 (standardized units) and for integration grid with spacing  $\delta = .3, .2, .1, .05$ . The MLEs for these 4 cases including the relative runtime are given in Table 9.1.

Table 9.1. Run time and MLEs for different integration grid resolutions.

δ	runtime	Estimates		
		$\alpha_0$	$\theta$	$log(n_0)$
0.30	8.4	-2.819786	1.258468	3.569731
0.20	22.6	-2.817610	1.254757	3.583690
0.10	99.0	-2.817570	1.255112	3.599040
0.05	403.0	-2.817559	1.255281	3.607158

We see the results change only slightly as the fineness of the integration grid increases. Conversely, the runtime on the platform of the day for the 4 cases increases rapidly which, as we have suggested before, could probably be regarded

in relative terms, across platforms, for gaging the decrease in speed as the fineness of the integration grid increases. The effect of this is that we anticipate some numerical error in approximating the integral on a mesh of points, and that error increases as the coarsenss of the mesh increases.

In section 6.9 back in chapt. 4 we used a discrete representation of the state-space in order to have control over its extent and shape, for example so that we could clip out "non-habitat". Clearly that formulation of the model is relevant to the use of integrated likelihood in the sense that such a representation of the state-space underlies the computation of the integral. Thus, for example, we could easily compute the MLE of parameters under some model with a restricted state-space merely by creating the required state-space at whatever grid resolution is desired, and then feed that state-space into the likelihood evaluation above. The R function scr which comes with the R package for this book accommodates an arbitrary state-space fashioned in this manner, as well as state-spaces created by polygons or GIS shapefiles<sup>3</sup>.

Next we studied the effect of the state-space buffer on the MLEs, using a fixed  $\delta=.2$  for all analyses. We used state-space buffers of 1 to 4 units stepped by .5. This produced the following results, given here are the state-space buffer, area of the state-space, the MLE of N for the prescribed state-space and the corresponding MLE of density:

4538		ssbuff	Ass	Nhat	Dhat
4539	[1,]	1.0	66.98212	37.73338	0.5633352
4540	[2,]	1.5	84.36242	46.21008	0.5477567
4541	[3,]	2.0	103.74272	57.00617	0.5494956
4542	[4,]	2.5	125.12302	69.03616	0.5517463
4543	[5,]	3.0	148.50332	82.17550	0.5533580
4544	[6,]	3.5	173.88362	96.44018	0.5546249
4545	[7,]	4.0	201.26392	111.83524	0.5556646

The estimates of D stabilize rapidly and the incremental difference is within the numerical error associated with approximating the integral. The results suggest that wolverine density is around 0.56 individuals per 100  $km^2$  (recall that a state-space unit is  $10 \times 10 \ km$ ). This is about 5.6 individuals per thousand  $km^2$  which compares with XYZ-lookup-XYZ reported in Royle et al. (2011c) based on a clipped state-space as described in section XYZ (XYZ chapter 4 XYZ).

## 9.4.1 Exercises

- 1. Compute the 95% confidence interval for wolverine density, somehow.
- 2. Compute the AIC of this model and modify intlik3 to consider alternative link

<sup>&</sup>lt;sup>3</sup>to be completed!

functions (at least one additional) and compare the AIC of the different models and the estimates. Comment.

#### 9.5 PROGRAM DENSITY AND THE R PACKAGE SECR

DENSITY is a software program developed by Efford (2004) for fitting spatial capture-recapture models based mostly on classical maximum likelihood estimation and related inference methods. Efford (2011) has also released an **R** package named secr, that contains many of the functions within **DENSITY** but also incorporates new models and features. Here, we will focus on secr as it will continue to be developed, contains more functionality and is based in **R**. To install and run models in secr, you must download the package and load itin **R**.

```
4564 > install.packages(secr)
4565 > library(secr)
```

secr allows the user to simulate data and fit a suite of models with various detection functions and covariate responses. secr uses the standard R model specification framework using tildes. E.g., the model command is secr.fit and is generally written as

4570 > secr.fit(capturedata, model = list(D~1, g0~1, sigma~1), buffer = 20000)

where we have  $g0^1$  indicating the intercept model. To include covariates, this would be written as  $g0^b$  where b is a behavioral response covariate. Possible predictors for detection probability include both pre-defined variables (e.g., t and b corresponding to "time" and "behavior"), and user-defined covariates of several kinds. The discussion of covariates is developed in chapter  $XX(8)^4$ 

Before we can fit the models, the data must first be entered into secr. Two input files are required: trap layout (location and identification information for each trap) and capture data (e.g., sampling session, animal identification, trap day, and trap location). SECR requires that you specify the trap type, the two most common for camera trapping/hair snares are proximity detectors and count detectors. The 'proximity' detector type allows, at most, one detection of each individual at a particular detector on any occasion. The count detector designation allows repeat encounters of each individual at a particular detector on any occasion. There are other detector types that one can select such as: 'polygon' detector type which allows for a trap to be a sampled polygon, e.g., scat surveys, and 'signal' detector which allows for traps that have a strength indicator, e.g., acoustic arrays. The detector types single and multi can be confusing as multi seems like it would appropriate for something like a camera trap, but instead these two designations refer to traps that retain individuals, thus precluding the ability for animals to be captured in other traps during the sampling occasion. The single type indicates

<sup>&</sup>lt;sup>4</sup>Beth: does secr fit a local trap-specific response or just a global behavioral response?

trap that can only catch one animal at a time, while multi indicates traps that may catch more than one animal at a time. For a full review of the detector types, one should look at the help manual, which can be accessed in R after installing the SECR package by using the command:

```
> RShowDoc("secr-manual", package = "secr")
```

As with all of the scr models, secr fits a detection function relating the probability of detection to the distance of a detector from an individual activity center. secr allows the user to specify one of a variety of detection functions including the commonly used half-normal, hazard rate, and exponential. There are 12 different functions, but some are only available for simulating data, and one should take caution when using different detection functions as the interpretation of the parameters, such as sigma, may not be consistent across formulations. The different detection functions are defined in the secr manual and can be found by calling the help function for the detection function:

#### > ?detectfn

It is useful to note that secr requires the buffer distance to be defined in meters and density will be returned as number of animals per hectare. Thus to make comparisons between secr and other models, we will often have to convert the density to the same units. Also, note that sigma is returned in units of meters.

# 9.5.1 Analysis using the secr package

To demonstrate the use of the secr package, we will show how to do the same analysis on the wolverine study as shown in section 4.6. To use the secr package, the data need to be formatted in a similar but slightly different manner than we use in WinBUGS. After installing the secr package, we first have to read in the trap locations and other related information, such as if the trap is operational during a sampling occasion. The secr package reads in the trap data through a command called read traps, which requires the detector type as input. The detector type is important because it will determine the likelihood that secr will use to fit the model. Here, we have selected proximity since individuals are captured at most once in each trap during each sampling occasion.

```
> traps= read.csv(wtraps.csv)
> colnames(traps)[1:3]<- c("trapID","x", "y") #name the first 3 columns
| to match the secr nomenclature</pre>
```

 $<sup>^5</sup>$ One question: SECR only ever reports sigma. What exactly is sigma? It is a scale parameter of a detection function and all detection functions have a scale parameter. But in what sense is this sigma parameter related to home range diameter? Efford doesnt explain this, does he? In some sections in chapter 4 or possibly 6 we get into this issue.

4625

```
> trapfile <- read.traps(data = traps, detector = "proximity")</pre>
4626
       After reading in the data, we now need to create the encounter matrix or array.
4627
    The secr package does this through the use of the make.capthist command, where
4628
    we provide the capture histories in raw data format (each line contains the session,
    identification number, occasion, and trap id for only 1 individual). This is the
4630
    format that was shown in the data input file wcaps, and we only need a line or
4631
    two to organize the data into the order that the make capthist command wants. In
4632
    creating the capture history, we provide also the trapfile with the trap information,
4633
    and the format (e.g., here fmt= trapID) so that secr knows how to match the
4634
    encounters to the trap, and finally, we provide the number of occasions.
4635
    > wolv.dat <- wcaps[,c(2, 3, 1)]
4636
                 #NEED TO UPDATE THIS WHEN I GET THE FILES,
4637
                 ### I JUST GUESSED AT THE CODE, BUT WOULD LIKE TO TRY IT.
4638
    > wolv.dat <- cbind(rep(1, dim(wolv.dat)[1], wolv.dat)
4639
    > colnames(wolv.dat) <- c("Session", "ID", "Occasion", "trapID")</pre>
4640
4641
    > wolvcapt=make.capthist(wolv.dat, trapfile, fmt = "trapID", noccasions = 165)
4642
       Calling the secr.fit command, will run the model. We are using the basic model
4643
    (SCR0), so we do not need to make any specifications in the command line except
4644
    for the providing the buffer size (in m). To specify different models, you can change
4645
    the default D~1, g0~1, sigma~1, which the interested reader can do with very little
4646
    difficulty.
4647
    > wolv.secr=secr.fit(wolvcapt, model = list(D~1, g0~1, sigma~1), buffer = 20000)
4648
4649
    > wolv.secr
4650
4651
    secr.fit( capthist = wolvcapt, buffer = 20000, binomN = 1 )
4652
    secr 2.0.0, 18:26:39 05 Jul 2011
4653
4654
    Detector type
                         proximity
4655
    Detector number
                         37
4656
    Average spacing
                         4415.693 m
4657
                         593498 652294 m
    x-range
4658
                         6296796 6361803 m
    y-range
                          21
    N animals
4660
    N detections
                          115
    N occasions
                          165
4662
    Mask area
                          1037069 ha
4664
```

```
D~1 g0~1 sigma~1
    Model
4665
    Fixed (real)
                         none
    Detection fn
                         halfnormal
4667
    Distribution
                         poisson
    N parameters
                      :
                         3
4669
    Log likelihood
                         -746.754
                         1499.508
    AIC
4671
    AICc
                         1500.920
4672
4673
    Beta parameters (coefficients)
4674
                beta
                         SE.beta
                                         lcl
                                                     ucl
4675
           -9.749576 0.23027860 -10.200913 -9.298238
4676
           -4.275736 0.15846104
                                   -4.586313 -3.965158
4677
    sigma 8.699202 0.07868944
                                    8.544973
                                               8.853430
4678
    Variance-covariance matrix of beta parameters
4680
                       D
                                    g0
4681
                                               sigma
            0.053028233
                          0.000546922 -0.005226926
4682
    g0
            0.000546922
                          0.025109900 -0.005885213
    sigma -0.005226926 -0.005885213 0.006192027
4684
4685
    Fitted (real) parameters evaluated at base levels of covariates
4686
            link
                      estimate
                                SE.estimate
                                                        lcl
             log 5.831941e-05 1.360973e-05 3.713638e-05 9.158548e-05
    D
4688
          logit 1.371121e-02 2.142902e-03 1.008756e-02 1.861207e-02
    g0
4689
             log 5.998124e+03 4.727205e+02 5.140849e+03 6.998355e+03
    sigma
4690
```

Under the fitted (real) parameters, we find D, the density, given in units of individuals/hectare (1 hectare = 100 m2). To convert this into individuals/1000km2, we multiply by 100000, thus our density estimate is 5.83 individuals/1000 km2. Sigma is given in units of meters, to convert to kilometers, we divide by 1000, which puts sigma at 5.99 km. Both of these estimates are very similar to those provided in section 4.6 for the buffer size equal to 20 km. As an exercise, run this analysis for 30 and 40 km buffers and compare those found in section 4.6 under **Winbugs**. NOTE: The function secr.fit will return a warning when the buffer size appears to be too small. This is useful particularly with the different units being used between programs and packages.

#### 9.6 SUMMARY AND OUTLOOK

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In this chapter, we showed that classical analysis of SCR models based on likelihood methods is a relatively simple proposition. Analysis is based on the so-called integrated likelihood in which the individual activity centers (random effects) are

removed from the conditional-on-s likelihood by integration. We showed how to construct the integrated likelihood and fit some simple models in the R programming language. In addition, likelihood analysis for some broad classes of SCR models can be accomplished in the software package DENSITY or in the equivalent R library secr which we provided an illustration of here. In later chapters we provide more detailed analyses of SCR data using the secr package.

To compute the integrated likelihood we have to precisely describe the state-space of the underlying point process. In practice, this leads to a buffer around the trap array. We note that this is not really a buffer strip in the sense of Wilson et al. (XYZ) which is a feature of the analysis but it is somewhat more general here. In particular, it establishes the support of the integrand which we generally require to compute any integral. It might be that the integrand itself is finite even if the support is infinity but that may or may not be the case depending on the choice of detection function. As a practical matter then, it will typically be the case that, while estimates of N increase with the size of the buffer, estimates of density stabilize. This is not a feature of the classical methods based on using model M0 or model Mh and buffering the trap array.

Why or why not use likelihood inference exclusively? For certain specific models, it is probably more computationally efficient to produce MLEs. However, Win-**BUGS** is extremely flexible in terms of describing models, although it sometimes can be quite slow. We can devise models in WinBUGS easily that we cannot fit in secr. E.g., random individual effects of various types (see next chapter), we can handle missing covariates in complete generality and seamlessly, and impose arbitrary distributions on random variables. Moreover, models can easily be adapted to include auxiliary data types. For example, we might have camera trapping and genetic data and we can describe the models directly in WinBUGS and fit a joint model. For the MLE we have to write a custom new piece of code for each model or hope someone has done it for us. Later we consider open population models which are straightforward to develop in WinBUGS but, so far, there is no available platform for doing MLE although we imagine one could develop this. Another thing that is more conceptual here is non-CSR point processes (see chapter XYZ) and generating predictions of how many individuals have home range centers in any particular polygon. Basic benefits of Bayesian analysis have been discussed elsewhere (Chapter 2? BPA book? Link and Barker?) and we believe these are compelling. On the other hand, likelihood analysis makes it easy to do model-selection by AIC. Goodness-of-fit is probably no more difficult or easy under either paradigm (see next chapter?).

In summary, basic SCR models are easy to implement by either likelihood or Bayesian methods but we feel that the typical user will realize much more flexibility in model development using existing platforms for Bayesian analysis. While these tend to be slow (sometimes excruitatingly slow), this will probably not be an impediment in most problems, especially at some near point in the future. Since we spent a lot of time here talking about specific technical details on how to implement

likelihood analysis of SCR models, we provided a corresponding treatment in the next chapter on how to devise MCMC algorithms for SCR models. This is a bit more tedious and requires more coding, but is not technically challenging (exept perhaps to develop highly efficient algorithms which we dont excel at).

