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### CHAPTER



# Closed Population Models

models presented in this book, including the models for open populations discussed closed population CR models serve as the basis fouthe development of the rest of the Although few populations are actually closed except during very short time intervals, which states that no individuals move onto or off of the sampled area during the study. demographic closure, meaning that no births or deaths occur, and geographic closure, size, N, does not change during the study. Two forms of closure are often discussed: estimating population size in closed populations. A closed population is one whose pooos In this chapter we introduce ordinary non-spatial capture-recapture (CR) models for

models. In fact, when M'is known, they are precisely logistic regression models. closed population models are closely related to binomial (or logistic) regression-type the most primitive spatial capture-recapture model. All of these different types of of individual covariate models is distance sampling, which could be thought of as random effect, which represent unobserved or unmeasured covariates. A special type (sex, age, body mass) or unstructured "heterogeneity" in the form of an individual ous types of "individual effects" either in the form of explicit, observed covariates important extensions of ordinary closed population models that accommodate varistructure of closed population models as binomial GLMs. We then consider some all respects (across individuals, and replicates). This allows us to highlight the basic "model M<sub>6</sub>" (Otis et al., 1978), in which encounter probability is strictly constant in We begin with the most basic capture-recapture model, colloquially referred to as in Chapter 16.

models. tary development of Bayesian analysis of ordinary, i.e., non-spatial closed population the reader to Kéry and Schaub (2012, chapter 6) for an accessible and complementechnical origins and demonstrate applications to closed population models. We refer We use data augmentation throughout this book and thus emphasize its conceptual and from the standpoint of doing MCMC (Royle et al., 2007; Royle and Dorazio, 2012). mentation across classes of capture-recapture models and one that is really convenient context of capture-recapture models where N is unknown, it has a consistent impleature (e.g., Tanner and Wong, 1987). This is a general concept in statistics but, in the this using a method related to classical "data augmentation" from the statistics liter-We emphasize Bayesian analysis of capture-recapture models and we accomplish

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# 4,1 The simplest closed population model: model Mo

Bernoulli distribution to a binomial distribution based on a sample of size K. That is ters for each individual,  $y_i = \sum_i y_i$ , and the observation model changes from a original binary detection variables can be aggregated into the total number of encounp is constant (i.e., there are no individual or temporal covariates that affect p) the we may write  $y_{ik} \sim \text{Bernoulli}(p)$ . Consequently, for this very simple model in which are independent and identically distributed ("itd") Bernoulli random variables and dent within individuals and among individuals. Formally, this allows us to say that yik Table 4.1. Except where noted explicitly, we suppose that observations are indepen $k=1,\,2,\,\ldots,\,K$  . Usually this is organized as a row of a matrix with elements  $y_{ik}$ , see y, for each individual (i = 1, 2, ..., N) is a vector having elements denoted by yik for would have an encounter history bt the form  $\sqrt{y} = (0, 1, 1, 0, 0)$ . Thus, the observation occasions, then an individual captured during occasions 2 and 3 but not otherwise not (y = 0) during any sampling occasion. As an example, suppose K = 3 sampling and these are of the form of a sequence of 0's and 1's indicating capture (y = 1) or subsequently recaptured. We suppose that individual encounter histories are obtained, "occasions," such as trap nights, where individuals are captured, marked, released, and a population of N individuals which we subject to repeated sampling, say over K To start looking at the simplest capture-recapture model, let's suppose there exists

 $(\lambda, q)$  laimonid  $\sim \lambda_i \chi$ 

individuals in the population (i.e., population size). for every individual in the population i = 1, 2, ..., N, where N is the number of

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(the total fumber of detections, y/), which is shown in the rightmost column. the binary detection history data can be summarized in the detection frequency K=5 sample occasions. Under a model with constant encounter probability, Table 4.1 A toy capture-recapture data set with n=6 observed individuals and

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 $y_i = \sum_j y_{ij}, y_{ii} \neq \sum_j \sum_j y_{ij}, \text{etc.}$ coođu

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4.1 The Simplest Closed Population Model: Model Mo

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recapture models discussed in this book. of almost all of classical capture-recapture models, including many spatial captureindividual is either encountered in a sample, or not-which forms the cornerstone We emphasize the central importance of the basic Bernoulli encounter model—an \$200g

Assuming individuals in the population are encountered independently, the joint standard methods that you are already familiar with (i.e., GLMs - see Chapter 3) we can specify a simple model and then deal with the fact that W is unknown sing estimating W is actually the central objective. But, by proceeding as if W were known, In fact, we are proceeding here as if we knew M. In practice we don't, of course, and capture-recapture models are just logistic regressions is a slight oversimplification. regression model with only an intercept term  $(\log \operatorname{it}(p) = \operatorname{constant})$  To say that all Evidently, the basic capture-recapture model is a simplistic version of a logistic-

probability distribution of the observations is the product of M binomials

(4.1.1) 
$$\prod_{i=1}^{N} \operatorname{Binomial}(y_i|K, p).$$

model using a BUGS engine poses no difficulty. this is a binomial regression model of the variety described in Chapter 3, fitting this the  $y_i = 0$  observations and the resulting data are just iid binomial counts. Because We emphasize that this expression is conditional on N, in which case we get to observe

not captured. In this case, an alternative and equivalent expression to Eq. (4.1.1) is important to note is that if we know N, then we know no, i.e., the number of individuals illustration, we converted the data-from Table 4.1 to this format (Table 4.2). What is indicator function evaluating to Tilife argument is true and 0 otherwise. For sake of exactly k times after K survey occasions,  $n_k = \sum_{i=1}^N I(y_i = k)$  where I() is the encounter frequencies. Specifically, Jet.n., denote the number of individuals captured Equation (4.1.1) can be simplified even further if we reformat the observations as

capture frequencies  $n_k$  are no longer independent because  $n_0$  is a function of the other the number of uncaptured individuals (n0) is unknown. Consequently, the observed The essential problem in capture-recapture, however, is that N is not known because where  $\pi_k = \Pr(y = k)$  under the binomial model with parameter p and sample size K .

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~90D frequencies,  $n_0 = N - \sum_{k=1}^K n_k$ . Hence, their joint distribution is multinomial (e.g., see Illian et al. (2008, p. 61)):

 $(n_0, n_1, \dots, n_K)^{\sim}$  Multinomial  $(N, \pi_0, \pi_1, \dots, \pi_K)$ (£.1.4)

history probabilities. refer to as "cell probabilities" or in the specific context of capture-recapture, encounter not encountered at all. These possible outcomes occur with probabilities  $\pi_k$ , which we defined by the possible encounter frequencies: encountered  $y=1,2,\ldots,K$  times or the multinomial posits a population of N individuals with K + I possible outcomes number (K+1) in this case) of possible categories. In the context of capture-recapture, nomial distribution is the standard model for discrete responses that can fall into a fixed We gave a general overview of the multinomial distribution in Section 2.2. The multi-

for differences in detection among maividuals, which we address subsequently in this formulation of the model that can accommodate individual-level covariates to account many instances, including for spatial capture-recapture (SCR) models, we require a frequencies, which entail that capture probabilities are the same for all individuals. In because we seldom are concerned with models for the aggregated encounter history nomial model is straightforward, but that is not always sufficiently useful in practice maximize the multinomial likelihood directly. Direct likelihood analysis of the multi-To fit the model in which N is unknown, we can regard no as a parameter and

chapter, and in Chapter 7.