9751 9752	10
1752	MCMC DETAILS

# 

# MCMC DETAILS

#### 11.1 INTRODUCTION

In this chapter we will dive a little deeper into Markov chain Monte Carlo (MCMC) sampling. We will construct custom MCMC samplers in R, starting with easy-to-code GLMs and GLMMs and moving on to simple SCR models. We will also demonstrate some tricks and simple extensions to the 'spatial null model'. Finally, we will illustrate some alternative ready-to-use software packages for MCMC sam-pling. We will NOT provide exhaustive background information on the theory and justification of MCMC sampling there are entire books dedicated to that subject and we refer you to Robert and Casella (2004) and Robert and Casella (2010). Rather we aim to provide you with enough background and technical know-how to start building your own MCMC samplers for SCR models in R. 

### 11.1.1 Why build your own MCMC algorithm?

The standard program we have used so far to run MCMC analyses is WinBUGS (Gilks et al., 1994). The wonderful thing about WinBUGS is that it will automatically use the most appropriate and efficient form of MCMC sampling for the model specified by the user.

The fact that we have such a Swiss Army knife type of MCMC machine begs the question: Why would anyone want to build their own MCMC algorithm? For one, there are a limited number of distributions and functions implemented in WinBUGS. While OpenBUGS provides more options, some more complex models may be impossible to build within these programs. A very simple example from spatial capture-recapture that can give you a headache in WinBUGS is when your state-space is an irregular-shaped polygon, rather than an ideal rectangle that can be characterized by four pairs of coordinates. It is easy to restrict activity centers

to any arbitrary polygon in R using an ESRI shapefile (and we will show you an example in a little bit), but you cannot use a shape file in a BUGS model.

Sometimes implementing an MCMC algorithm in R may be faster than in Win-BUGS - especially if you want to run simulation studies where you have hundreds or more simulated data sets, several years' worth of data or other large models, this can be a big advantage.

Finally, building your own MCMC algorithm is a great exercise to understand how MCMC sampling works. So while using the BUGS language requires you to understand the structure of your model, building an MCMC algorithm requires you to think about the relationship between your data, priors and posteriors, and how these can be efficiently analyzed and characterized. Not to mention that, if you are an R junkie, it can actually be fun. However, if you don't think you will ever sit down and write your own MCMC sampler, consider skipping this chapter - apart from coding it will not cover anything SCR-related that is not covered by other, more model-oriented chapters as well.

#### 11.2 MCMC AND POSTERIOR DISTRIBUTIONS

As mentioned in Chapter 2, MCMC is a class of simulation methods for drawing (correlated) random numbers from a target distribution, which in Bayesian inference is the posterior distribution. As a reminder, the posterior distribution is a probability distribution for an unknown parameter, say  $\theta$ , given a set of observed data and its prior probability distribution (the probability distribution we assign to a parameter before we observe data). The great benefit of computing the posterior distribution of  $\theta$  is that it can be used to make probability statements about  $\theta$ , such as the probability that  $\theta$  is equal to some value, or the probability that  $\theta$  falls within some range of values. As an example, suppose we conducted a Bayesian analysis to estimate detection probability of some species at a study site (p), and we obtained a posterior distribution of beta(20,10) for the parameter p. The following R commands demonstrate how we make inferences based upon summaries of the posterior distribution. Fig 1 shows the posterior along with the summary statistics.

```
4809 > (post.median <- qbeta(0.5, 20, 10))

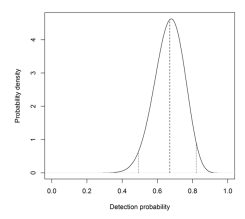
4810 [1] 0.6704151

4811 > (post.95ci <- qbeta(c(0.025, 0.975), 20, 10))

4812 [1] 0.4916766 0.8206164
```

Thus, we can state that there is a 95% probability that  $\theta$  lies between 0.49 and 0.82.

The posterior distribution summarizes all we know about a parameter and thus, is the central object of interest in Bayesian analysis. Unfortunately, in many if not most practical applications, it is nearly impossible to directly compute the posterior.



**Figure 11.1.** Probability density plot of a hypothetical posterior distribution of beta(20,10); dashed lines indicate mean and upper and lower 95% interval

Recall Bayes theorem:

$$p(theta|y) = p(y|\theta) * p(\theta)/p(y), \tag{11.2.1}$$

where  $\theta$  is the parameter of interest, y is the observed data,  $p(\theta|y)$  is the posterior,  $p(y|\theta)$  the likelihood of the data conditional on  $\theta$ ,  $p(\theta)$  the prior probability of  $\theta$ , and, finally, p(y) is the marginal probability of the data, which can also be written as

$$p(y) = \int p(y|\theta) * p(\theta)dtheta$$

This marginal probability is a normalizing constant that ensures that the posterior integrates to 1. You read in Chapter 2 that this integral is often hard or impossible to evaluate, unless you are dealing with a really simple model. For example, consider that you have a Normal model, with a set of n observations, y that come from a Normal distribution:

$$y \sim \text{Normal}(\mu, \sigma),$$

where  $\sigma$  is known and our objective is to obtain an estimate of  $\mu$  using Bayesian statistics. To fully specify the model in a Bayesian framework, we first have to define a prior distribution for  $\mu$ . Recall from Chapter 2 that for certain data models, certain priors lead to conjugacy i.e. if you choose the right prior for your

parameter, your posterior distribution will be of a known parametric form. The conjugate prior for the mean of a normal model is also a Normal distribution:

$$\mu \sim \text{Normal}(\mu_0, \sigma_0^2)$$

If  $\mu_0$  and  $\sigma_0^2$  are fixed, the posterior for  $\mu$  has the following form (for the algebraic proof, see XXX):

$$\mu|y \sim \text{Normal}(\mu_n, \sigma_n^2)$$
 (11.2.2)

4836 where

$$\mu_n = (sig^2/sig^2 + n * sig0^2) * mu0 + (n * sig0^2/sig^2 + n * sig0^2) * y - bar$$

4837 And

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$$sign^2 = sig^2 * sig0^2 / (sig^2 + n * sig0^2)$$

We can directly obtain estimates of interest from this Normal posterior distribution, such as the mean mu-hat and its variance; we do not need to apply MCMC, since we can recognize the posterior as a parametric distribution, including the normalizing constant p(y). But generally we will be interested in more complex models with several, say n, parameters. In this case, computing p(y) from Eq. 11.2.1 requires n-dimensional integration, which is can be difficult or impossible. Thus, the posterior distribution in generally only known up to a constant of proportionality:

$$p(\theta|y)proptop(y|\theta)*p(\theta)$$

The power of MCMC is that it allows us to approximate the posterior using simulation without evaluating the high dimensional integrals and to directly sample from the posterior, even when the posterior distribution is unknown! The price is that MCMC is computationally expensive. Although MCMC first appeared in the scientific literature in 1949 (Metropolis and Ulam, 1949), widespread use did not occur until the 1980s when computational power and speed increased (Gelfand and Smith, 1990). It is safe to say that the advent of practical MCMC methods is the primary reason why Bayesian inference has become so popular during the past three decades. In a nutshell, MCMC lets us generate sequential draws of  $\theta$  (the parameter(s) of interest) from distributions approximating the unknown posterior over T iterations. The distribution of the draw at t depends on the value drawn at t-1; hence, the draws from a Markov chain. <sup>1</sup> As T goes to infinity, the Markov chain converges to the desired distribution in our case the posterior distribution for  $\theta$ —y. Thus, once the Markov chain has reached its stationary distribution, the generated samples can be used to characterize the posterior distribution,  $p(\theta|y)$ , and point estimates of  $\theta$ , its standard error and confidence bounds, can be obtained directly from this approximation of the posterior. In practice, although we know that

<sup>&</sup>lt;sup>1</sup>In case you are not familiar with Markov chains, for t random samples  $\theta$  (1), ...  $\theta$  (t) from a Markov chain the distribution of  $\theta$  (t) depends only on the most recent value,  $\theta$  (t-1).

a Markov chain will eventually converge, we can only generate a limited number of samples a process that depending on the model can be quite time consuming.

Assessing whether our Markov chain has indeed converged is an important part of MCMC sampling and we will speak about some common diagnostics in Section XX.

#### 11.3 TYPES OF MCMC SAMPLING

There are several MCMC algorithms, the most popular being Gibbs sampling and Metropolis-Hastings sampling. We will be dealing with these two classes in more detail and use them to construct the MCMC algorithms for SCR models. Also, we will briefly review alternative techniques that are applicable in some situations.

#### 4871 11.3.1 Gibbs sampling

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Gibbs sampling was named after the physicist J.W. Gibbs by Geman and Geman (1984), who applied the algorithm to a Gibbs distribution <sup>2</sup>. The roots of Gibbs sampling can be traced back to work of Metropolis et al. (1953), and it is actually closely related to Metropolis sampling (see Chapter 11.5 in Gelman et al. (2004), for the link between the two samplers). We will focus on the technical aspects of this algorithm, but if you find yourself hungry for more background, Casella and George (1992) provide a more in-depth introduction to the Gibbs sampler.

In Chapter 2 you already heard about the basic principles of Gibbs sampling<sup>3</sup>. But as a refresher, let's go back to our simple example from above to understand the motivation and functioning of Gibbs sampling. Recall that for a Normal model with known variance and a Normal prior for  $\mu$ , the posterior distribution of  $\mu|y$  is also Normal. Conversely, with a fixed (known)  $\mu$ , but unknown variance, the conjugate prior for  $\sigma^2$  is an Inverse-Gamma distribution with shape and scale parameters a and b:

$$\sigma^2 \sim Iv - Gamma(a, b),$$

With fixed a and b, the posterior p(sig|mu,y) is also an Inverse Gamma distribution, namely:

$$sig|u, u \sim InvGamma(an, bn),$$
 (11.3.1)

where an = n/2 + a and  $bn = 1/2\sigma(yi - mu)^2 + b$  However, what if we know neither mu nor sig, which is probably the more common case? The joint posterior distribution of mu and sig now has the general structure

$$p(mu, sig|y) = \frac{p(y|mu) * p(mu) * p(sig)}{\int p(y|mu) * p(mu) * p(sig) dmudsig}$$

<sup>&</sup>lt;sup>2</sup>a distribution from physics we are not going to worry about, since it has no immediate connection with Gibbs sampling other than giving its name

<sup>&</sup>lt;sup>3</sup>maybe we should think out chapter 2 and concentrate that material here?

Or

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```
p(mu, sig|y) \propto p(y|mu) * p(mu) * p(sig)
```

This cannot easily be reduced to a distribution we recognize. However, we can condition mu on sig (i.e., we treat sig as fixed) and remove all terms from the joint posterior distribution that do not involve mu to construct the full conditional distribution,

$$p(mu|sig, y) \propto p(y|mu) * p(mu)$$

The full conditional of mu again takes the form of the Normal distribution shown in Eq. ??; similarly, p(sig|mu, y) takes the form of the Inverse Gamma distribution shown in Eq. Eq. 11.3.1 both distribution we can easily sample from. And this is precisely what we do when using Gibbs sampling we break down high-dimensional problems into convenient one-dimensional problems by constructing the full conditional distributions for each model parameter separately; and we sample from these full conditionals, which, if we choose conjugate priors, are known parametric distributions. Let's put the concept of Gibbs sampling into the MCMC framework of generating successive samples, using our simple Normal model with unknown mu and sig and conjugate priors as an example. These are the steps you need to build a Gibbs sampler:

**Step 0:** Begin with some initial values for  $\theta$ ,  $\theta(0)$ . In our example, we have to specify initial values for mu and sig, for example by drawing a random number from some uniform distribution, or by setting them close to what we think they might be. (Note: This step is required in any MCMC sampling chains have to start from somewhere. We will get back to these technical details a little later.)

Step 1: Draw  $\theta 1(1)$  from the conditional distribution  $p(\theta 1(1) - \theta 2(0), \theta d(0))$  Here, 4912  $\theta$ 1 is mu, which we draw from the Normal distribution in Eq. ?? using sig(0) as 4913 value for sig. 4914

Step 2: Draw  $\theta 2(1)$  from the conditional distribution  $p(\theta 2(1) - \theta 1(1), \theta 3(0), \theta d(0))$ Here,  $\theta 2$  is sig, which we draw from the Inverse Gamma distribution of Eq. 11.3.1, 4916 using mu(1) as value for mu...

**Step d:** Draw  $\theta d(1)$  from the conditional distribution  $p(\theta d(1) - \theta 1(1), ..., \theta d-1(1))$ In our example we have no additional parameters, so we only need step 0 through to 2. Repeat Steps 1 to d for K = a large number of samples. In terms of R coding, this means we have to write Gibbs updaters for mu and sig and embed them into a loop over K iterations. The final code in the form of an R function is shown in Panel 1.

Andy will build the panel environment here soon.

Panel 1: R-code for a Gibbs sampler for a Normal model with unknown mu

```
and sig and conjugate (Normal and Inverse Gamma, respectively) priors
4927
    for both parameters.
4928
4929
    Normal.Gibbs<-function(y=y,mu0=mu0, sig0=sig0, a=a,b=b,niter=niter) {
4930
4931
    ybar<-mean(y)</pre>
4932
    n<-length(y)
4933
    mu<-runif(1) #mean initial value
4934
    sig<-runif(1) #sd initial value
4935
    an<-n/2 + a
4936
4937
    out<-matrix(nrow=niter, ncol=2)</pre>
4938
    colnames(out)<-c('mu', 'sig')</pre>
4939
4940
    for (i in 1:niter) {
4941
4942
    #update mu
4943
    mun \leftarrow (sig/(sig+n*sig0))*mu0 + (n*sig0/(sig+n*sig0))*ybar
4944
    sign <- (sig*sig0)/ (sig+n*sig0)</pre>
    mu<-rnorm(1,mun, sqrt(sign))</pre>
4946
4947
    #update sig
4948
    bn < -0.5 * (sum((y-mu)^2)) + b
    sig<-1/rgamma(1,shape=an, rate=bn)</pre>
4950
    out[i,]<-c(mu,sqrt(sig))</pre>
4951
4952
    }
4953
    return(out)
4954
    }
4955
        This is it! You can use the code NormalGibbs.R in the R package scrbook to
4956
    simulate some data, y \sim \text{Normal}(5, 0.5) and run your first Gibbs sampler. Your
4957
    output will be a table with two columns, one per parameter, and K rows, one per
4958
    iteration. For this 2-parameter example you can visualize the joint posterior by
4959
    plotting samples of \mu against samples of \sigma (Fig. 2 XXX):
4960
    plot(out[,1], out[,2])
4961
    The marginal distribution of each parameter is approximated by just examining the
4962
    samples of this particular parameter you can visualize it by plotting a histogram
    of the samples (Fig. 3 a, b XXX):
4964
    par(mfrow=c(1,2))
    hist(out[,1]); hist (out[,2])
```

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Finally, recall an important characteristic of Markov chains, namely, that the chain has to have converged (reached its stationary distribution) for samples to come from the posterior distribution. In practice, that means you have to throw out some of the initial samples called the burn-in. We will talk about this in more when we talk about convergence diagnostics. For now, you can use the plot(out[,1]) or plot(out[,2]) command to make a time series plot of the samples of each parameter and visually assess how many of the initial samples you should discard. Figure 3 c and d shows plots for the estimates of mu and sigma from our simulated data set; you see that in this simple example the Markov chain apparently reaches its stationary distribution very quickly the chains look 'grassy' seemingly from the start. It is hard to discern a burn-in phase visually (but we will see examples further on where the burn-in is clearer) and you may just discard the first 500 draws to be sure you only use samples from the posterior distribution. The mean of the remaining samples are your estimates of mu and sig:

```
summary(mod[501:10000,])
                                        sig
            mu
4982
             : 4.936
                                     : 0.4569
4983
     Min.
                            Min.
     1st Qu.: 4.984
                           1st Qu.: 0.4889
4984
     Median: 4.994
                                   0.4961
                         Median :
     Mean
             :
               4.994
                          Mean
                                    0.4964
4986
                          3rd Qu.: 0.5037
     3rd Qu.: 5.005
     Max.
             : 5.062
                            Max.
                                     : 0.5356
4988
```

## 11.3.2 Metropolis-Hastings sampling

Although it is applicable to a wide range of problems, the limitations of Gibbs sampling are immediately obvious what if we do not want to use conjugate priors (or what if we cannot recognize the full conditional distribution as a parametric distribution, or simply do not want to worry about these issues)? The most general solution is to use the Metropolis-Hastings (MH) algorithm, which also goes back to the work by Metropolis et al. (1953). You saw the basics of this algorithm in Chapter 2. In a nutshell, because we do not recognize the posterior  $p(\theta|y)$  as a parametric distribution, the MH algorithm generates samples from a known proposal distribution, say  $h(\theta)$ , that depends on  $\theta$  at t-1. The  $t^{th}$  sample is accepted or rejected based on its joint posterior probability density compared to the density of the sample at t-1. The original Metropolis algorithm requires  $h(\theta)$  to be symmetric so that  $h(\theta^t|\theta^{t-1}) = h(\theta^{t-1}|\theta^t)$ ; but a later development of the algorithm by Hastings (1970) lifted this condition. Using a symmetric proposal distribution makes life a little easier and we are going to limit our coverage of the Metropolis-Hastings sampler to this specific case. Specifically, we are going to use a Normal proposal distribution, which is also referred to as 'random walk Metropolis-Hastings

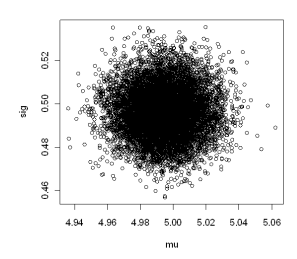


Figure 11.2. Joint posterior distribution of mu and sig from a Normal Model

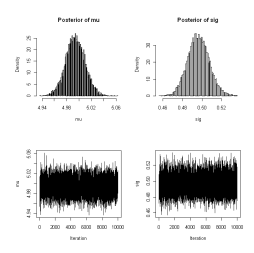


Figure 11.3. Plots of the posterior distributions of mu(a) and sig(b) from a Normal model and time series plots of mu(c) and sig(d).

sampling'. It is worth knowing that there are alternative formulations of the algorithm. For example, in the independent M-H,  $\theta^t$  does not depend on  $\theta^{t-1}$ , while the Langevin algorithm (Roberts and Rosenthal, 1998) aims at avoiding the random walk by favoring moves towards regions of higher posterior probability density. The interested reader should look up these algorithms in Robert and Casella (2004) or Robert and Casella (2010).

Building a MH sampler can be broken down into several steps. We are going to demonstrate these steps using a different but still simple and common model the logit-normal or logistic regression model. For simplicity, assume that

$$y \sim \text{Bern}(\exp(\theta)/(1 + \exp(\theta)))$$

5015 and

5012

5013

5014

$$\theta \sim \text{Normal}(\mu_0, \sigma)$$

The following steps are required to set up a random walk MH algorithm:

Step 0: Choose initial values,  $\theta(0)$ .

Step 1: Generate a proposed value of  $\theta$  at t from h(thetat—thetat-1). We often use a Normal proposal distribution, so we draw  $\theta$ 1 from  $Normal(theta0, sigh^2)$ , where  $sigh^2$  is the variance of the Normal proposal distribution, a tuning parameter that we have to set.

Step 2: Calculate the ratio of posterior densities for the proposed and the original value for  $\theta$ :

$$r = p(\theta^t|y)/p(\theta^{t-1}|y)$$

5024 In our example,

 $r = \text{Bern}(y|\theta^t) * Normal(\theta^t|\mu_0, \sigma_0) / Bernoulli(y|thetat-1) * Normal(thetat-1|mu_0, siq_0)$ 

5025 Step 3: Set

5034

5026 \begin{eqnarray\*}
5027 \$\theta\$ (t) &= & \$\theta\$ (t) \mbox{with probability min(r,1)}//
5028 & = & \$\theta\$ (t-1) \mbox{ otherwise }
5029 \end{eqnarray\*}

We can do that by drawing a random number u from a Unif(0,1) and accept  $\theta^t$  if u < r. Repeat for  $t = 1, 2, \ldots$  a large number of samples. The  $\mathbf R$  code for this MH sampler is provided in Panel 2 XXXX.

5033 Panel 2: R code to run a Metropolis sampler on a simple Logit-Normal model.

Logreg.MH<-function(y=y, mu0=mu0, sig0=sig0, niter=niter) {

```
out<-c()
5037
5038
     theta<-runif(1, -3,3) #initial value
5039
5040
     for (iter in 1:niter){
5041
     theta.cand<-rnorm(1, theta, 0.2)
5042
5043
     loglike<-sum(dbinom(y, 1, exp(theta)/(1+exp(theta)), log=TRUE))</pre>
5044
     logprior <- dnorm(theta,mu0 ,sig0, log=TRUE)</pre>
5045
     loglike.cand<-sum(dbinom(y, 1, exp(theta.cand)/(1+exp(theta.cand)), log=TRUE))
5046
     logprior.cand <- dnorm(theta.cand, mu0, sig0, log=TRUE)</pre>
5047
     if (runif(1) < exp((loglike.cand+logprior.cand)-(loglike+logprior))){</pre>
5049
     theta<-theta.cand
5050
     }
5051
     out[iter]<-theta
5052
     }
5053
    return(out)
5055
     }
5056
```

The reason we sum the logs of the likelihood and the prior, rather than multiplying the original values, is simply computational. The product of small probabilities can be numbers very close to 0, which computers do not handle well. Thus we add the logarithms, sum, and exponentiate to achieve the desired result. Similarly, in case you have forgotten some elementary math,  $x/y = \exp(\log(x) - \log(y))$ , with the latter being favored for computational reasons.

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Comparing MH sampling to Gibbs sampling, where all draws from the conditional distribution are used, in the MH algorithm we discard a portion of the candidate values, which inherently makes in less efficient than Gibbs sampling the price you pay for its increased generality. In Step 1 of the MH sampler we had to choose a variance for the Normal proposal distribution. Choice of the parameters that define our candidate distribution is also referred to as 'tuning', and it is important since adequate tuning will make your algorithm more efficient, i.e. your Markov chain will converge faster. The variance should be chosen so that (a) each step of drawing a new proposal value for  $\theta$  can cover a reasonable distance in the parameter space, as otherwise, the random walk moves too slowly; and (b) proposal values are not rejected too often, as otherwise the random walk will 'get stuck' at specific values for too long. As a rule of thumb, your candidate value should be accepted in about 40% of all cases. Acceptance rates of 20 80% are probably ok, but anything below or above may well render your algorithm inefficient (this does not mean that it will give you wrong results only that you will need more iterations to converge to the posterior distribution). In practice, tuning will require some 'trialand-error' and some common sense. Or, one can use an adaptive phase, where the tuning parameter is automatically adjusted until it reaches a user-defined accep-

tance rate, at which point the adaptive phase ends and the actual Markov chain begins. This is computationally a little more advanced. Link and Barker (2009) discuss this in more detail. It is important the samples drawn during the adaptive phase are discarded. You can easily check acceptance rates for the parameters you monitor (that are part of your output) using the rejectionRate() function of the package coda (we will talk more about this package a little later on). Do not let the term 'rejection rate' confuse you; it is simply 1 acceptance rate. There may be parameters for example, individual values of a random effect or latent variables that you do not want to save, though, and in our next example we will show you a way to monitor their acceptance rates with a few extra lines of code.

#### 11.3.3 Metropolis-within-Gibbs

One weakness of the MH sampler is that formulating the joint posterior when evaluating whether to accept or reject the candidate values for  $\theta$  becomes increasingly complex or inefficient as the number of parameters in a model increases. It is probably going to sound like MCMC sampling is too good to be true but in these cases you can simply combine MH sampling and Gibbs sampling. You can use Gibbs sampling to break down your high-dimensional parameter space into easy-to-handle one-dimensional conditional distributions and use MH sampling for these conditional distributions. Better yet if you have some conjugacy in your model, you can use the more efficient Gibbs sampling for these parameters and one-dimensional MH for all the others. You have already seen the basics of how to build both types of algorithms, so we can jump straight into an example here and build a Metropolis-within-Gibbs algorithm.

#### 11.4 GLMMS POISSON REGRESSION WITH A RANDOM EFFECT

Let's assume a model that gets us closer to the problem we ultimately want to deal with a GLMM. Here, we assume we have Poisson counts, y, from i plots in j different study sites, and we believe that the counts are influenced by some plot-specific covariate, x, but that there is also a random site effect. So our model is:

$$yij \sim Poisson(lamij)$$

$$lamij = exp(aj + b * xi)$$

Let's use Normal priors on a and b,

 $aj \sim Normal(mua, siga)$ 

5111 and

 $b \sim Normal(mub, sigb)$ 

. <sup>4</sup> Since we want to estimate the random effect in this model, we do not specify  $\mu_a$  and  $\sigma_a$ , but instead, estimate them as well, so we have to specify hyperpriors for these parameters:

```
\mu_a \sim Normal(mu0, sig0)
\sigma_a \sim InvGamma(a0, b0)
```

With the model fully specified, we can compile the full conditionals, breaking the multi-dimensional parameter space into one-dimensional components:

```
\begin{eqnarray*}
5117
   p(a1|a2,a3,aj,b,y) & propto & p(yi1|a1,b) * p(a1|mua, siga) \
5118
    & \propto & Poisson(yi1| exp(a1 + b*x[j=1])) * Normal(a1|mua, siga)
5119
    \end{eqnarray*}
5120
    \begin{eqnarray*}
5121
    p(a2|a1,a3,aj,b,y) & p(yi2|a2,b) * p(a2|mua, siga) 
5122
    & \propto & Poisson(yi2|exp(a2 + b*x[j=1])) * Normal(a2|mua, siga)
    \end{eqnarray}
5124
    and so on for all elements of a.
5125
    \begin{eqnarray*}
5126
    p(b|a,y) &\propto & p(y|a,b) * p(b) \
5127
    &\propto& Poisson(y|exp(a + b*x)) *Normal(b|mub, sigb)
5128
    \end{eqnarray*}
5129
```

Finally, we need to update the hyperparameters for a:

```
p(mua|a) \propto p(a|mua, siga) * p(mua)
p(siga|a) \propto p(a|mua, siga) * p(siga)
```

Since we assumed a to come from a Normal distribution, the choice of priors for mua Normal and siga Inverse Gamma leads to the same conjucagy we observed in our initial Normal model, so that both hyperparameters can be updated using Gibbs sampling.

Now let' build the updating steps for these full conditionals. Again, for the MH steps that update a and b we use Normal proposal distributions with standard deviations sigha and sighb.

First, we set the initial values a(0) and b(0). Then, starting with a1, we draw a1(1) from Normal (a1(0), sigha), calculate the conditional posterior density of a1(0) and a1(1) and compare their ratios,