4.1.1 The core capture-recapture assumptions

closure, Otis et al (1978) list the following: logical and statistical assumptions. In addition to the basic assumption of population This basic capture-recapture model—model Mo—comes with it a host of specific bio-

1. animals do not lose their marks during the experiment,

3. each animal has a constant and equal probability of capture on each trapping 2. all marks are correctly noted and recorded at each trapping occasion, and

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make clear statements such as occur, but it is not necessary to enumerate each possibility. Rather, it is necessary to example, closed population models also assume that other sources of error do not is both redundant and impossible to list all the things not allowed by the model. For is that all model assumptions are apparent when a model is clearly specified, and it following chapters, we refrain from repeatedly making such statements. Our opinion to be valid, and while they are also assumed by most of the models we present in the assumptions I and 2 are undoubtedly necessary for inference from basic CR methods The remainder of their classic work is dedicated to relaxing assumption 3. While

$$N, \ldots, I = i \text{ for } i = 1, \ldots, N$$

4.1 The Simplest Closed Population Model: Model Mo

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approximation to truth—but actual truth. on our model, statistical inference proceeds by assuming the model is truth—not an be discarded in favor of a more realistic alternative. However, once we have settled biological and sampling issues. If this assumption is deemed false, the model can eters and obtain parameter estimates; however, whether or not it is true depends upon encounter. Under this assumption, it is easy to write down the likelihood of the paramnot affect the encounter of another individual, and encounter does not affect future is critical for valid inference. It implies that the encounter of one individual does show the iid symbol, it will be assumed unless otherwise noted, and this assumption leaves very little left to say with respect to assumptions. Although we will not always This simple model description carries a tremendous amount of information, and it

and time. Furthermore, in practice, we will rarely have enough data to estimate the then we will never be able to study general phenomena that hold true across space ecological hypotheses. If we make these assumptions too complex or too specific, as a necessary evil. In fact, one way to view assumptions is as embodiments of our that all models are wrong due to their assumptions, assumptions should not be viewed In spite of the fact that we assume that all models are fruth, but we acknowledge

parameters of highly complex models.

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4.1.2 Conditional likelihood

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provide equivalent inferences, at least in large sample sizes (Sanathanan, 1972). They provide equally valid descriptions of the data and, for many practical purposes tional models are legitimate modes of analysis in all capture-recapture types of studies. we cannot observe the value y = 0 in the data set. Both the conditional and uncondidistribution is a "zero truncated" binomial distribution which accounts for the fact that based on  $\Pr(y|y) > 0$ . For the simple case of model M<sub>0</sub>, the resulting conditional random variable withat is, if and only if yw 0. The observation model is therefore have to recognize that individuals appear or not in the sample based on the value of the the event that an individual is captured at least once. To construct this likelihood, we in which the likelihood of the observed encounter histories is devised conditional on 2002). This formulation differs from the so-called "conditional likelihood" approach standerings model in place of "ikelihood" (Sanathanan, 1972; Borchers et al., to as the "joint likelihood" the "full likelihood," or the "unconditional likelihood" parameter NcThis multinomial model, being conditional on N, is sometimes referred known and, when W is unknown, the model is multinomial with index or sample size We saw that the closed population model is a simple logistic regression model if N is

(2) zero-truncated binomial that arises "conditional on n"; and (3) the zero-inflated models based on the (1) binomial model for the joint or unconditional specification; mulations of the model—or modes of analysis—for analyzing all capture-recapture distribution that we describe in the next section. Thus, there are three distinct fortinct formulation of capture-recapture models based on the zero-inflated binomial data augmentation (described in Section 4.2 below), which produces yet a third dis-In this book we emphasize Bayesian analysis of capture-recapture models using

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### CHAPTER 4 Closed Population Models

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Table 4.3 Modes of analysis of capture-recapture models. Closed population models can be analyzed using the joint or "full likelihood" which contains M, as an explicit parameter, the conditional likelihood which does not involve M, or by data augmentation which replaces M with ₩. Each approach yields a distinct likelihood.

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Isimonid bətsifini-orəz	ı∱ a	Data augmentation
zero-truncated binomial	d	Conditional likelihood
W xebni diw laimonillum	N,q	Joint likelihood
eters in model Statistical model	швява	Mode of analysis

binomial that arises under data augmentation. Each formulation has distinct model parameters (shown in Table 4.3 for model  $M_0$ ).

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We consider a method of analyzing closed population models using parameter-expanded data sugmentation (PX-DA), which we abbreviate to "data augmentation" or DA, which is useful for Bayesian analysis and, in particular, analysis of models using the various BUGS engines and other Bayesian model fitting software. Data sugmentation 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 work 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 was introduced for closed population models (Royle and Dorazio, 2008, 2012; Gardner (Royle, 2009b), open population models (Royle and Dorazio, 2008; Royle et al., 2010a), spatial capture-recapture models (Royle and Schaub (2012, Chapters 6 and 10) provide a good introduction to data augmentation in the context of closed and and 10) provide a good introduction to data augmentation in the context of closed and and 10) provide a good introduction to data augmentation in the context of closed and

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Onceptually, the technique of data augmentation represents a reparameterization of the "complete data" model—i.e., that conditional on M. The reparameterization is achieved by embedding this data set into a larger data set having M > N "rows" (individuals) and re-expressing the model conditional on M instead of M. The great thing about data augmentation is that we do not need to know M for this reparameterization. Although this has a whiff of arbitrariness or even outright ad hockery for it is always possible, in practice, to choose M pretty easily for a given problem and context and results will be insensitive to choice of M. Then, a given problem and context and results will be insensitive to choice of M. Then, under data sugmentation, analysis is focused on the "sugmented data set." That is, we under data sugmentation, analysis is focused on the "sugmented data set." That is, we

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<sup>2</sup>Unless the data set is sufficiently small that parameters are weakly identified.

analyze the bigger data set—the one having M rows—with an appropriate model that accounts for the augmentation. This is achieved by a Bernoulli sampling process that determines whether an individual in M is also a member of N. Inference is focused directly on estimating the proportion  $\psi = E[N]/M$  instead of directly on where  $\psi$  is the "data augmentation parameter."

## 4.2.1 DA links occupancy models and closed population models

There is a close correspondence between so-called "occupancy" models and closed population models (see Royle and Dorazio, 2008, Section 5:6). 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 the sites. This yields encounter data such as that illustrated in the left panel of Table 4.4. The important problem is that a species may occur at a site, but go undetected, yielding an all-zero encounter history for the site, which in the case of occupancy studies, are observed.

Table 4.4 Hypothetical occupancy data set (left), capture-recapture data in standard form (center), and capture recapture data augmented with all-zero capture histories (right).

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In contrast to occupancy studies, in classical closed gopulation studies, we observe mally, inference is focused on the parameter  $\psi$ , the probability that a site is occupied. sites and the latter with occupied sites where the species went undetected. More for-"sampling" (or stochastic) zeros, where the former are associated with unoccupied in a sense, the inference problem is to partition the zeros into "structural" (fixed) and in fact does occur. Thus, while the zeros are observed, there are too many of them and, However, some of the zero vectors will typically correspond to sites where the species

to the data set and regarding the augmented data set, essentially, as a site-occupancy can analyze capture-recapture models by adding M-n all-zero encounter histories induces a uniform prior for N on the integers [0, M] (Royle et al., 2007). Then, one of M for capture-recapture models may notibe obvious. However, the choice of M M is set by design (i.e., the number of sites in the sample), whereas a natural choice the closed population model. The only distinction being that, in occupancy models, precisely the same for both types of problems if an upper limit M is specified for should be—in a "complete" data set. This objective (how many sampling zeros?) is ence problem is, essentially, to estimate how many sampling zeros there are—or a data set as in the middle panel of Table 4.4 where no zeros are observed. The infer-

data set, where the occupancy or data augmentation parameter  $(\psi)$  takes the place of

Thus, the heuristic motivation of data augmentation is to fix the size of the data the abundance parameter (V).