```
r = Poisson(y(j=1)|exp(a1(1)+b*x))*Normal(a1(1)|mua, siga)/Poisson(y(j=1)|exp(a1(0)+b*x))*Normal(a1(1)|mua, siga)/Poisson(y(j=1)|exp(a1(0)+b*x))*Normal(a1(1)|exp(a1(0)+b*x))*Normal(a1(1)|exp(a1(0)+b*x))*Normal(a1(1)|exp(a1(0)+b*x))*Normal(a1(1)|exp(a1(0)+b*x))*Normal(a1(1)|exp(a1(0)+b*x))*Normal(a1(1)|exp(a1(0)+b*x))*Normal(a1(1)|exp(a1(0)+b*x))*Normal(a1(1)|exp(a1(0)+b*x))*Normal(a1(1)|exp(a1(0)+b*x))*Normal(a1(1)|exp(a1(0)+b*x))*Normal(a1(1)|exp(a1(0)+b*x))*Normal(a1(1)|exp(a1(0)+b*x))*Normal(a1(1)|exp(a1(0)+b*x))*Normal(a1(1)|exp(a1(0)+b*x))*Normal(a1(1)|exp(a1(0)+b*x))*Normal(a1(1)|exp(a1(0)+b*x))*Normal(a1(1)|exp(a1(0)+b*x))*Normal(a1(1)|exp(a1(0)+b*x))*Normal(a1(1)|exp(a1(0)+b*x))*Normal(a1(1)|exp(a1(0)+b*x))*Normal(a1(1)|exp(a1(0)+b*x))*Normal(a1(1)|exp(a1(0)+b*x))*Normal(a1(1)|exp(a1(0)+b*x))*Normal(a1(1)|exp(a1(0)+b*x))*Normal(a1(1)|exp(a1(0)+b*x))*Normal(a1(1)|exp(a1(0)+b*x))*Normal(a1(1)|exp(a1(0)+b*x))*Normal(a1(1)|exp(a1(0)+b*x))*Normal(a1(1)|exp(a1(0)+b*x))*Normal(a1(1)|exp(a1(0)+b*x))*Normal(a1(1)|exp(a1(0)+b*x))*Normal(a1(1)|exp(a1(0)+b*x))*Normal(a1(1)|exp(a1(0)+b*x))*Normal(a1(1)|exp(a1(0)+b*x))*Normal(a1(1)|exp(a1(0)+b*x))*Normal(a1(1)|exp(a1(0)+b*x))*Normal(a1(1)|exp(a1(0)+b*x))*Normal(a1(1)|exp(a1(0)+b*x))*Normal(a1(1)|exp(a1(0)+b*x)
```

and accept a1(1) with probability min(r,1). We repeat this for all a's.

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<sup>&</sup>lt;sup>4</sup>Why is b a hyperparameter?

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out <- matrix (nrow=niter, ncol=3)

for (iter in 1:niter) {

for (j in 1:lev) {

#update a

colnames(out)<-c('mua','siga','b')</pre>

#initiate counter for acceptance rate of a

a.cand<-rnorm(1, a[j], 0.1) #update intercepts a one at a time

#loop over sites

```
For b, we draw b(1) from Normal (b(0), sigbh), compare the posterior densities
5143
    of b(0) and b(1),
    r = Poisson(y|exp(a+b(1)*x))*Normal(b(1)|mub, sigb)/Poisson(y|exp(a+b(0)*x))*Normal(b(0)|mub, sigb),
    and accept b (1) with probability min(r,1).
5145
        For mua and siga, we sample directly from the full conditional distributions (Eq.
5146
    XX and Eq XX):
                              mua(1) \sim Normal(mun, sign)
    where mun = (siqa(0)/siqa(0) + na * siq0) * mu0 + (na * siq0/siqa(0) + na *
    sig0) * abar(1) and sign = siga(0) * sig0/(siga(0) + n * sig0) Here, abar is the
5149
    current mean of the vector a, which we updated before, and na is the length of
5150
    a. For siga we use siga(1) \sim InvGamma(an, bn), where an = na/2 + a0, and
5151
    bn = 1/2\Sigma(a(1) - mua(1))^2 + b0.
5152
        We repeat these steps over K iterations of the MCMC algorithm. In this example
5153
    we may not want to save each value for a, but are only interested in their mean and
5154
    standard deviation. Since these two parameters will change as soon as the value for
5155
    one element in a changes, their acceptance rates will always be close to 1 and are
5156
    not representative of how well your algorithm performs. To monitor the acceptance
    rates of parameters you do not want to save, you simply need to add a few lines
5158
    of code into your updater to see how often the individual parameters are accepted.
    The full code for the MCMC algorithm of our Poisson GLMM in Panel 3 shows
5160
    one way how to monitor acceptance of individual a's.
    Panel 3: R code for the Metropolis-within-Gibbs sampler for
5162
    a Poisson regression with random intercepts.
5163
    Pois.reg<-function(y=y,site=site,mu0=mu0,sig0=sig0,a0=a0,b0=b0,
5165
               mub=mub, sigb=sigb, niter=niter){
5166
5167
    lev<-length(unique(site))</pre>
                                     #number of sites
5168
    a<-runif(lev,-5,5) #initial values a
5169
    b<-runif(1,0,5) #initial value b
5170
    mua <-mean(a)
    siga<-sd(a)
```

```
loglike<- sum(dpois (y[site==j], exp(a[j] + b*x[site==j]), log=TRUE))</pre>
5183
     logprior<- dnorm(a[j], mua,siga, log=TRUE)</pre>
5184
     loglike.cand<- sum(dpois (y[site==j], exp(a.cand + b *x[site==j]), log=TRUE))</pre>
5185
     logprior.cand<- dnorm(a.cand, mua,siga, log=TRUE)</pre>
     if (runif(1) < exp((loglike.cand+logprior.cand) (loglike+logprior))) {</pre>
5187
     a[j] < -a.cand
5188
     aUps<-aUps+1
5189
    }
5190
    }
5191
5192
     if(iter %% 100 == 0) { #this lets you check the acceptance rate of a at every 100th iteration
5194
                   cat("
                            Acceptance rates\n")
                   cat("
                              a = ", aUps/lev, "\n")
5195
    }
5196
5197
     #update b
5198
     b.cand<-rnorm(1, b, 0.1)
5199
     avec<-rep(a, times=c(rep(10,10)))</pre>
     loglike<- sum(dpois (y, exp(avec + b*x), log=TRUE))</pre>
     logprior<- dnorm(b, mub,sigb, log=TRUE)</pre>
5202
     loglike.cand<- sum(dpois (y, exp(avec + b.cand *x), log=TRUE))</pre>
5203
     logprior.cand<- dunif(b.cand, mub,sigb, log=TRUE)</pre>
5204
     if (runif(1) < exp((loglike.cand+logprior.cand) (loglike+logprior) )) {</pre>
5206
     b<-b.cand
     }
     #update mua using Gibbs sampling
5209
     abar <- mean(a)
5210
     mun<- (siga/(siga+lev*sig0))*mu0 + (lev*sig0/(siga+lev* sig0))*abar</pre>
5211
     sign <- (siga*sig0)/ (siga+lev*sig0)</pre>
5212
     mua<-rnorm(1,mun, sqrt(sign))</pre>
5213
5214
     #update siga using Gibbs sampling
5215
     a0n<-lev/2 + a0
5216
     b0n<-0.5 * (sum((a-mua)^2)) +b0
5217
     siga<-1/rgamma(1,shape=a0n, rate=b0n)</pre>
5218
5219
     out[iter,]<-c(mua, sqrt(siga), b)</pre>
5221
5222
5223
    return(out)
5224
    }
5225
```

### 11.4.1 Rejection sampling and slice sampling

While MH and Gibbs sampling are probably the most widely applied algorithms for posterior approximation, there are other options that work under certain circumstances and may be more efficient when applicable. WinBUGS applies these algorithms and we want you to be aware that there is more out there to approximate posterior distributions than Gibbs and MH. One alternative algorithm is rejection sampling. Rejection sampling is not an MCMC method, since each draw is independent of the others. The method can be used when the posterior  $p(\theta|y)$  is not a known parametric distribution but can be expressed in closed form. Then, we can use a so-called envelope function, say,  $q(\theta)$ , that we can easily sample from, with the restriction that  $p(\theta|y) < M * q(\theta)$ . We then sample a candidate value for  $\theta$  from  $q(\theta)$ , calculate  $r = p(\theta|y)/M * q(\theta)$  and keep the sample with the probability r. M is a constant that has to be picked so that  $r \in [0,1]$ , for example by evaluating both  $p(\theta|y)$  and  $q(\theta)$  at n points and looking at their ratios. Rejection sampling only works well if  $g(\theta)$  is similar to  $p(\theta|y)$ , and packages like WinBUGS use adaptive rejection sampling (Gilks and Wild, 1992), where a complex algorithm is used to fit an adequate and efficient g(theta) based on the first few draws. Though efficient in some situations, rejection sampling does not work well with high-dimensional problems, since it becomes increasingly hard to define a reasonable envelope function. For an example of rejection sampling in the context of SCR models, see Chapter 9. Another alternative is slice sampling (Neal, 2003). In slice sampling, we sample uniformly from the area under the plot of  $p(\theta|y)$ . Considering a single univariate theta. Let's define an auxiliary variable,  $U \sim Uniform(0, p(\theta|y))$ . Then,  $\theta$  can be sampled from the vertical slice of  $p(\theta|y)$  at U (Figure 4):

o \theta|U \sim \mbox{Unif}(B),

```
where B = \theta : U < p(\theta|y)
```

Slice sampling can be applied in many situations; however, implementing an efficient slice sampling procedure can be complicated. We refer the interested reader to chapter 7 of Robert and Casella (2010) for a simple example. Both rejection sampling and slice sampling can be applied on one-dimensional conditional distributions within a Gibbs sampling setup.

#### 11.5 MCMC FOR CLOSED CAPTURE-RECAPTURE MODEL MH

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<sup>&</sup>lt;sup>5</sup>there are supposed to be equations in the caption of figure 4 but it kept causing errors

<sup>&</sup>lt;sup>6</sup>Andy could move material from chapter 3 to here.

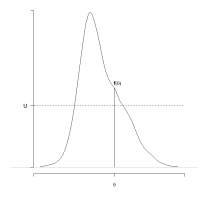


Figure 11.4. Slice sampling. For...

# 11.6 MCMC ALGORITHM FOR THE BASIC SPATIAL CAPTURE-RECAPTURE MODEL

By now you have seen how to build MCMC algorithms for some basic generalized linear models. Now, we'll walk you through the steps of building your own MCMC sampler for the basic SCR model (i.e. without any individual, site or time specific covariates) with both a Poisson and a binomial encounter process. As usual, we will have to go through two general steps before we write the MCMC algorithm:

- (1) Identify your model with all its components (including priors)
- (2) Recognize and express the full conditional distributions for all parameters

It is worthwhile to go through all of step 1 for an SCR model, but you have probably seen enough of step 2 in our previous examples to get the essence of how to express a full conditional distribution. Therefore, we will exemplify step 2 for some parameters and tie these examples directly to the respective R code.

#### Step 1 Identify your model

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Recall the components of the basic SCR model with a Poisson encounter process from Chapter 3: We assume that individuals i, or rather, their activity centers si, are uniformly distributed across our state space S,

$$si \sim U(S)$$

and that the number of times individual i encounters trap j, yij, is a random Poisson variable with mean lamij,

$$yij \sim Poisson(lamij)$$

The tie between individual location, movement and trap encounter rates is made by the assumption that lamij, is a decreasing function of the distance between si

and j, Dij, of the half-normal form

$$Lamij = lam0 * exp(-Dij2/2 * sig2),$$

where lam0 is the baseline trap encounter rate at Dij = 0 and sig controls the shape of the half-normal function.

In order to estimate the number of si in S, N, we use data augmentation (sect. 3.XYZ) and create M-n all-0 encounter histories, where n is the number of individuals we observed and M is an arbitrary number that is larger than N. We estimate N by summing over the auxiliary data augmentation variables, zi, which is 1 if the individual is part of the population and 0 if not, and assume that zi is a random Bernoulli variable,

$$z_i \sim \mathrm{Bern}(\psi)$$

To link the two model components, we modify our trap encounter model to

$$Lamij = lam0 * exp(-Dij2/2 * sig2) * zi.$$

The model has the following structural parameters, for which we need to specify priors  $\psi$  the Uniform (0,1) is required as part of the data augmentation procedure and in general is a natural choice of an uninformative prior for a probability; note that this is equivalent to a Beta(1,1) prior, which will come in handy later.  $s_i$  since si is a pair of coordinates it is two-dimensional and we use a uniform prior limited by the extent of our state-space over both dimensions.  $\sigma$  we can conceive several priors for sigma but let's assume an improper prior one that is Uniform over (-Inf, Inf). We will see why this is convenient when we construct the full conditionals for sigma.  $\lambda_0$  analogous, we will use a Uniform (-Inf, Inf) improper prior for sigma. The parameter that is the objective of our modeling, N, is a derived parameter that we can simply obtain by summing all z's:

$$N = sum(z)$$

Step 2 - Construct the full conditionals Having completed step 1, let's look at the full conditional distributions for some of these parameters. We find that with improper priors, full conditionals are proportional only to the likelihood of the observations; for example, take the movement parameter sigma:

$$Sig|s, lam0, z, ypropto[y|s, lam0, z, sig] * [sig]$$

Since the improper prior implies that [sig] propto 1, we can reduce this further to

$$Sig|s, lam0, z, ypropto[y|s, lam0, z, sig]$$

The R code to update sigma is shown in Panel 4. <sup>7</sup>

 $<sup>^7</sup>$  Somewhere in chapter 2 i added a comment about rejecting parameters outside of the parameter space as being an ok thing to do. Richard said he read something in Robert and Casellas book on that. Hopefully he can remember where and we can cite it back in Ch 2 and again here. It could be mentioned in a sentence or two up in the MCMC section.

```
Panel 4: R code to update sigma within an MCMC algorithm for
5305
     an SCR model when using an improper prior
5306
5307
5308
     sig.cand <- rnorm(1, sigma, 0.1) #draw candidate value
5309
      if(sig.cand>0){
                          #automatically reject sig.cand that are <0
5310
          lam.cand <- lam0*exp(-(D*D)/(2*sig.cand*sig.cand))</pre>
5311
          11<- sum(dpois(y, lam*z, log=TRUE))</pre>
5312
          llcand <- sum(dpois(y, lam.cand*z, log=TRUE))</pre>
5313
          if(runif(1) < exp(llcand - ll)){
5314
               11<-11cand
5315
5316
               lam<-lam.cand
               sigma<-sig.cand
5317
           }
5318
       }
5319
5320
```

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These steps are analogous for lam0 and si and we will use MH steps for all of these parameters. Similar to the random intercepts in our Poisson GLMM, we update each si individually. Note that to be fully correct, the full conditional for si contains both the likelihood and prior component, since we did not specify an improper, but a Uniform prior on si. However, with a Uniform distribution the probability density of any value is 1/(upper limit lower limit) = constant. Thus, the prior components are identical for both the current and the candidate value and can be ignored (formally, when you calculate the ratio of posterior densities, r, the identical prior component appears both in the numerator and denominator, so that they cancel each other out).

We still have to update zi. The full conditional for zi is

```
zi|y, sigma, lam0, spropto[y|z, sigma, lam0, s] * [zi]
```

and since  $zi \sim Bernoulli(psi)$ , the term has to be taken into account when updating zi. The R code for updating zi is shown in Panel 5.

```
Panel 5: R code to update z
5334
5335
              zUps <- 0 #set counter to monitor acceptance rate
5336
              for(i in 1:M) {
5337
                   if(seen[i]) #no need to update seen individuals, since their z =1
5338
                       next.
5339
                  zcand \leftarrow ifelse(z[i]==0, 1, 0)
                  11z <- sum(dpois(y[i,],lam[i,]*z[i], log=TRUE))</pre>
                  llcand <- sum(dpois(y[i,], lam[i,]*zcand, log=TRUE))</pre>
5342
5343
                  prior <- dbinom(z[i], 1, psi, log=TRUE)</pre>
5344
                  prior.cand <- dbinom(zcand, 1, psi, log=TRUE)</pre>
5345
```

```
if(runif(1) < exp( (llcand+prior.cand) - (llz+prior) )) {
    z[i] <- zcand
    zUps <- zUps+1
}
</pre>
```

 $\psi$  itself is a hyperparameter of the model, with an uninformative prior distribution of Unif(0,1) or Beta(1,1), so that

$$Psi|z \propto [z|psi] * Beta(1,1)$$

The Beta distribution is the conjugate prior to the Binomial and Bernoulli distributions (remember that  $z \sim Bernoulli(psi)$ ). The general form of a full conditional of a Beta-Binomial model with  $yi \sim Bernoulli(p)$  and  $p \sim Beta(a, b)$  is

$$p(p|y) \propto Beta(a + sum(yi), b + n - sum(yi)))$$

5356 In our case, this means we update psi as follows:

```
si<-rbeta(1, 1+sum(z), 1 + M-sum(z))
```

5351

These are all the building blocks you need to write the MCMC algorithm for the spatial null model with a Poisson encounter process. You can find the full R code (SCR0pois.R) in the online supplementary material.

#### 11.6.1 SCR model with binomial encounter process

The equivalent SCR model with a binomial encounter process is very similar. Here, each individual i can only be detected once at any given trap j during a sampling occasion k. Thus

$$yij \sim Binomial(pij, K)$$

Where  $p_{ij}$  is some function of distance between  $\mathbf{s}_i$  and trap location  $\mathbf{x}_j$ . Here we use:

$$pij = 1 - exp(-lamij)$$

Recall from Chapter 2 that this is the complementary log-log (cloglog) link function, which constrains pij to fall between 0 and 1. For our MCMC algorithm that means that, instead of using a Poisson likelihood, Poisson(y|sigma,lam0,s,z), we use a Binomial likelihood, Binomial(y,K|sigma,lam0,s,z), in all the conditional distributions. As an example, Panel 6 shows the updating step for lam0 under a binomial encounter model. The full MCMC code for the binomial SCR can be found in the online supplements.

```
Panel 6: MCMC updater for lamO in a SCR model with Binomial encounter process and cloglog link function on detection. Here, pmat =
```

```
1-exp(-lam).
5376
5377
             lam0.cand <- rnorm(1, lam0, 0.1)
5378
             if(lam0.cand >0){
                                 #automatically reject lam0.cand that are <0
5379
                 lam.cand <- lam0.cand*exp(-(D*D)/(2*sigma*sigma))</pre>
5380
                 p.cand <- 1-exp(-lam.cand)</pre>
5381
                 11<- sum(dbinom(y, K, pmat *z, log=TRUE))</pre>
5382
                 llcand <- sum(dbinom(y, K, p.cand *z, log=TRUE))</pre>
5383
                 if(runif(1) < exp( llcand - ll) ){</pre>
5384
                      11<-llcand
                      pmat<-p.cand
                      lam0<- lam0.cand
5387
                 }
5388
             }
5389
        Another possibility is to model variation in the individual and site specific de-
5390
    tection probability, pij, directly, without any transformation, such that
5391
    pij<-p0 * exp(-Dij2/(2*sig^2))
5392
    and p0 = \{0,1\}. This formulation is analogous to how detection probability is
5393
    modeled in distance sampling under a half-normal detection function; however, in
5394
    distance sampling p0 - detection of an individual on the transect line - is assumed
5395
    to be 1 (Buckland, 2001). Under this formulation the updater for lam0 (equivalent
    to p0 in Eq XX) becomes:
5397
              lam0.cand <- rnorm(1, lam0, 0.1)
5398
              if(lam0.cand >0 & lam0.cand < 1 ){
                                                           #automatically reject lam0.cand that are not
5399
                   lam.cand <- lam0.cand*exp(-(D*D)/(2*sigma*sigma))</pre>
5400
                   11<- sum(dbinom(y, K, lam *z, log=TRUE)) #no transformation needed
5401
                   llcand <- sum(dbinom(y, K, lam.cand *z, log=TRUE))</pre>
                   if(runif(1) < exp( llcand - ll) ){</pre>
5403
                        11<-11cand
                        lam<-lam.cand
5405
                        lam0<- lam0.cand
5406
                   }
5407
              }
```

#### 11.6.2 Looking at model output

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Now that you have an MCMC algorithm to analyze spatial capture-recapture data with, let's run an actual analysis so we can look at the output. As an example, we will use the bear data ... <sup>8</sup> You can use the same script provided back in

<sup>&</sup>lt;sup>8</sup>Does this data set come up before Ch6? If not, introduce data here. Or, Andy, would you rather use simulated data?

Chapter XX to read in the data and build the augmented encounter history array; then source the MCMC code for the binomial encounter model algorithm with the cloglog link and run 5000 iterations. This should take approximately 25 minutes.

```
5416 > source('SCRObinom.txt')
5417 > modO<-SCR.0(y=bigTrap, X=trapmat, M=M, xl=xl, xu=xu, yl=yl, yu=yu, K=8, niter=5000)</pre>
```

Before, we used simple R commands to look at model results. However, there is a specific R package to summarize MCMC simulation output and perform some convergence diagnostics package coda (Plummer et al., 2006). Download and install coda, then convert your model output to an memo object

> chain<-mcmc(mod0)

# Markov chain time series plots

Start by looking at time series plots of your Markov chains using plot(chain). This command produces a time series plot and marginal posterior density plots for each monitored parameter, similar to what we did before using the hist() and plot() commands (Fig. 5). Time series plots will tell you several things: First, the way the chains move through the parameter space gives you an idea of whether your MH steps are well tuned. If chains were constant over many iterations you would probably need to decrease the tuning parameter of the (Normal) proposal distribution. If a chain moves along some gradient to a stationary state very slowly, you may want to increase the tuning parameter so that the parameter space is explored more efficiently.

Second, you will be able to see if your chains converged and how many initial simulations you have to discard as burn-in. In the case of the chains shown in Figure 5, we would probably consider the first 750 - 1000 iterations as burn-in, as afterwards the chains seem to be fairly stationary.

#### A word of caution about chain convergence

Since we do not know what the stationary posterior distribution of our Markov chain should look like (this is the whole point of doing an MCMC approximation), we effectively have no means to assess whether it has converged to this desired distribution or not. As mentioned before, the only certainty is that a Markov chain will eventually converge to its stationary distribution, but no-one can tell us how long this will take. Also, you only now the part of your posterior distribution that the Markov chain has explored so far for all you know the chain could be stuck in a local maximum, while other maxima remain completely undiscovered. Acknowledging that there is truly nothing we can do to ever proof convergence of our MCMC chains, there are several things we can do to increase the degree of confidence we have about the convergence of our chains. One option, and that advocated by what we will loosely call the WinBUGS community, is to run several Markov chains and to start them off at different initial values that are overdispersed relative to the posterior distribution. Such initial values help to explore different

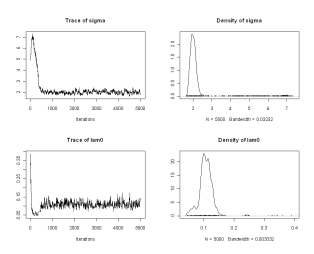


Figure 11.5. Time series and posterior density plots for sigma and lam0.

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areas of the parameter space simultaneously; if after a while all chains oscillate around the same average value, chances are good that they indeed converged to the posterior distribution. Gelman and Rubin came up with a diagnostic statistic that essentially compares within-chain and between-chain variance to check for convergence of multiple chains (Gelman et al., 2004). Of course, running several parallel chains is computationally expensive. Extra computational demands are not the only and by no means the major concern some people voice when it comes to running several parallel MCMC chains to assess convergence. Again, consider the fact that we do not know anything about the true form of the posterior distribution we are trying to approximate. How do we, then, know how to pick overdispersed initial values? We dont all we can do is pick overdispersed values relative to our expectations of what the posterior should look like. To use a quote from the home page of Charlie Geyer, a Bayesian statistician from the University of Minnesota, "If you don't know any good starting points [...], then restarting the sampler at many bad starting points is [...] part of the problem, not part of the solution." (http://users.stat.umn.edu/charlie/mcmc/diag.html). His suggestion is that your only chance to discover a potential problem with your MCMC sampler is to run it for a very long time. But again, there is no way of knowing how long is long enough. It is up to you to decide, which school of thoughts appeals more to you one long versus several parallel Markov chains. Irrespectively, part of developing an MCMC sampler should be to make sure (within reasonable limits) that you are not missing regions of high posterior density because of the way you specify your starting values. Once you have explored the behavior of your chain under a reasonable range of

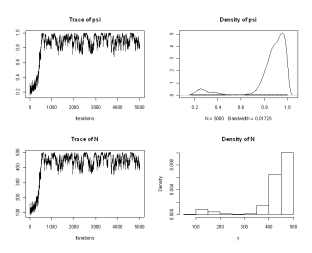
starting values, you may feel comfortable enough to run only one long chain. The fact that convergence cannot be proven does not mean that you should not look for potential problems in your MCMC sampler. Some problems are easily detected using simple plots, such as the time series plots we discussed above. If the overall trajectory of your chain at the end of your simulations is still upward or downward, your chain clearly has not converged and you need to run your model much longer. If you run several parallel chains and their stationary distributions look different, you may be looking at a multi-modal posterior or a problem with your sampler. With these words of caution, let's get back to looking at our model output.

#### 11.6.3 Posterior density plots

The plot() command also produces posterior density plots and it is worthwhile to look at those carefully. For parameters with priors that have bounds (e.g. Uniform over some interval), you will be able to see if your choice of the prior is truncating the posterior distribution. In the context of SCR models, this will mostly involve our choice of M, the size of the augmented data set. If the posterior of N has a lot of mass concentrated close to M (or equivalently the posterior of psi has a lot of mass concentrated close to 1), as in the example in Figure 6, we have to re-run the analysis with a larger M. A flat posterior plot shows you that the parameter essentially cannot be identified there may not be enough information in your data to estimate model parameters and you may have to consider a simpler model. Finally, posterior density plots will show you if the posterior distribution is symmetrical or skewed if the distribution has a heavy tail, using the mean as a point estimate of your parameter of interest may be biased and you may want to opt for the median or mode instead.

#### 11.6.4 Serial autocorrelation and effective sample size

Even when we can be relatively confident that our chains have converged, the subsequent samples generated from a Markov chain are not iid samples from the posterior distribution, due to the correlation amongst samples introduced by the Markov process. As a consequence, the variance of the mean cannot simply be derived with the standard variance estimator, which takes into account the sample size (here, number of iterations). Rather, the sample size has to be adjusted to account for the autocorrelation in subsequent samples (see Chapter 8 in Robert and Casella (2010) for more details). This adjusted sample size is referred to as the effective sample size. Checking the degree of autocorrelation in your Markov chains and estimating the effective sample size your chain has generated should be part of evaluating your model output. If you use WinBUGS through the R2WinBUGS package, the print() command will automatically return the effective sample size for all monitored parameters. In the coda package there are several functions you



**Figure 11.6.** Time series and posterior density plots of psi and N for the bear data set truncated by the upper limit of M (500).

can use to do so. effectiveSize() will directly give you an estimate of the effective sample size for you parameters:

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Alternatively, you can use the autocorr.diag() function, which will show you the degree of autocorrelation for different lag values (which you can specify within the function call, we use the defaults below):

```
        5522
        > autocorr.diag(mcmc(mod))

        5523
        sigma
        lam0
        psi
        N

        5524
        Lag
        0
        1.0000000
        1.0000000
        1.0000000
        1.0000000

        5525
        Lag
        1
        0.9979948
        0.9494134
        0.9847503
        0.9774201

        5526
        Lag
        5
        0.9915567
        0.8038168
        0.9111951
        0.9113525

        5527
        Lag
        10
        0.9836016
        0.6714021
        0.8462108
        0.8509803

        5528
        Lag
        50
        0.8985337
        0.1983780
        0.6138516
        0.6233994
```

Whichever function you use, if you find that your supposedly long Markov chain has not generated enough pseudo-iid samples, you should consider a longer run. In the present case we see that autocorrelation is especially high for the parameter sigma and our effective sample size for this parameter is 4! <sup>9</sup> This means we would

<sup>&</sup>lt;sup>9</sup>Anyone have any idea how the autocorrelation in sigma could be reduced?

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have to run the model for much longer to obtain a reasonable effective sample size. Unfortunately, with many SCR models we observe high degrees of serial autocorrelation, which means we have to run long chains to obtain enough samples that can be considered iid, in order to obtain reasonable estimates of our parameters and their variances. What exactly constitutes a reasonable effective sample size is hard to say, but as a rule of thumb you should probably aim at several hundreds of these pseudo-iid samples. A more meaningful measure of whether you've run your chain for enough iterations is the time-series or Monte Carlo error the 'noise' introduced into your samples by the stochastic MCMC process which we introduced in Chapter 2. The MC error decreases with increasing sample size and its magnitude can thus be controlled by adjusting the length of the Markov chain. As a rule of thumb, the MC error should be 1% or less of the parameter estimate. Once you have reached this level, the estimates of the mean, standard error and 95% quantiles should no longer change significantly with additional iterations. For highly correlated samples, it will take more iterations to reduce the MC error. In coda, the MC error is given as part of the summary results (see below). Another option to deal with the serial autocorrelation of samples is to 'thin' Markov chains by some rate r and save only every r-th iteration. But as discussed in Chapter 2, this is not efficient and should only be applied if needed for practical reasons (e.g. a large number of parameters and iterations may force you to thin your samples so you object storing the model output does not become unmanageably large). For now, let's continue using this small set of samples to continue looking at the output.

#### 11.6.5 Summary results

Now that we checked that our chains apparently have converged and pretending that we have generated enough samples from the posterior distribution, we can look at the actual parameter estimates. The summary() function will return two sets of results: the mean parameter estimates, with their standard deviation, the nave standard error - i.e. your regular standard error calculated for K (= number of iterations) samples without accounting for serial autocorrelation - and the corrected MC error (Time-series SE), which accounts for autocorrelation. In WinBUGS, this latter value is referred to as MC error and is only given in the log output within BUGS itself. You should adjust the summary() call by removing the burnin from calculating parameter summary statistics. To do so, use the window() command, which lets you specify at which iteration to start 'counting'. In contrast to WinBUGS, which requires you to set the burn-in length before you run the model, this command gives us full flexibility to make decisions about the burnin after we have seen the trajectories of our Markov chains. For our example, summary(window(chain, start=1001)) returns the following output:

```
71 Iterations = 1001:5000
72 Thinning interval = 1
```

```
Number of chains = 1
5573
    Sample size per chain = 4000
5574
5575
    1. Empirical mean and standard deviation for each variable,
5576
       plus standard error of the mean:
5577
5578
                                Naive SE Time-series SE
                Mean
5579
             1.9986
    sigma
                       0.13805 0.0021827
                                                   0.016091
5580
    lam0
             0.1096
                       0.01523 0.0002407
                                                   0.001401
5581
                                                   0.010734
    psi
             0.6113 0.09148 0.0014465
5582
    N
           489.8535 71.79695 1.1352094
                                                   8.431119
5583
5584
    2. Quantiles for each variable:
5585
5586
                 2.5%
                                                   75%
                                                           97.5%
                              25%
                                        50%
5587
    sigma
             1.75780
                         1.89847
                                     1.9900
                                               2.0944
                                                          2.2772
5588
             0.08357
                         0.09824
                                     0.1087
                                               0.1192
                                                          0.1427
    lam0
5589
             0.45110
                         0.54838
                                     0.6052
                                               0.6639
                                                          0.8192
    psi
5590
    N
           366.00000 440.00000 485.0000 530.0000 654.0000
5591
       Looking at the MC errors, we see that in spite of the high autocorrelation, the
5592
    MC error for sigma is below the 10ur algorithm gives us a posterior distribution of
5593
    N, but we are usually interested in the density, D. Density itself is not a parameter
5594
    of our model, but we can derive a posterior distribution for D by dividing each
5595
    value of N (N at each iteration) by the area of the state-space (here 3032.719 km<sup>2</sup>)
5596
    and we can use summary statistics of this distribution to characterize D:
    > summary(window(chain[,4]/ 3032.719, start=1001))
5598
    Iterations = 1001:5000
5599
    Thinning interval = 1
5600
    Number of chains = 1
5601
    Sample size per chain = 4000
5602
5603
    1. Empirical mean and standard deviation for each variable,
5604
       plus standard error of the mean:
5605
                                              Naive SE Time-series SE
                Mean
5607
                                                               0.0027801
          0.1615229
                           0.0236741
                                             0.0003743
5609
    2. Quantiles for each variable:
5610
5611
                25%
                        50%
      2.5%
                                75% 97.5%
5612
    0.1207 0.1451 0.1599 0.1748 0.2156
```

If we compare our mean density of 0.16/km2 (and other parameters) with results from the same model run in secr and WinBUGS in Chapter XX, we see that estimates are almost identical (Table 1).

#### 5617 11.6.6 Other useful commands

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While inspecting the time series plot gives you a first idea of how well you tuned your MH algorithm, use rejectionRate() to obtain the rejection rates (1 acceptance rates) of the parameters that are written to your output:

```
      5621
      > rejectionRate(chain)

      5622
      sigma
      lam0
      psi
      N

      5623
      0.44108822
      0.77675535
      0.00000000
      0.01940388
```

Recall that rejection rates should lie between 0.2 and 0.8, so our tuning seems to have been appropriate here. Psi is never rejected since we update it with Gibbs sampling, where all candidate values are kept. And since N is the sum of all z, all it takes for N to change from one iteration to the next are small changes in the z-vector, so the rejection rate of N is always low. If you have run several parallel chains, you can combine them into a single mcmc object using the mcmc.list() command on the individual chains (note that each chain has to be converted to an mcmc object before combining them with mcmc.list()). You can then easily obtain the Gelman-Rubin diagnostic (Gelman et al., 2004), in WinBUGS called R-hat, using gelman.diag(), which will indicate if all chains have converged to the same stationary distribution. For details on these and other functions, see the coda manual, which can be found together with the package on the CRAN mirror.

#### 11.7 MANIPULATING THE STATE-SPACE

So far, we have constrained the location of the activity centers to fall within the outermost coordinates of our rectangular state space by posing upper and lower bounds for x and y. But what if S has an irregular shape maybe there is a large water body we would like to remove from S, because we know our terrestrial study species does not occur there. Or the study takes place in a clearly defined area such as an island. As mentioned before, this situation is difficult to handle in WinBUGS. In some simple cases we can adjust the state space by setting SXi to be some function of SYi or vice versa. In this manner, we can cut off corners of the rectangle to approximate the actual state space. In R, we are much more flexible, as we can use the actual state-space polygon to constrain out si. <sup>10</sup>To illustrate that, let's look at a camera trapping study of Florida panthers (Puma concolor coryi) conducted in the Picayune Strand Restoration Project (PSRP) area, southwest Florida (Fig. 7), by XXX, and financed by XXX. In the 1960ies the PSRP area was slated for

<sup>&</sup>lt;sup>10</sup> Have to check if we can use panther stuff for the book; otherwise, use raccoon example.

housing development, but then bought back by the State of Florida and is currently being restored to its original hydrology and vegetation. In an effort to estimate the density of the local Florida panther population, 98 camera traps were operated in the area for 21 months between 2005 and 2007. Florida panthers are wide-ranging animals and in order to account for their wide movements, the state-space was defined as the trapping grid buffered by 15 km around its outermost coordinates. However, the resulting rectangle contained some ocean in its southwestern corner (Fig. 7). In order to precisely describe the state-space, the ocean has to be removed. You can create a precise state-space polygon in ArcGIS and read it into R, or create the polygon directly within R. In the present case we intersected two shape files one of the state of Florida and one of the rectangle defined by a strip of 15 km around the camera-trapping grid. While you will most likely have to obtain the shapefile describing the landscape of and around your trapping grid (coastlines, water bodies etc.) from some external source, a polygon shapefile buffering your outermost trapping grid coordinates can easily be written in R.

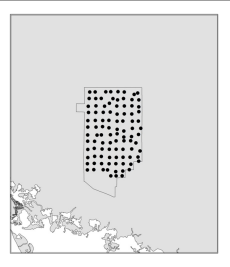
If xmin, xmax, ymin and ymax, mark the outermost x and y coordinates of your trapping grid and b is the distance you want to buffer with, load the package shapefiles (Stabler, 2006) and use:

You can read shapefiles into R loading the package maptools (Lewin-Koh et al., 2011) and using the function readShapeSpatial(). Make sure you read in shapefiles in UTM format, so that units of the trap array, the movement parameter sigma and the state-space are all identical. Intersection of polygons can be done in R also, using the package rgeos (Bivand and Rundel, 2011) and the function gIntersect(). The area of your single - polygon can be extracted directly from the state-space object SSp:

write.shapefile(ddShapefile, 'c:/, arcgis=T) # save to location of #choice

```
> area <- SSp@polygons[[1]]@Polygons[[1]]@area /1000000</pre>
```

Note that dividing by 1000000 will return the area in km2 if your coordinates describing the polygon are in UTM. If your state-space consists of several disjunct polygons, you will have to sum the areas of all polygons to obtain the size of the state-space. To include this polygon into our MCMC sampler we need one last spatial R package sp (Pebesma and Bivand, 2011), which has a function, over(),



**Figure 11.7.** Rectangular state-space for a Florida panther camera trapping study in the PSRP area (grey outline, red block inset map of Florida) contain some ocean (white) that needs to be removed from the state-space.

which allows us to check if a pair of coordinates falls within a polygon or not. All we have to do is embed this new check into the updating steps for s:

Scoord <- Spatial Points (Scand \* 1000) #convert to spatial points on UTM (m) scale Sin Poly <- over (Scoord, SSp) # check if scand is within the polygon

```
for(i in 1:M) {
if(is.na(SinPoly[i])==FALSE) { #if scand falls within polygon, continue update
[rest of the updating step remains the same]
```

Note that it is much more time-efficient to draw all M candidate values for s and check once if they fall within the state-space, rather than running the over() command for every individual pair of coordinates. To make sure that our initial values for s also fall within the polygon of S, we use the function runifpoint() from the package spatstat (Baddeley and Turner, 2005), which generates random uniform points within a specified polygon. You'll find this modified MCMC algorithm in the online supplementary material (SCR0poisSSp). Finally, observe that we are converting candidate coordinates of S back to meters to match the UTM polygon. In all previous examples, for both the trap locations and the activity centers we

have used UTM coordinates divided by 1000 to estimate sigma on a km scale. This is adequate for wide ranging individuals like bears. In other cases you may center all coordinates on 0. No matter what kind of transformation you use on your coordinates, make sure to always convert candidate values for S back to the original scale (UTM) before running the over() command.

#### 11.8 MCMC SOFTWARE PACKAGES

Throughout most of this book we will use WinBUGS and, occasionally, JAGS to run MCMC analyses. Here, we will briefly discuss the main pros and cons of these two programs as well as WinBUGS successor OpenBUGS. You can find scripts to simulate data and run the basic SCR model in all three programs in the online supplementary material (simSCR0poisBUGS).

#### 11.8.1 WinBUGS

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In a nutshell, WinBUGS (and the other programs) do everything that we just went 5720 through in this chapter (and quite a bit more). Looking through your model, Win-5721 BUGS determines which parameters it can use standard Gibbs sampling for (i.e. for conjugate full conditional distributions). Then, it determines, in the following 5723 hierarchy, whether to use adaptive rejection sampling, slice sampling or in the 5724 'worst' case Metropolis-Hastings sampling for the other full conditionals (Spiegel-5725 halter et al., 2003). If it uses MH sampling, it will automatically tune the updater 5726 so that it works efficiently. While WinBUGS is a convenient piece of software that 5727 is still widely used, its major drawback is that it is no longer being developed, i.e. 5728 no new functions or distributions are added and no bugs are fixed. 5729

#### 11.8.2 OpenBUGS

OpenBUGS is essentially the successor of WinBUGS. While the latter is no longer worked on, OpenBUGS is constantly developed further. The name 'OpenBUGS' refers to the software being open source, so users do not need to download a license key, like they have to for WinBUGS (although the license key for WinBUGS is free and valid for life).

Compared to WinBUGS, OpenBUGS has a lot more built-in functions. The method of how to determine the right updater for each model parameter has changed and the user can manually control the MCMC algorithm used to update model parameters. Several other changes have been implemented in OpenBUGS and a detailed list of differences between the two BUGS versions, can be found at http://www.openbugs.info/w/OpenVsWin

While OpenBUGS is a useful program for a lot of MCMC sampling applications, for reasons we do not understand, simple SCR models do not converge in OpenBUGS. It is therefore advisable that you check any OpenBUGS SCR model results

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5782 5783 against result from WinBUGS. Also, currently, the R package BRugs (Thomas et al., 2006) necessary for running OpenBUGS through R has problems with 64-bit machines, so you may have to use the 32-bit version of R and OpenBUGS in order to make it work. The BUGS project site at http://www.openbugs.info provides a lot of information on and download links for OpenBUGS.

There is an extensive help archive for both WinBUGS and OpenBUGS and you can subscribe to a mailing list, where people pose and answer questions of how to use these programs at http://www.mrc-bsu.cam.ac.uk/bugs/overview/list.shtml

#### 11.8.3 JAGS Just Another Gibbs Sampler

JAGS, currently at Version 3.1.0, is another free program for analysis of Bayesian hierarchical models using MCMC simulation. Originally, JAGS was the only program using the BUGS language that would run on operating systems other than the 32 bit Windows platforms. By now, there are OpenBUGS versions for Linux or Macintosh machines. JAGS 'only' generates samples from the posterior distribution; analysis of the output is done in R either by running JAGS through R using either the packages rjags (Plummer, 2011) or R2jags (Su and Yajima, 2011), or by using coda on your JAGS output. The program, manuals and rjags can be downloaded at http://sourceforge.net/projects/mcmc-jags/files/ When run from within R using the package rjags or R2jags, writing a JAGS model is virtually identical to writing a WinBUGS model. However, some functions may have slightly different names and you can look up available functions and their use in the JAGS manual. One potential downside is that JAGS can be very particular when it comes to initial values. These may have to be set as close to truth as possible for the model to start. Although JAGS lets you run several parallel Markov chains, this characteristic interferes with the idea of using overdispersed initial values for the different chains. Also, we have occasionally experienced JAGS to crash and take the R GUI with it. Only re-installing both JAGS and rjags seemed to solve this problem. On the plus side, JAGS usually runs a little faster than WinBUGS, sometimes considerably faster (see section 4.XYZ), is constantly being developed and improved and it has a variety of functions that are not available in WinBUGS. For example, JAGS allows you to supply observed data for some deterministic functions of unobserved variables. In BUGS we cannot supply data to logical nodes. Another useful feature is that the adaptive phase of the model (the burn-in) is run separately from the sampling from the stationary Markov chains. This allows you to easily add more iterations to the adaptive phase if necessary without the need to start from 0. There are other, more subtle differences and there is an entire manual section on differences between JAGS and OpenBUGS. For questions and problems there is a JAGS forum online at http://sourceforge.net/projects/mcmc-jags/forums/forum/610037.

 $<sup>^{11}\</sup>mathrm{As}$  we make progress on the book, lets be sure to add linkages to places where we use JAGS in examples.

#### 11.9 SUMMARY AND OUTLOOK

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While there are a number of flexible and extremely useful software packages to perform MCMC simulations, it sometimes is more efficient to develop your own MCMC algorithm. Building an MCMC code follows three basic steps: Identify your model including priors and express full conditional distributions for each model parameter. If full conditionals are parametric distributions, use Gibbs sampling to draw candidate parameter values from this distributions; otherwise use Metropolis-Hastings sampling to draw candidate values from a proposal distribution and accept or reject them based on their posterior probability densities. These custom-made MCMC algorithms give you more modeling flexibility than existing software packages, especially when it comes to handling the state-space: In BUGS (and JAGS for that matter) we define a continuous rectangular state-space using the corner coordinates to constrain the Uniform priors on the activity centers s. But what if a continuous rectangle isn't an adequate description of the state-space? In this chapter we saw that in R it only takes a few lines of code to use any arbitrary polygon shapefile as the state-space, which is especially useful when you are dealing with coastlines or large bodies of water that need removing from the state-space. Another example is the SCR R package SPACECAP (Gopalaswamy et al., 2011) that was developed because implementation of an SCR model with a discrete state-space was inefficient in WinBUGS. Another situations in which using BUGS/JAGS becomes increasingly complicated or inefficient is when using point processes other than the Uniform Poisson point process which underlies the basic SCR model (see Chapter X). In the Chapters 9 and XX you will see examples of different point processes, implemented using custom-made MCMC algorithms. 12 Finally, the Chapters XX and XX deal with unmarked or partially marked populations using hand-made MCMC algorithms to handle the (partially) latent individual encounter histories. While some of these models can be written in BUGS/JAGS, <sup>13</sup>, they are painstakingly slow; others cannot be implemented in BUGS/JAGS at all. In conclusion, while you can certainly get by using BUGS/JAGS for standard SCR models, knowing how to write your own MCMC sampler allows you to tailor these models to your specific needs.

<sup>&</sup>lt;sup>12</sup>Richard, Beth expand on that?

<sup>&</sup>lt;sup>13</sup>the Poisson one for partially marked we wrote in BUGS and it should work with a known number of marked; the Bernoulli in JAGS with the dsum() function should work for the fully unknown; maybe some others? I dont remember. We may have to try writing the others before saying that they dont work in BUGS/JAGS; they are certainly much faster in R, though.

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# 

### STATE-SPACE COVARIATES

Underlying all spatial capture recapture models is a point process model describing the distribution of individual activity centers  $(\mathbf{s}_i)$  within the state space  $(\mathcal{S})$ . So far we have focused our discussion on the homogeneous binomial point process,  $\mathbf{s}_i \sim Uniform(\mathcal{S}), i=1,2,\ldots,N$ , where N is the size of the population. This is a model of "spatial-randomness" because the intensity of the activity centers is constant across the study area and the activity centers are distributed independently of each other.

The spatial-randomness assumption is often viewed as restrictive because ecological processes such as territoriality and habitat selection can result in non-random distributions of organisms. We have argued, however, that this assumption is less restrictive than may be recognized because the homogeneous point process actually allows for infinite possible configurations of activity centers. Furthermore, given enough data, the uniform prior will have very little influence on the estimated locations of activity centers. Nonetheless, the homogeneous point process model does not allow one to model population density using covariates—a central objective of much ecological research. For example, a homogeneous point process model may result in a density surface map indicating that individuals were more abundant in one habitat than another, but it does not do so explicitly. A more direct approach would be to model density using covariates as is done in generalized linear models (GLMs).

In this chapter we will present a method for fitting inhomogeneous binomial point process models using covariates in much the same way as is done with GLMs. The covariates we consider differ from those covered in previous chapters, which were typically attributes of the animal (e.g. sex, age) and were used to model movement or encounter rate. In contrast, here we wish to model covariates that

 $<sup>^{1}\</sup>mathrm{The}$  phrase "complete spatial-randomness" is reserved for the homogeneous Poisson point process

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are defined for all points in S, which we will refer to as state-space, or density, covariates. These may include continuous covariates such as elevation, or discrete covariates such as habitat type.

Borchers and Efford (2008) were the first to propose an inhomogeneous point process model for SCR models, and our approach is similar to theirs with the exception that we will use a binomial rather than a Poisson model because the binomial model is easily integrated into our data augmentation scheme and is consistent with the objective of determining how a *fixed* number of activity centers are distributed with respect to covariates.

The method we use to accommodate inhomogeneous binomial point process models within our MCMC algorithm is simple—we replace the uniform prior with a prior describing the distribution of the N activity centers conditional on the covariates. Development of this prior, which does not have a standard form, is a central component of this chapter. First we will begin with a review of homogeneous point process models.

#### 14.1 HOMOGENEOUS POINT PROCESS REVISITED

The homogeneous Poisson point process is the model of "complete spatial randomness" and is often used in ecology as a null model to test for departures from randomness. Given its central role in the analysis of point processes, it is helpful to compare it with the binomial model that we use in our SCR models. The primary descriptor of the homogeneous point process model is the "intensity" parameter,  $\mu$  which describles the expected number of points in an infinitesimally small area. The intensity parameter can also be used to determine the expected number of points in any region of the state-space  $\mathcal{S}$ . To denote this, we say that the expected number of points in region  $B \in \mathcal{S}$  is  $n(B) = A(B)\mu$  where A(B) is the area of region B. In words, the expected number of points in B is simply the area of Bmuliplied by the intensity parameter. One property of the Poisson model is that if we divide the entire state-space into  $k = 1, \ldots, K$  disjunct regions, the counts  $\mathbf{n}(\mathbf{B})$ are independent and identically distributed, (i.i.d.). This is one of the distinctions between the Poisson model and the binomial model, for which the counts  $n(B_k)$ are not i.i.d. as we will explain shortly. This difference is also related to another distinction between the two models, namely that the binomial model conditions on the number of points to be simulated N; whereas under the Poisson model N is random. Here is some simple **R** code to illustrate this point.