To analyze the augmented data set, we recognize that it is a zero-inflated version non-biological sense) into the population, for example during an analysis by MCMC. M-n all-zero histories" potential individuals" because they exist to be recruited (in a but that were underected by sampling) at which detections did not occur. We call these "occupied sites" (in capture-recapture, individuals that are members of the population individuals that are not members of the real population that was sampled) as well as type model which includes both "unoccupied sites" (in capture-recapture, augmented right panel of Table 4.4, and then analyze the augmented data set using an occupancyset by adding too many all-zero encounter histories to create the data set shown in the

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the following set of assumptions (we include typical priors for a Bayesian analysis): binomial model which arises under data augmentation can be formally expressed by is the probability of a structural zero in the augmented data set. The zero-inflated data set of size M is a member of the sampled population—in the sense that  $1-\psi$ assume that  $z_i\sim {
m Bernoulli}(\psi)$  where  $\psi$  is the probability that an individual in the We member of the population of N individuals exposed to sampling. We variables,  $z_1, z_2, \ldots, z_M$ , to indicate whether each individual i is  $(z_i = 1)$  or is not binomial model can be described "hierarchically," by introducing a set of binary latent data set of size M-is a simple zero-inflated binomial likelihood. The zero-inflated population model, the resulting likelihood under data augmentation—that is, for the zeros, which do not correspond to individuals in the population. For a basic closed zeros (corresponding to actual individuals that were missed) and some are structural of the known-Waata set. That is, some of the augmented all-zero rows are sampling

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$$0 = 0$$
  $0 = I(y = 0)$   
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Bernoulli( $\psi$ )
 $i = iz$ 
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to express the conditional-on-z observation model concisely in just one step: for  $i=1,\ldots,M$ , where I(y=0) is a point mass at y=0. It is sometimes convenient

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$$|y_i|z_i \sim \text{Binomial}(K, z_i p)$$

and we understand this to mean, if  $z_i = 0$ , then  $y_i$  is necessarily 0 because its success

probability is  $z_i p = 0$ .

a function of these latent variables: under the assumptions of the zero-inflated model,  $z_i \stackrel{iid}{\sim} Bernoulli(\psi)$ ; therefore, N is model. In its place, we estimate wand functions of the latent variables z. In particular, Note that, under data augmentation, N is no longer an explicit parameter of this

$$\frac{1}{2}\sum_{i=1}^{M}=N$$

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

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

(1.2.4)

on-z" form using methods of MCMC either in the BUGS engines or using our own and Dorazio, 2012). Normally, however, we will analyze the model in its "conditionalobtain an estimator of Musing the so-called "Best unbiased predictor" (see Royle (e.g., see Royle, 2006). We could estimate these parameters and then use them to MLEs of the structural parameters  $\psi$  and p or those of other more complex models Interpreted as a likelihood, we can directly maximize this expression to obtain the

MCMC algorithms (see Chapter 17).

is shown in Panel 4.1. The last line of the model specification provides the expression  $z_i \sim {
m Bernoulli}(\psi)$ . The BUGS model description of the closed population model M<sub>0</sub> inflated model of the augmented data combines the model of the latent variables, represent the encounter frequency for potential individuals  $y_{n+1}, \ldots, y_M$ . The zerovector of frequencies (y<sub>1</sub>, ..., y<sub>n</sub>, 0, 0, ..., 0) where the augmented values of y = 0to individual-specific encounter frequencies, the augmented data are given by the menting model Mo. For this model, in which we can aggregate the encounter data polito It is helpful to understand data augmentation by seeing what its effect is on imple-4.2.2 Model Mo in BUGS

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#### PANEL 4.1

Model  $M_0$  under data augmentation. Here y,  $\kappa$ ,  $\min$ d, and  $\max$  are provided as data. The population size,  $M_0$  is computed as a function of the data augmentation variables z.

for computing N from the data augmentation variables  $z_i$ . Note that, to improve readability of code snippets (especially of large ones), we will sometimes deviate from our standard notation abit. In this case we use  $n \pm nd$  for n (the number of encountered individuals), and  $M = n \pm nd + nz$  is the total size of the augmented data set. In other cases we might also use noce in place of K and  $n \pm r \approx ps$  in place of J. We find that word definitions make code easier to understand, especially without having to read surrounding text.

Specification of a more general model in terms of the individual encounter observations yie 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 quan-

In this manner, it is straightforward to incorporate covariates on p for both individuals and sampling occasions (see discussion of this below and Chapter 7) as well as to devise other extensions of the model, including models for open populations

(see Chapter 16).

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M. One way of inducing the Uniform(0, M) prior on N is by assuming the following the posterior associated with this prior is equal to the likelihood for sufficiently large a uniform prior on N. The Uniform(0, M) prior for N is innocuous in the sense that inference problems with unknown N can be justified as originating from the choice of Use of parameter-expanded data augmentation (PX-DA), or DA for short, for solving

pierarchical prior:

(4.2.2)

 $\sim 0$ mrotin $0 \sim \psi$  $(\psi, M)$ lsimoni $\mathbb{A} \sim N$ 

that is appropriate for the augmented data set of known size M: Eq. (4.2.2), may be combined to yield a reparameterization of the conventional model The model assumptions, specifically the multinomial model (Eq. (4.1.3)) and

 $(\lambda \pi \psi, \dots, \zeta \pi \psi, I \pi \psi, M)$ lsimoniilu $M \sim (\lambda n, \dots, \zeta n, I n)$  $(\xi, 2, 4)$ 

for M. model, which is conditional on W, by integrating over a binomial prior distribution arise formally by removing the parameter Wefrom the ordinary closed population mial prior distribution for M. Thus, the models we analyze under data augmentation This expression arises by removing N from Eq. (4.1.3) by integrating over the bino-

data augmentation parameter. However, the full likelihood containing both N and  $\psi$ model by marginalization (integration) and replaced with the new parameter \$\psi\$, the probability  $1-\psi$ ). In Eq. (4.2.3), W has been eliminated as a formal parameter of the N, and others that are "structural zeros" (occurring in the augmented data set with of individuals that are sampling zeros  $(\psi \pi_0)$ , and belong to the population of size ability  $\psi_{\overline{M}}(0) + (1-\psi)$ , indicating that these unobserved individuals are a mixture Note that the M-n unobserved individuals in the augmented data set have prob-

can also be analyzed (see Royle et al., 2007).

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Other such "integrated likelihood" models are sensible. For example, we could place could be obtained from the ordinary likelihood, integrated over a binomial prior. resulting model as the "binomial-integrated likelihood" to reflect that an estimator on-W) model (by summation over a binomial prior). As a result, we could refer to the capture-recapture models in which M is marginalized out of the ordinary (conditionalcapture-recapture models. However, it really is just a formal reparameterization of is unfamiliar to most people, even to many of those with substantial experience with poiss Data augmentation may seem like a strange and mysterious black-box, and likely it 4.2.4 Remarks on data augmentation

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this type of marginalization (over a Poisson prior) is done by the R package secr produces a likelihood in which A replaces M, instead of \$\psi\$ replacing \$M\$. We note that a Poisson prior on N with mean A and marginalize N over the Poisson prior. This

for analysis of spatial capture-recapture models (see Section 6.5.3).

nothing more sophisticated than that is required

rameters need to be sampled using a Metropolis-Hastings step (e.g., Chapter 17), but MCMC is not much more difficult for complicated models—sometimes the hyperpavariables, i.e., the collection of z's, can be sampled from Bernoulli full conditionals. tributions) that can be sampled from directly. Furthermore, the data augmentation the parameters p and  $\psi$  have known full conditional distributions (in fact, beta dis-Appenned In this case, DA converts model Mo to a basic occupancy model, and in BUGS. Consider the simplest context—analyzing model Mo using the occupancysampling. And, in particular, capture-recapture models become trivial to implement also fixed. As a result, MCMC is a relatively simple proposition using standard Gibbs set of fixed size, so that the parameter dimension in any capture-recapture model is We emphasize the motivation for data augmentation being that it produces a data pot40

be affected by M and so some optimal choice of M might exist (Copalaswamy, 2012). computational burden. It seems likely that the properties of the Markov chains should the posterior for W is not truncated, but it should not be too large due to the increased for some concern. The guiding principle is that it should be chosen large enough so that Potential sensitivity of parameter estimates to M (especially of M) might be cause

(King and Brooks, 2001; Durban and Elaton, 2005; King et al., 2008; Schoffeld and jump MCMC (RIMCMC) or other so-called "trans-dimensional" (TD) algorithms There are other approaches to analyzing models with unknown M, using reversible Formal analysis of this is needed.

details of the DA implementation are the same for all capture-recapture problems. Furthermore, data augmentation is often easier to implement than RJMCMC, and the Royle et al \$2012c; Royle and Converse, in review) which we consider in Chapter 14. develop higrarchical models that involve structure on M (Converse and Royle, 2012; However, despite that N is removed as an explicit parameter in DA, it is possible to the advantage that one can model N explicitly or consider alternative priors for N. iteration of the MCMC algorithm. TD/RIMCMC approaches might appear to have the parameter space is a function of M, and will therefore typically vary at each approaches seek to analyze the conditional-on-N model in which the dimension of on M and we (usually) analyze the unconditional model. The various TD/RJMCMC TD methods is that DA is used to create a distinctly new model that is unconditional Barker, 2008; Wright et al. 2009). What distinguishes DA from RJMCMC and related

### 4.2.5 Example: Black bear study on Fort Drum

ples each week for 8 weeks, thus we distinguished K=8 weekly sample intervals. during June and July 2006. Barbed wire traps were baited and checked for hair samries on 47 individuals obtained from an array of 38 baited "hair snares" (Figure 4.1) (2009) and Gardner et al. (2010b). The specific data used here are encounter histotion. These data have been analyzed in various forms by Wegan (2008); Gardner et al. Wegan of Cornell University and their colleagues at the Fort Drum Military Installalected at Fort Drum Military Installation in upstate New York by P.D. Curtis and M.T. To illustrate the analysis of model M<sub>0</sub> using data augmentation, we use a data set col-

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4.2 Data Augmentation

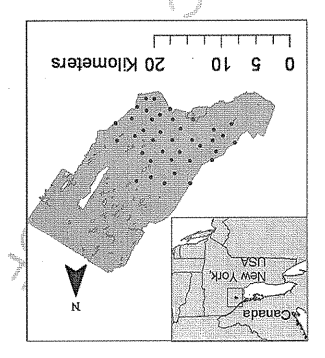


FIGURE 4.1

8 weeks during June and July, 2006. Fort Drum Black bear study area and the 38 baited hair snare locations operated for

with 128 all-zero encounter histories, resulting in a total sample size of M = 175. ?beardata for the commands to do the analysis). Here, the data were augmented (beardata) at the R prompt, and the analysis can be set up and run as follows (see The data are provided in the R package serbook, can be loaded by typing data

```
# p/c traditional CR models ignore space
                                                                              $1100
                                                              00110 > Y[Y > I] < - I
# toss out multiple encounters per occasion
                                              00102 > Y <- apply(Yaug,c(1,3),sum)
                   # summarize by ind x rep
                                      00100 > Yaug[1:nind,] <- beardata$bearArray
                                      00005 > Yaug < apray(0, dim = c(M,ntraps,K))
                                                               bmin-M -> sn < 00000
                                      00000 > ntraps <- dim(beardata$bearArray)[2]
                                           > K <- dim(beardata$bearArray)[3]
                                        00000 > nind -> dim(beardata$bearArray)[1]
                                               o0005 > trapmat -- beatdatagestrapmat
                                                            o0000 > data(beardata)
     # load the bear data and extract components
                                                           00022 > Jiprary(scrbook)
        108 MALL
```

 $\lambda = 1$  if individual encounter events (i.e.,  $\lambda_{ijk} = 1$  if individual i was The raw data object, beardata\$bearArray, is a three-dimensional array

encountered in trap j during occasion k, and 0 otherwise). For fitting model  $M_h$ , see below), it is sufficient to reduce the data to individual encounter frequencies which we have re-labeled 'Y' above. The **BUGS** model file along with commands to fit the model are as follows:

```
n.iter = 2000, n.burnin = 1000, n.thin = 1,debug = TRUE,working.directory = getwd()
00230
                    .t. and chara. n. chairs, parameo, model.file="modelMo.txt",n.chains = 3,
90229
                                                                                    ## gnu fue model:
00350
                                                                                ",file="modelM0.txt")
61200
01200
                                                                                       ([W:[]z)wns->N
00200
00200
$6100
                                                                       " dbin(tmp(i,k),l)
                                                                                           Y[1,k]
06100
                                                                              thg(i,k) <- p*z(i)
č8100
                                                                                     for(k in 1:K) {
08100
                                                                                  z[i] ~ dbern(psi)
SL100
                                                                                       }(M:1 ni i) rol
07100
                                                                                        (I'0) Jiump _ d
$9100
                                                                                     (I ,0)linub ~ izq
09100
                                                                                                 woger{
55100
                                                                                                 > csr ("
09100
                             list(z = zst, psi = runif(l) \frac{1}{2} p = runif(l) \frac{1}{2}
                                                                              } () uotabung -> satut <</pre>
$$100
                                                            .f ,bnin-M)monidx, (bnin, I) qey) > -> dez <
                                     * inice...
                                                      ((5)
04100
                                                                        > params0 <- c('pai','p','N')
                                          # parameters
25100
                                                                   > date < - list (y = y, M = M, K = K)
00130
                                     # Toad R2WinBUGS, set-up:
                                                                                  > Tiprary (RawinBuds)
90109
                                                                                       > set .seed(2013)
                      # to obtain the same results each time
00170
```

This produces the following posterior summary statistics:

> print(fito,digits = 2)
Inference for Bugs model at "modelMo.txt", fit using WinBUGS,
3 chains, each with 2000 iterations(first 1000 discarded)
n.sims = 3000 iterations saved

bavsa anoijsrāji 0065 = amis.n

делтаись S0'689 01.261 08.889 SP'08F 0011/1 8Z'II OL'ETS 00.48 00'TS 00.8₽ 00.7£ 66'I \$6'6\$ đ 60..0 08.0 25.0 08.0 82.0 52,0 58.0 92.0 0.22 FO. 0 62'0 98.0 TE:0 крас 85.2 **%**\$4.76 808 328

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.<sup>3</sup> 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.

[... some output deleted ...]

This is even a little suspicious....