```
mu <- 4 # intensity
Np <- rpois(1, mu) # Np is random
PPP <- cbind(runif(Np), runif(Np)) # Poisson point process
Nb <- 4 # Nb is fixed
BPP <- cbind(runif(Nb), runif(Nb)) # Binomial point process
```

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Note that in both models, the N points are independent of one another and distributed uniformly throughout S. Thus, the intensity at any point  $x \in S$  is  $\mu = 1/A(S)$  where A(S) denotes the area of the state-space. In the  $\mathbf{R}$  code above, the area of the state-space is 1 unit, and thus the intensity is  $\mu = 1/1$ .

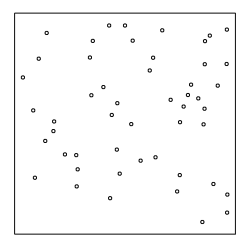
Although the Poisson model is typically described in terms of  $\mu$ , the binomial model is not; rather, it is more common to consider a discrete state space, such as a grid with with K pixels. Under the binomial model, the number of points in each region is  $n(B_k) \sim Bin(N, p_k)$  where  $p_k = A(B)/A(S)$ , ie  $p_k$  is simply the fraction of the state-space area in  $B_k$ . This discrete space representation of the binomial point process is shown in Fig. 14.1. The state-space in this case is the unit square, and thus the probability of a point falling in each of the 25 disjunct regions is  $p_k = 1/25$  and thus the expected counts are simply  $\mathbb{E}(n(B_k)) = Np_k$ . In the figure N = 50 and thus we would expect 2 points per pixel, which happens to be the empirical mean of the data in Fig. 14.1. Note also that these counts are not independent realizations from a binomial distribution since  $\sum_k n(B_k) = N$ . Instead, the model for the entire vector is  $\mathbf{n}(\mathbf{B}) \sim Multinomial(N, \pi = (p_1, p_2, \dots, p_K))$ (Illian, 2008b). The dependence among counts has virtually no practical consequence when the number of pixels is large. For example, if we have 100 pixels, the number of counts in one pixels tells you very little about the expected count in another pixel. However, if there are only 2 pixels, then clearly the number of points in one pixel tells you exactly how many will occur in the remaining pixel. To gain familiarity with the multinomial distribution and the discrete representation of space, use the rmultinom function in R to simulate counts similar to those shown in Fig. 14.1, for example using a command such as:

```
n.B_k <- rmultinom(1, size=50, prob=rep(1/25, 25))
matrix(n.B_k, 5, 5)</pre>
```

The discrete space representation of the binomial point process is of practical importance when fitting SCR models because spatial covariates are almost always represented in a discrete format, often called "rasters" in GIS-speak. In such cases, we often need to change our definition of the prior for an activity center from  $s_i \sim Uniform(\mathcal{S})$  to  $s_i \sim Multinomial(1,\pi)$ . In the latter case, the activity center is simply defined as an integer representing pixel "id". Note also that the multinomial distribution with an index of 1 (i.e. size=1 in rmultinom) is referred to as the categorical distribution, which we will frequently use in the BUGS language.

#### 14.2 INHOMOGENEOUS BINOMIAL POINT PROCESS

As with the homogeneous model, the inhomogeneous binomial point process model is developed conditional on N. The primary distinction is that the uniform distribution is replaced with another distribution allowing for the intensity parameter to vary spatially. To arrive at this new distribution, define  $\mu(x,\beta)$  to be a function of



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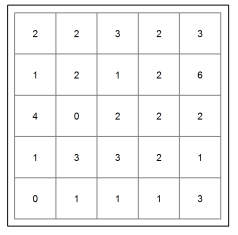


Figure 14.1. Homogeneous binomial point process with N=50 points represented in continuous and discrete space.

spatially-referenced covariates  $(\beta)$  available at all points of the state space. To be concise we will subsequently drop the vector of cofficients from our notation, and simply use  $\mu(x)$ . Since an intensity must be strictly positive, it is natural to model  $\mu(x)$  using the log-link.

$$\log(\mu(x)) = \sum_{j=1}^{J} \beta_j v_j(x), \quad x \in \mathcal{S}$$

where  $\beta_j$  is the regression coefficient for covariate  $v_j(x)$ . To be clear, v(x) is the value of any covariate, such as habitat type or elevation, at location x. This equation should look familiar because it is the standard linear model used in log-linear GLMs. Note, however, that we have no need for an intercept because it would be confounded with N. This should be intuitive since an intercept would represent the expected value of N when  $\beta=0$ , but we already have a parameter in the model for expected abundance, namely  $\mathbb{E}[N]=\psi M$ . Thus an intercept would be redundant, and without it we are still able to achieve our goal of describing the distribution of N activity centers as a function of spatial covariates.

Now that we have a model of the intensity parameter  $\mu(x)$ , we need to develop the associated probability density function to use in place of the uniform prior. Remembering that the integral of a pdf must be unity, we can create a pdf by dividing  $\mu(x)$  by a normalizing constant, which in this case is the integral of  $\mu(x)$ evaluated over the entire state-space ANDY, is there a better justification for this? EXAMPLES 201

The probability density function is therefore

$$f(x) = \frac{\mu(x)}{\int_{x \in S} \mu(x) \, \mathrm{d}x}$$
 (14.2.1)

Substituting this distribution for the uniform prior allows us to fit inhomogeneous binomial point process models to spatial capture-recapture data. We can also use this distribution to obtain the expected number of individuals in any given region. Specifically, the proportion of N expected to occur in any region B when heterogeneity in density is present is  $p(B) = \int_B f(x) \, \mathrm{d}x$ . These are also the multinomial cell probabilities if the regions are disjoint and compose the entire state-space.

As a practical matter, note that the integral in the demoninator of f(x) is evaluated over space, and since we almost always regard space as two-dimensional, this is a two-dimensional integral that can be approximated using the methods discussed in refChXXX. These methods include Monte Carlo integration, Gaussian quadrature, etc... Alternatively, if our state-space covariates are in raster format, *i.e* they are in discrete space, the integral can be replaced with a sum over all pixels, which is much more efficient computationally.

We now have all the tools needed to fit inhomogeneous point process (IPP) models. Before doing so, we note that the IPP for the activity centers results in another IPP for the observation process,  $\lambda(x)$ . As a reminder,  $\lambda(x)$  is the expected number of captures for a trap at point x. As was true for the homogeneous model, this intensity function is a product of the point process intensity and the encounter rate function,  $\lambda(x) = \mu(x)g(x)$ .

In the next section we walk through a few examples, building up from the simplest case where we actually observe the activity centers as though they were data. In the second example, we fit our new model to simulated data in which density is a function of a single continuous covariate. Example three shows an analysis in discrete space using both secr (Efford, 2011) and JAGS (Plummer, 2003). In the last example, we model the intensity of activity centers for a real dataset collected on jaguars (*Panthera onca*) in Argentina.

#### 14.3 EXAMPLES

#### 14.3.1 Simulation and analysis of inhomogeneous point processes

In SCR models, the point process is not directly observed, but in other contexts it is. Examples include the locations of disease outbreaks or the locations of trees in a forest. Fitting inhomogeneous point process models to such data is straightforward and illustrates the fundamental process that we will later embed in our MCMC algorithm used to fit SCR models.

Suppose we knew the locations of 100 animals' activity centers. To estimate the intensity surface  $\mu(x)$  underlying these points, we need to derive the likelihood for our data under this model. Given the pdf f(x) (Eq. 14.2.1) and assuming that

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the points are mutually independent of one another, we may write the likelihood as the product of R such terms, where R = 100 is the sample size in this case, ie the observed number of activity centers.

$$\mathcal{L}(\beta|\mathbf{x}_i) = \prod_{i=1}^R f(x_i)$$

Having defined the likelihood we could choose a prior and obtain the posterior for  $\beta$  using Bayesian methods, or we can find the maximum likelihood estimates (MLEs) using standard numerical methods as is demonstrated below.

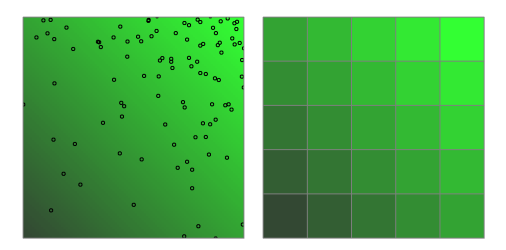
First, let's simulate some data. Simulating data under an inhomogeneous point process model is often accomplished using indirect methods such as rejection sampling. Rejection sampling proceeds by simulating data from a standard distribution and then accepting or rejecting each sample using probabilities defined by the distribution of interest. For more information, readers should consult an accessible text such as Robert and Casella (2004). In our example, we simulate from a uniform distribution and then accept or reject using the (scaled) probability density function f(x). Note that we first define a spatial covariate (elevation) that is a simple function of the spatial coordinates increasing from the southwest to the northeast of our state-space.<sup>2</sup>

The following **R** commands demonstrate the use of rejection sampling to simulate an inhomogeneous point process for the covariate depicted in Fig. 14.3.1. The code uses the cuhre function in the R2Cuba package to integrate the intensity function over space (Hahn et al., 2011).