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4.2 Data Augmentation

In order to obtain an estimate of density, D, we need an area to associate with the estimate of V, and in Chapter I we already went through a number of commonly used procedures to conjure up such an area, including buffering the trap array by the home range radius, often estimated by the mean maximum distance moved (MMDM) (Parmenter et al., 2003), I/2 MMDM (Dice, 1938) or directly from telemetry data (Wallace et al., 2003). Typically, the trap array is defined by the convex full around the trap locations, and this is what we applied a buffer to. We computed the buffer by using a telemetry-based estimate of the mean female home range radius (2.19 km) (Bales et al., 2005) instead of using an estimate based on our relatively more sparse recapture data. For the Fort Drum study, the convex hull has an area of LS7.135 km², and the buffered convex hull has an area of 277.011 km². To create this we used functions contained in the R package receos and created a utility function behaves which is in our R package ecrebook. The commands are as follows:

```
20500
                                         dyrea (bpl)
                                                          00500
plot (plch, border black', lwd = 2, add = TRUE)
                                                          26200
                            bjor(pbj' coj=, drsk()
                                                          06700
           pby <- (gBuffer(plch_width = buff))
                                                          28200
                         ъ́јср <- dcomo€xHull(p3)}
                                                          08200
           p3 <- SpatialPolygons(list(p2 = p2))
                                                          2720o
            DS \leftarrow Polygons(list(pl = pl)), ID = l)
                                                          00570
  pl <- Polygon(rbind(traplocs_braplocs[1,]))
                                                          59200
             00000 > bcharea <- function(buff, traplocs) {
                                      > Jiprary (rgeos)
```

The resulting buffered convex hull is shown in Figure 4.2.

omio > bcharea(2.19, traplocs = trapmat)

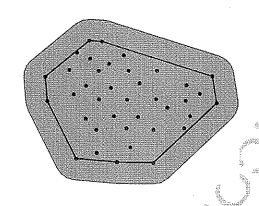


FIGURE 4.2

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Convex hull of the bear hair snare array at Fort Drum, NY, buffered by mean female home range radius (2.19 km).

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CHAPTER 4 Closed Population Models

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posterior summary of the ratio of N and the prescribed area (277.011 km²): To conjure up a density estimate under model Mo, we compute the appropriate polso

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L69T'0 T#8T:0 £081.0 S081.0 O.LLSO

Median

> TIO. (YYS/N\$Jail.amia\$0Jil) eliJnaup <

90320

Mean

1856£61.0 #89969T'0 %2.S

and not something we can formally evaluate (or estimate) from the data based on in our stated value of the "sampled area." Clearly though this is largely subjective, interval of (0.170, 0.195). Our estimate of density should be reliable if we have faith which yields a density estimate of about 0.18 ind/km2, and a 95% Bayesian confidence

# 4.3 Temporally varying and behavioral effects

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term  $\alpha_2 x_{ik}$ . Because spatial capture-recapture models allow us to include trap-specific  $\alpha_1 y_{i,k-1}$  or in any previous period (persistent behavioral response), described by the or ephemeral behavioral response; (Yang and Chao, 2005)), described by the term whether an individual was captured in the immediate previous period (a Markovian tured in any previous period). Therefore, encounter probability changes depending on where  $x_{ik}$  is a covariate indicator variable of previous capture (i.e.,  $x_{ik} = 1$  if cap-

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

Behavioral response is an important concept in animal studies because individuals polyo

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

"behavioral response" to trapping (even if the animal is not physically trapped). behavior. A common situation in many animal studies is that in which there exists a varying over time, as a function of date or season (Kery et al., 2010) due to species amount of search time, number of observers, or trap nights, or encounter probability responses that are relevant in many capture-recapture studies are "effort" such as such as when encounter probability varies over time among samples. Time-varying K occasions. Sometimes we can't aggregate the encounter data for each individual, vidual encounter frequencies—the number of times individuals are captured out of binomial model in capture-recapture and so we have considered models for indi-

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the BUCS language and we provide a number of examples in Chapter and elsewhere. capture in that trap. Models with temporal effects are easy to describe and analyze in the encounter probability is modified for an individual trap depending on previous response that is trap-specific (Royle et al., 2011b). In this local behavioral response, covariates, we can describe a third type of behavioral response-a local behavioral

Poles Models in which encounter probability varies by individual have a long history in 4.4 Models with individual heterogeneity

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model Mh see Royle and Dorazio (2008, Chapter 6) and Kéry and Schaub (2012, traps or other encounter mechanism. For additional background and applications of modeling heterogenetive due to the spatial organization of individuals in relation to ate models (See 4.5 below). Indeed, spatial capture-recapture models do just that, by model explicit factors that contribute to heterogeneity, e.g., using individual covariunderstood before fitting any such model. One solution to this problem is to seek to is not identifiable across classes of different heterogeneity models, and this should be produce wildly different estimates of W. In that sense, M for most practical purposes cases it's possible to find models that yield precisely the same expected data, yet Dorazio and Royle, 2003; Link, 2003). Indeed, Link (2003) showed that in some be extremely sensitive to the choice of heterogeneity model (Fienberg et al., 1999; lock, 1996; Pledger, 2004). One important practical matter is that estimates of N can various non-parametric approaches (Burnham and Overton, 1978; Norris and Polfor  $p_i$ . There are many different varieties of model  $M_h$  including parametric and broad class of models, each being distinguished by the specific distribution assumed models is often referred to as "model  $M_h$ " ("h" for heterogeneity), but really this is a to a CHAMM but in the capture-recapture context W is unknown. The basic class of assumption as  $p_i \not\sim \{\theta\}$  This type of model is similar in concept to extending a GLM tributed according to some probability distribution; [1] We denote this basic model individual-specific encounter probability parameters; p1, are random variables discapture-recapture models (Otis et al., 1978). Conceptually, we imagine that the capture-recapture and, indeed, this so-called "model Mr" is one of the elemental

We will work with a specific type of model Mh here which is a natural extension Chapter 6).

of the basic binomial observation model of model Mo so that

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random effect assumed to be normally distributed: where  $\mu$  is a fixed parameter (the mean) to be estimated, and  $\eta_i$  is an individual

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

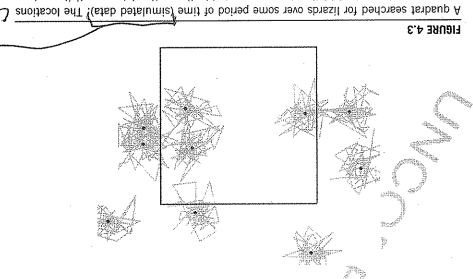
"logit-normal mixture" was analyzed by Coull and Agresti (1999) and elsewhere. It We could as well combine these two steps and write  $\log \operatorname{it}(p_i) \sim \operatorname{Normal}(\mu, \sigma_p^2)$ . This

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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 consider a beta prior distribution for  $p_1$  (Dorazio and Royle, 2003) and so-called "finite-mixture" models are also popular (Norris and Pollock, 1996; Pledger, 2004). In the latter, individuals are assumed to belong to a finite number of latent classes, each of which has its own capture probability.

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might model the exposure or availability of an individual to capture by supposing proportion of times the individual is within the sample plot boundaries, say  $\phi$ . We due to the overlap of its home range with the sampled area—essentially the long-run in the vicinity of the study area is liable to experience variable exposure to encounter dots) in some kind of random fashion. Heuristically, we imagine that each individual individuals move about their home range (home range centroids are given by the solid exhibits some sense of spatial fidelify in the form of a home range or territory, and quadrat searched repeatedly over a period of time. Further, suppose that the species for some species of reptile (see Royle and Young, 2008). Figure 4.3 shows a sample models. Imagine a quadrat that can be uniformly searched by a crew of biologists heterogeneity, we have to understand the relevance of movement in capture-recapture the spatial aspect of capture-recapture studies: To see how this juxtaposition induces area, it does yield an estimator that accommodates the heterogeneity in p induced by estimate M. While this doesn't resolve the problem of not knowing the effective sample thus it became common to use some version of model Mh in spatial trapping arrays to with the array of trap locations should yield heterogeneity in encounter probability, and (Karanth, 1995) because investigators recognized that the juxtaposition of individuals Model  $M_h$  has important historical relevance to spatial capture-recapture situations



A quadrat searched for lizards over some period of time (simulated data): The locations of encounter for each of 10 lizards are connected by lines—the dots are activity centers.

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of "random temporaty-emigration" (Kendall et al., 1997) where  $\phi_i$  is the individualbe individual-speciffe  $\phi_i$  Chandler et al., 2011). This system describes, precisely, that studies, it is natural that  $\phi$  should depend on where an individual lives, i.e., it should during any sample, and 0 otherwise. Then,  $\Pr(\alpha_i = 1) = \phi$ . In the context of spatial that  $a_i = 1$  if individual i is available to be captured (i.e., within the survey plot)

 $p_0$ , conditional on  $a_i = 1$ , then the marginal probability of detecting individual i is formally with auxiliary spatial information. If individuals are detected with probability to sampling due to movement in the proximity of the trapping array explicitly and Conceptually, SCR models aim to deal with this problem of vailable exposure specific probability of being "available" for capture.

 $!\phi 0d = !d$ 

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

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Royle (2003), and others data augmentation, has been considered by Coull and Agresti (1999), Dorazio and more traditional analysis of model Mh based on integrated likelihood, without using ventional likelihood methods as zero-inflated binomial mixtures (Royle, 2006) and we note that heterogeneity models formulated under DA are easily analyzed by contion. Although we use data augmentation in the context of Bayesian methods here, challenging. We address that here using Bayesian analysis based on data augmenta-Bayesian methods. However, begause W is not known, inference is somewhat more and Ware, 1982; Berger et al., 1999), which we discuss in Chapter 6, or standard tics using standard methods of inference based either on integrated likelihood (Laird binomial GLMM. This is a type of model that is widely applied throughout statispozis If W is known, it is worth taking note of the essential simplicity of model Mh as a

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

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a distribution for  $p_l$ . Here we assume and otherwise  $y_i|z_i=0 \sim I(y=0)$ , i.e., a point mass at y=0. Further, we prescribe

 $(^2$ o,  $\mu$ ) Normal $(\mu, \sigma^2)$ 

prior is to assume  $\tau = 1/\sigma^2 \sim Gamma(.1,.1)$ , although we usually choose  $\sigma \sim \sigma$ standard deviation  $\sigma \sim \mathrm{Uniform}(0,B)$  for some large B. Another common default For prior distributions we assume  $p_0 = \log i t^{-1}(\mu) \sim U$ niform(0, 1) and, for the

Uniform(0, B).

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# 4.4.2 Analysis of the Fort Drum data with model Mh

tion which we demonstrate here for setting up and running model Mh for the Port The function jage from the R2jage package runs essentially like the bugs functhe use of  ${
m JAGS}$  run from within  ${
m f R}$  using the useful packages R2jags or rjags. individuals. We have so far mostly used WinBUGS but we are now transitioning to heterogeneity model, and we used data augmentation to produce a data set of M=700Here we provide an analysis of the Fort Drum bear survey data using the logit-normal

Drum bear data:

```
> wbout ** jags (datal, inits, params), model.file = "modelMh.txt", n.chains
90442
                                                                                > TIPKSIA (RS] 938)
00440
                                        sigmap = runif(1,.7,1,2), lp = rnorm(M,-2)) }
                   . (I) lint = 0q ,0.=isq ,(!=<!), psi=10,() { list(z = as.numeric(y>=1), psi=2 -> sini <
06400

','ieq','qempis','0q') -> Lemeraq <</pre>
90459
                                             > data1 <- list(y = y nz = nz, nind = nind, K = K)
00420
                                                                           , file="ModelMh.t%t")
51100
01400
                                                                       ([(zn + baia):1]z)mue->W
20400
00400
                                                                        y[i] ~ dbin(mu[i],Κ)
                                             opeciarron model
26500
                                                                          [t]q*[t]s \rightarrow [t]um
06600
                                                                        ]od;r(b[;]) <- jb[;]
C8500
                                             individual effect
                                                                    lp[i] ~ dnorm(mup,taup)
08500
                                                                           z[i] ~ dbern(psi)
                                     zero fullation variables
$7£00
                                                                       }((zu + puru): t u; t) xo3
00370
                                                                     taup <- 1/(sigmap*sigmap)
psi ~ dunif(0,1)</pre>
59500
09600
                                                                           (01,0)linub ~ qsmgiz
55500
                                                                          wnb <- jod(b0/(j-b0))
00320
                                            # brior distributions
                                                                                (1,0)linub ~ 0q
54500
                                                                                            } Tabom
00340
                                                                                            ") JED <
25500
                                                                                 > sef.seed(2013)
06500
                                                                  [.... get data as before ....]
00372
```

posterior distribution for N shown in Figure 4.4. Posterior summaries of parameters million iterations (mixing is poor for this model and this data set), which produced the for spatial capture-recapture models. Using modelMhBUGS, we ran 3 chains of 1 algorithm for model Mh in Chapter 17, where we also develop MCMC algorithms for fun, we construct our own MCMC algorithm using a Metropolis-within-Cibbs firthe model using either JAGS or WinBUGS as specified by the user. In addition, We provide an R function model MABUGS in the package scrbook which will

iter = 1010000, n.burnin = 10000, working.directory = getwd())

We used M = 700 for this analysis and we note that while the posterior mass posses are given in Table 4.5.

extremely long right tail, with some MCMC draws at the upper boundary N=700, of N is concentrated away from this upper bound (Figure 4.4), the posterior has an

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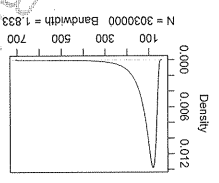


FIGURE 4.4

Posterior of M for Fort Drum bear study data under the legit-normal version of model M<sub>h</sub>.

posterior samples. 1010000 iterations, discarding the first 10000 for a total of three million bear data. Results were obtained using WinBUGS running 3 chains, each with Table 4.5 Posterior summaries from model Mn fitted to the Fort Drum black

						bring)
029	1.006	319,000	102,000	000.2a 768.6a	122.695	N
099	900.1	824.0	741.0	480.0 101.0	971.0	u w
820	1.003	878.8	2.025	312.1 738.0	2.096	d <sub>o</sub>
0+9	800.1	602.0	090 0	0.056 0.002	S70.0	0d
ŋə <sup>,</sup> u	16AA	%9'Z6	%0 <b>9</b>	. S.S.%	Mean	Parameter
NATIONAL SAME AND ADDRESS OF THE PARTY OF TH	MANAGEMENT OF THE PROPERTY OF	(1)	Sal.	. The state of the		

277.11 km<sup>2</sup>): rior distribution of D = N/277.11 (recall the buffered area of the convex hull is posterior distribution of density we produce the relevant summaries of the postesuggesting that an even higher value of M may be called for. To characterize the

(II. TTS/N\$Jail.amia\$Juodw).Yrammua < 22400

((279.0,02.0,20.0)p,tf.772\N\$paif.emia\$puodw)9firmsp < 00000

bears per square km and a 95% Bayesian credible interval is (0.224, 1.151). Therefore, the point estimate, characterized by the posterior median, is around 0.37