```
# spatial covariate (with mean 0)
     elev.fn <- function(x) x[1]+x[2]-1
6000
     # intensity function
6001
    mu <- function(x, beta) exp(beta*elev.fn(x=x))</pre>
6002
6003
     # Simulate PP using rejection sampling
6004
     set.seed(300225)
     N <- 100
6006
     count <- 1
6007
     s <- matrix(NA, N, 2)
6008
     beta <- 2 # parameter of interest
6009
     int.mu <- R2Cuba:::cuhre(2, 1, mu, beta=beta)$value
     elev.min \leftarrow elev.fn(c(0,0)) #elev.fn(cbind(0,0))
     elev.max \leftarrow elev.fn(c(1,1)) #elev.fn(cbind(1,1))
     Q <- max(c(exp(beta*elev.min) / int.mu,
6013
                 exp(beta*elev.max) / int.mu))
6014
    while(count <= 100) {
6015
```

<sup>&</sup>lt;sup>2</sup>Such functional forms of covariates are rarely available, which is why continuous spatial covariates are more often measured on a discrete grid.

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**Figure 14.2.** An example of a spatial covariate, say elevation, and a realization of a inhomogeneous binomial point process with N=100 and  $\mu(x) = exp(\beta Elev)$  where  $\beta = 2$ .

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The simulated data are shown in Fig 14.3.1. High elevations are represented by light green and low elevations by dark green. The activity centers of one hundred animals are shown as points, and it is clear that these simulated animals prefer the high elevations. Perhaps they are mountain goats. The underlying model describing this preference is  $\log(\mu(x)) = \exp(\beta \times Elevation(x))$  where  $\beta = 2$  is the parameter to be estimated and Elevation(x) is a function of the coordinates at x, as displayed on the map.

Given these points, we will now estimate  $\beta$  by minimizing the negative-log-likelihood using R's optim function.

```
# Negative log-likelihood

nll <- function(beta) {

int.mu <- cuhre(2, 1, mu, beta=beta)$value

-sum(beta*elev.fn(s) - log(int.mu))
```

```
6037 }
6038 starting.value <- 0
6039 fm <- optim(starting.value, nll, method="Brent",
6040 lower=-5, upper=5, hessian=TRUE)
6041 c(Est=fm$par, SE=sqrt(1/fm$hessian)) # estimates and SEs
```

Maximizing the likelihood took a small fraction of a second, and we obtained an estimate of  $\hat{\beta} = 1.99$ . We could plug in this estimate to our linear model at each point in the state-space to obtain the MLE for the intensity surface.

This example demonstrates that if we had the data we wish we had, *i.e.* if we knew the coordinates of the activity centers, we could easily estimate the parameters governing the underlying point process. Unfortunately, in SCR models, the activity centers cannot be directly observed, but spatial re-captures, that is captures of individuals at multiple locations in space, provide us with the information needed to estimate these latent parameters.

#### 14.3.2 Fitting inhomogeneous point process SCR models

#### Continuous space

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One of the nice things about hierarchical models is that they allow us to break a problem up into a series of simple conditional relationships. Thus, we can simply add the methods described above into our existing MCMC algorithm to simulate the posteriors of  $\beta$  conditional on the simulated values of  $\mathbf{s}_i$ . To demonstrate, we will continue with the previous example. Specifically, we will overlay a grid of traps upon the map shown in Fig. 14.3.1. We will then simulate capture histories conditional upon the activity centers shown on the map. Then, we will attempt to estimate the activity center locations as though we did not know where they were, as is the case in real applications.

Here is some  $\mathbf{R}$  code to simulate the encounter histories under a Poisson observation model, which would be appropriate if animals could be detected multiple times at a trap during a single occassion.

```
# Create trap locations
6065
     xsp \leftarrow seq(-0.8, 0.8, by=0.2)
     len <- length(xsp)</pre>
6067
     X <- cbind(rep(xsp, each=len), rep(xsp, times=len))</pre>
6068
6069
     # Simulate capture histories, and augment the data
6070
     ntraps <- nrow(X)</pre>
6071
     T <- 5
     y <- array(NA, c(N, ntraps, T))
     nz <- 50 # augmentation
6075
     M <- nz+nrow(y)</pre>
6076
     yz <- array(0, c(M, ntraps, T))
```

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```
6078
     sigma <- 0.1 # half-normal scale parameter
6079
     lam0 <- 0.5
                      # basal encounter rate
6080
     lam <- matrix(NA, N, ntraps)</pre>
6081
6082
     set.seed(5588)
6083
     for(i in 1:N) {
6084
          for(j in 1:ntraps) {
6085
              distSq \leftarrow (s[i,1]-X[j,1])^2 + (s[i,2] - X[j,2])^2
6086
              lam[i,j] \leftarrow exp(-distSq/(2*sigma^2)) * lam0
6087
              y[i,j,] <- rpois(T, lam[i,j])
          }
6090
     yz[1:nrow(y),,] <- y # Fill</pre>
6091
6092
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```

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Now that we have a simulated capture-recapture dataset y, and we have augmented it to create the new data object yz, we are ready to begin sampling from the posteriors. A commented Gibbs sampler written in  $\mathbf{R}$  is available in the accompanying  $\mathbf{R}$  package scrbook (see ?scrIPP). There are two small parts of the  $\mathbf{R}$  code that distinguish it from previous code we have shown to fit homogeneous point processes. First, we need to update the parameter  $\beta$  conditional on all other parameters in the model. The code to do so is:

```
D1 <- cuhre(2, 1, mu, lower=c(xlims[1], ylims[1]),
6099
                  upper=c(xlims[2], ylims[2]), beta=beta1)$value
6100
     beta1.cand <- rnorm(1, beta1, tune[3])</pre>
6101
     D1.cand <- cuhre(2, 1, mu, lower=c(xlims[1], ylims[1]),
6102
                        upper=c(xlims[2], ylims[2]), beta=beta1.cand)$value
6103
    11.beta1 <- sum( beta1*elev.fn.v(S) - log(D1) )</pre>
6104
     11.beta1.cand <- sum( beta1.cand*elev.fn.v(S) - log(D1.cand) )</pre>
6105
     if(runif(1) < exp(ll.beta1.cand - ll.beta1) ) {</pre>
6106
          beta1<-beta1.cand
6108
    }
        Next, we need to put the new prior on the activity centers:
6109
     #ln(prior), denominator is constant
6110
    prior.S <- beta1*cov(S[i,1], S[i,2]) # - log(D1)</pre>
6111
    prior.S.cand <- beta1*(Scand[1] + Scand[2]) # - log(D1)</pre>
6112
     if(runif(1) < exp((11.S.cand+prior.S.cand) - (11.S+prior.S))) {</pre>
6113
         S[i,] <- Scand
         lam <- lam.cand
         D[i,] <- dtmp
6116
```

We can apply this modified sampler to our data using the code shown in the help file for scrIPP. We obtain posterior distributions summarized in Table 14.2.

Mixing is good, and as usual, life is very nice when we are working with simulated data.

Fitting continuous space IPP models is somewhat difficult in **BUGS** because our prior f(x) is not one of the available distributions that come with the software<sup>3</sup> secr allows users to fit continuous space using polynomials of the x- and y- coordinates, but not for truly continuous covariates. However, these are not really important limitations because discrete space versions are straight-forward, and virtually all spatial covariates are defined as such.

#### Discrete space

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To fit discrete space models, we follow the same steps as outlined in Chapter XXX—we define  $s_i$  as pixel ID, and we use the categorical distribution as a prior. A good example of this is in +citeKery capricaillie. Here we present an analysis of the simulated data shown in the right panel of Fig. 14.3.1. The spatial covariate, let's call it elevation again, was simulated from a kriging type of model as shown on the help page ch9simData in scrbook. The points are the number of activity centers in each pixel, generated from a single realization of the IPP mu(x) = 2elev.

The **BUGS** code to fit an IPP model to these data is shown in the following panel.

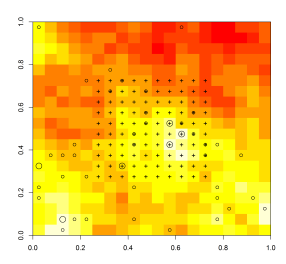
```
model{
     sigma ~ dunif(0, 1)
6139
     lam0 ~ dunif(0, 5)
6140
     beta \sim dnorm(0,0.1)
6141
     psi ~ dbeta(1,1)
6142
6143
     for(j in 1:nPix) {
6144
       theta[j] <- exp(beta*elevation[j])</pre>
6145
       probs[j] <- theta[j]/sum(theta[])</pre>
6146
6147
6148
     for(i in 1:M) {
6149
       w[i] ~ dbern(psi)
6150
       s[i] ~ dcat(probs[])
       x0g[i] <- Sgrid[s[i],1]</pre>
6152
```

Table 14.1. Posterior summaries from inhomogeneous point proces model

	Mean	SD	2.5%	50%	97.5%
$\sigma = 0.10$	0.1026	0.0048	0.0935	0.1025	0.1123
$\lambda_0 = 0.50$	0.4419	0.0493	0.3496	0.4400	0.5390
$\psi = 0.66$	0.6826	0.0554	0.5762	0.6820	0.7923
$\beta = 2.00$	2.1601	0.3390	1.5193	2.1583	2.8043
N = 100	102.7696	6.2689	92.0000	102.0000	117.0000

<sup>&</sup>lt;sup>3</sup>It is possible, if somewhat cumerbersome, to add new distributions in BUGS.

EXAMPLES 207



**Figure 14.3.** Simulated activity centers in discrete space. The spatial covariate, elevation, is highest in the ligher areas. Density of activity centers (circles) increases with elevation. Trap locations are shown as crosses.

```
y0g[i] <- Sgrid[s[i],2]</pre>
6153
       for(j in 1:ntraps) {
6154
          dist[i,j] \leftarrow sqrt(pow(x0g[i]-grid[j,1],2) + pow(y0g[i]-grid[j,2],2))
6155
          lambda[i,j] <- lam0*exp(-dist[i,j]*dist[i,j]/(2*sigma*sigma)) * w[i]</pre>
          y[i,j] ~ dpois(lambda[i,j])
6157
6158
       }
6159
6160
     N <- sum(w[])</pre>
6161
     Density <- N/1 # unit square
6162
6163
     }
```

This model can also be fit in secr, which refers to the pixel locations as a "mask". R code to fit the models using secr and JAGS is available in scrbook, see help(ch9secrYjags). Results of the comparision are shown in Table ?? and are very similar as expected.

Density surface maps can be created for fun, and of course to inform management decisions. [describe how to do this]

#### 14.3.3 The jaguar data

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Estimating density of large felines has been a priority for many conservation organizations, but no robust methodologies existed before the advent of SCR. Distance sampling is not feasible for such rare and cryptic species, and traditional capture-recapture methods yield estimates that are highly sensitive to the subjective choice of the effective survey area. In this example, we demonstrate how readily density can be estimated for a globally imperilled species using SCR. Furthermore, we show how inhomogeneous point process models can be used to test important hypotheses regarding the factors affecting density.

[describe study]

A few aspects of this design are noteworthy. First, the dimensions and configuration of the trap array differed among the regions of the trap array. This fact alone could explain variation in the number of animals exposed to sampling, which

Software	Par	Est.	SD	lower	upper
secr	N	49.2803	5.7535	41.0087	64.3879
	$\beta$	2.1772	0.5628	1.0741	3.2804
	$\lambda_0$	0.9203	0.0764	0.7824	1.0825
	$\sigma$	0.0990	0.0038	0.0918	0.1068
$_{ m JAGS}$	N	48.2072	5.4053	39.0000	60.0000
	$\beta$	2.1026	0.5323	1.0889	3.1506

0.9328 0.0766

 $0.1004 \quad 0.0041$ 

 $\lambda_0$ 

1.0921

0.1089

0.7898

0.0929

Table 14.2. Comparision of secr and JAGS results

SUMMARY 209

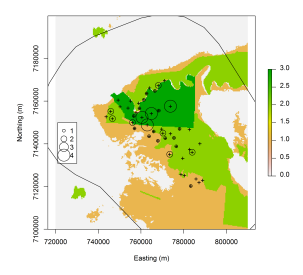


Figure 14.4. Jaguar detections

would have no biological meaning. Furthermore, the area of inference is an irregular polygon that was not sampled uniformly. Only by estimating density can we hope to extrapolate our estiamates from the sampled region to get what we are after. In this case, this is readily accomplished since the entire state-space can be classified as one of the 3 levels of protection from poaching. Of course, it general it is always preferable to sample more uniformly throughout the area of interset in case some unmeasured covariate biases the extrapolation.

To assess the influence of poaching on jaguar density, we considered 2 metrics of poaching pressure, one political and one continuous measure of accessibility (Fig xxx).

### 14.4 SUMMARY

When state-space covariates are available, we can model density by replacing the uniform prior on the activity centers with a prior based on a normalized log-linear function of covariates. This yields a model of the inhomogeneous point process describing the location of activity centers, which can be used to test hypotheses about covariates affecting density. In rare cases, these covariates are truly continuous in the sense that they are defined as a function of space. More often, covariates are represented on rasters, which simplifies the analysis. Fitting these models can be accomplished using **BUGS**, **secr**, or the custom **R** code presented in this chapter

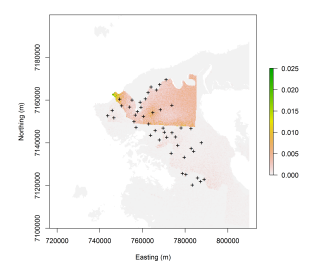


Figure 14.5. Estimated density surface for the jaguar dataset

and found in the package scrbook.

All the examples in this section included a single state-space covariate, but this was for simplicity only. Including multiple covariates poses no additional challenges. Likewise, additional model structure such sex-specific encounter rate parameters or behavioral responses can be accommodated.

#### 14.5 OTHER IDEAS

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Should have some discussion on some ideas for building flexible models. Might be cool to use the Ickstadt/Wolpert as a model for the inhomogeneous point process. Dont have to do it, just mention it. Also some kind of a spline model or similar.

209	15	
211	INHOMOGENEOUS POINT PROCESS	

5212 5213	16
5214	OPEN MODELS

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