4.4.3 Comparison with MLE appreciation of N is highly skewed; therefore, we see that the posterior mean pozen

O 1056-OP CV = TRUE argument):

> N <- table (jout\$BUGSoutput\$sims listsN)

> xg <- as.numeric(names(N))

sp <- smooth.spline (xg,N,cy=TRUE)

MOOCH Spline (x = xg, y = N, cv = TRUE)

Smoothing Parameter apar = 00.09339815 lambda = 8.201724e-09 (17 iterations)

Equivalent Degrees of Preedom (Df): 121.1825

Definition Criterion: 2544481

Penalized Criterion: 2544481

00480 90482 00482

SL+00

07.500

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Penalized Criterion: 2544481

PRESS: 5903.4

PRESS: 5903.4

We obtain the mode of the smoothed frequencies as follows:

(1) 85 (2D\$\(\frac{1}{2}\) = \(\frac{1}{2}\) \(\frac{1}2\) \(\frac{1}2\) \(\frac{1}2\) \(\frac{1}2\) \(\frac{1

We don't dwell too much on the difference between the MLE and features of the posterior, but we do note here that the posterior distribution for the parameters of this model, for the Fort Drum data set, is very sensitive to the prior distributions. In the present case, the use of a Uniform(0, 1) prior for  $p_0 = \log i - 1(\mu)$  is somewhat informative—in particular, it is not at all "flat" on the scale of  $\mu$ , and this affects the posterior. We generally always recommend use of a Uniform(0, 1) prior for logit<sup>-1</sup>( $\mu$ ) in such models. That said, we were surprised at this result, and we experimented with other prior configurations including putting a flat prior on  $\mu$  directly. This kind of small sample instability has been widely noted in model  $M_h$  (Fienberg et al., 1999; Dorazio and Royle, 2003), as has extreme sensitivity to the specific form of model M<sub>h</sub> (Link, 2003). In summary, while the mode is well defined, the data set is relatively sparse and hence inferences are poor and sensitive to model choice.

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# Individual covariate models: toward spatial

capture-recapture

models, we begin with the basic binomial observation model: as "model  $M_x$ " (the x here being an explicit covariate). As with office closed population on closed population models, Kéry and Schaub (2012) referred to this class of models called "individual covariate models" but, in keeping with the classical nomenclature to influence encounter probability is measured for each individual. These are counter Pozza A standard situation in capture-recapture models is when a covariate which is thought

 $\gamma_i \sim \text{Binomial}(K, p_i)$ .

To model the covariate, we use a logit model for encounter probability of the form:

2008; Langtimm et al., 2011). group as a function of the observation-level individual covariate, "group size" (Royle, in models of aerial survey data, it is natural to model the detection probability of a such as body mass, or a discrete covariate such as group or cluster size. For example, probability are type of animal (juvenile/adult or male/female), a continuous covariate are the regression coefficients. Classical examples of covariates influencing detection where  $x_i$  is the covariate value for individual i and the parameters  $\alpha = (\alpha_0, \alpha_1)$ 

have some direct information about the latent variable, which comes from the spatial each captured individual. Unlike model  $M_h$ , in SCR models (and model  $M_x$ ) we do where the individual covariate is only observed imperfectly (or partially observed) for to the trapping array. Specifically, SCR models are individual covariate models, but model  $M_x$ , where the covariate describes where the individual is located in relation here because spatial capture-recapture models can be described precisely as a form of observed for the n individuals that appear in the sample. These models are important Model  $M_x$  is similar in structure to model  $M_h$ , except that the individual effects are

(1989), Alho (1990), and Borchers et al. (2002)). An estimator of N is ideas of unequal probability sampling (i.e., Horvitz-Thompson estimation<sup>5</sup>; Huggins Traditionally, estimation of N in model Mx is achieved using methods based on locations/distribution of individual recaptures.

$$\frac{1}{iq} \sum_{i=1}^{n} = \hat{N}$$

combnied as: is  $\vec{p}_i = \Pr(y_i > 0)$  and, in closed population capture-recapture models, it can be where pi is the probability that individual i appeared in the sample. This quantity

$$^{N}(iq-1)-1=(0$$

parameters are estimated from the conditional likelihood of the observed encounter where  $p_i$  is a function of parameters  $\alpha_0$  and  $\alpha_1$  according to Eq. (4.5.1). In practice,

npozo SPor a quick summary of the idea see: http://en.wikipedia.org/wiki/Horvitz-Thompson\_estimator.

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### CHAPTER 4 Closed Population Models

histories which is, for observation yi,

$$\mathcal{L}_{\mathcal{C}}(\boldsymbol{\omega}|\boldsymbol{y}_{i}) = \frac{|\boldsymbol{\omega}|_{\boldsymbol{y}_{i}}|\boldsymbol{\omega}|_{\boldsymbol{y}_{i}}|\boldsymbol{\omega}|_{\boldsymbol{z}_{i}}}{i\tilde{q}} = (i\boldsymbol{y}|\boldsymbol{\omega})_{o}\boldsymbol{\omega}$$

This derives from a straightforward application of the law of testal probability. Conceptually, we partition  $\Pr(y)$  according to  $\Pr(y) = \Pr(y|y > 0) \Pr(y > 0) + \Pr(y|y = 0)$  For any positive value of y the second term is necessarily 0, and so we rearrange to obtain  $\Pr(y|y > 0) = \Pr(y) / \Pr(y > 0)$  which, in the specific case where  $\Pr(y)$  is the binomial probability mass function (pmf) produces Eq. (4.5.2).

Here we take a formal model-based approach to Bayesian analysis of such models based on the joint likelihood using data augmentation (Royle, 2009b). Classical likelihood analysis of the so-called "full likelihood" is covered by Borchers et al. (2002). For Bayesian analysis of model  $M_x$ , because the individual covariate is unobserved for the  $n_0 = N - n$  uncaptured individuals, we require a model to describe variation in x among individuals, essentially allowing the sample to be extrapolated to the population. For example, if we have a continuous trait measured on each individual, then we might assume that x has a hormal distribution:

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

Data sugmentation 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 following form, for each i=1,2,...,M:

$$z_i \sim ext{Bernoulli}(\psi)$$
 $\gamma_i | z_i = 1 \sim ext{Binomial}(K, p_i(x_i))$ 
 $\gamma_i | z_i = 1 \sim 1 (\gamma = 0)$ 
 $\gamma_i | z_i = 0 \sim 1 (\gamma = 0)$ 
 $x_i \sim ext{Normal}(\mu, \sigma^2)$ 

Fully spatial capture-recapture models 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 BUCS language. The BUCS specification is very similar to that for model  $M_{\rm h}$ , but we require the distribution of the covariate to be specified, along with priors for the parameters of that distribution.

### 4.51 Example: location of capture as a covariate

Here we consider a special type of model M<sub>x</sub> that is especially relevant to spatial capture-recapture. Intuitively, some measure of distance from home range center to traps for an individual should be a reasonable covariate to explain heterogeneity in encounter probability, i.e., individuals with more exposure to traps should have higher encounter probabilities and vice versa. So we can imagine estimating such a higher encounter probabilities and vice versa. So we can imagine estimating such a duantity, say average distance from home range center to "the trap array", and then using it as an individual covariate in capture-recapture models. A version of this idea

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4.5 Individual Covariate Models: Toward Spatial Capture-Recapture 111

that SCR models represent a formalization of this heuristic procedure. ill-defined sample area. After you read later chapters of this book, it will be apparent distribution on the individual covariate, we will resolve the problem of having an Boulanger and McLellan (2001). By analyzing the full likelihood and placing a prior by data augmentation. We take a slightly different approach than that adopted by a fully model-based analysis of the version of model Mx described above, analyzed individual's home range center. We provide an example of this type of approach using that the average location of capture would be a decent estimate (heuristically) of an of an ideal situation where we have a dense grid of traps over some geographic region, capture location. This is intuitively appealing because we can imagine, in some kind edge" of the trapping array, where the home range center is estimated by the average the Huggins-Alho estimator and with covariate "distance from home range center to was put forth by Boulanger and McLellan (2001) (see also Ivan, 2012), but using

range center but, clearly this is biased for individuals that live around the edge (or we might expect the sample mean encounter location to be a good estimate of home of individual covariate models in the literature. For an expansive, dense trapping grid observed mean encounter location—which, while ad hoc, is consistent with the use therefore decrease. We have defined si in terms of a sample quantity—the array should have a higher probability of encounter and, as x<sub>i</sub> increases, p<sub>i</sub> should "edge" itself is not precisely defined. Conceptually, individuals in the middle of the have used distance from edge of the trap array but that is less easy to quantify, as or magnitude of the vector u, and we use it throughout the book. In practice, people array,  $x_0$ :  $x_i = ||s_i - x_0||$ . Note that ||u|| is standard notation for Euclidean norm from the average encounter location of individual i, say s;, to the centroid of the trap For our purposes here, we define the scalar individual covariate  $x_i$  to be the distance

individual being captured by the trap array: 0 to some large number, say B, beyond which it would be difficult to imagine an would not be captured. Therefore, let's assume that  $x_i$  is uniformly distributed from and it seems sensible that an individual located extremely far from the array of traps distribution for it. As a measurement of distance we know it must be positive-valued, is also missing. Therefore, it is a latent variable, and we need to specify a probability A key point is that  $s_i$  is missing for each individual that is not encountered and so  $x_i$ off) the trapping array.

(a,0)mrolinU  $\sim ix$ 

centroid of the array. example; B should be at least a home range diameter past the furthest trap from the where B is a specified constant, which we may choose to be arbitrarily large. For

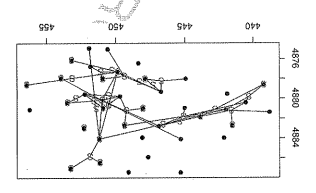
the centroid of the trapping array) using the R function spiderplot provided in the Fort Drum data. We need to compute the individual covariate  $x_{ij}$  (distance from We have to do a little bit of data processing to fit this individual covariate model to

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4.5.2 Fort Drum bear study



#### FIGURE 4.5

home range center with a line. home range center. All traps in which a bear was captured are connected to its estimated with the "S" symbols being the average capture location of each bear, i.e., its estimated Spider plot of the Fort Drum study data. The black dots represent the 47 trap locations

spider plot are as follows: from trap centroid" (the variable xeent returned by spiderplot) and making the call a "spider plot." The R commands for obtaining the individual covariate "distance SCYDOOK. This function also produces the keen plot shown in Figure 4.5 which we

> xceur <- posqsxceur 04500 > toad <- Sepiderplot (beardata\$bearArray, beardata\$trapmat) 0033 > data(beardata) 06200 > Jiprary(scrbook)

P0295

and fit the model are as follows: model reside. The BUCS model specification and R commands to package the data with our choice of B to reflect the changing area over which the W individuals of the the population of M individuals resides. We will see shortly that M does, in fact, scale units of the trap centroid. Therefore, the model associates a precise area within which implication is that the population size parameter, W, applies to the area within 11.5 from the array center to the furthest trap. Once we choose a value for B, the direct troid" we used  $x_{\rm g} \sim {\rm Uniform}(0,B)$  with  $B=11.5~{\rm km}^2$ , which is about the distance For the analysis of these data using the individual covariate "distance from cen-

beta ~ dnorm(0,.01) 00210 (I,0)linub ~ isq 59500 alpha0 -- log(p0/(l-p0)) 09500 (I,0)linub ~ 0q # prior distributions  $\varsigma \varsigma \varsigma \varsigma 0 o$ model { 05500 ಇಕರ (" 54500

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### 4.5 Individual Covariate Models: Toward Spatial Capture-Recapture

```
This produces the posterior summary statistics in Table 4.6.
                                    n.chains = 3, n.iter = 11000, n.burnin = 1000, n.thin = 1)
                                                                                                              00900
                                         fit2 <- bugs (data2, inits, parama2, model.file="modelMcov.txt",
                                                                                                              51900
                              inits <- function() (list(z = zst, psi = psi, p0 = runif(l), beta = rnorm(l)
                                                                      params2 <- c('po','psi','N','beta')
                                                                                                              56900
                                    data2 <- list(y = y,nz = nz, nind = nind, K = K, xcent, B = ll.5)
                                                                                                              06900
                                                                                    [ile="modelMcov.txt"]
                                                                                                              $2900
                                                                                                              00900
                                                                                 ([(zu + puțu):[]z)wns ->
                                                                                                              $1900
DEIP
DEIP
                                                                                                              01900
                                                                                    Y[i] ~ dbin(mu[i],K)
                                                                                                              50900
                                                                                      [t]q^{*}[t]s -> [t]um
                                                                                                              00900
                                                                                    logit(p[i]) <- lp[i]
                                                                                                              56500
                                                                        jp[i] <- alpha0 + beta*xcent[i]</pre>
                                                 # individual effect
                                                                                                              06900
                                                                                       z[i] _ qpexu(bai)
                                                                                                              58500
                                                          # Dy Astisbles
                                                                                   xcent[i] ~ dunif(0,B)
                                                                                                              08500
                                                                                   )((zu + puru): t ur r) xog $2500
```

notwithstanding, it is worth pondering how this model could be an improvement there is a good explanation for this which we discuss in the next section. That issue

We note that the estimated M is much lower than obtained by model M<sub>h</sub> but

as the number of individuals with home range centers less than B from the centroid component of the model and the parameter N therefore has explicit spatial context, explicit area which is defined by our prescribed value of B. That is, this area is a fixed In addition, and importantly, using our new model, the estimated N applies to an on an explicit mechanism, as opposed to a phenomenological one such as model Mh. by the centroid of the array. Moreover, we have done so using a model that is based due to spatial location of individuals relative to exposure to the trap array, characterized considered previously. Well, for one, we have accounted formally for heterogeneity (conceptually or technically) over some other model/estimator including Mo and Mr

of the trap array. As such the implied "effective area" of the trap array for a given B

is a precisely defined quantity—it is that of a circle with radius B.

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30000 posterior samples. 3 chains, each with 11000 iterations, discarding the first 1000 for a total of Fort Drum black bear data. Results were obtained using WinBUGS running M<sub>x</sub>) with covariate "distance from the centroid of the trap array," fitted to the Table 4.6 Posterior summaries from the individual covariate model (model

Jal	100					J	57 N9 D
087		Sr.0-	92:0-	96.0-	90.0	9Z.0	-FA
0061	i	00.17	00.88	00.03	67.3	26,83	. N
3200	l	44.0	48.0	92.0	90.0	46.0	1/4
1100	L	79.0	49.0	04.0	70.0	46.0	0d
iie.n	16dA	%9'26	%09	5'2%	as	Mean	Parameter
						duina iaila	and anana

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4.5.3 Extension of the model

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than that obtained previously from model Mh (Section 4.2) and also why, if we were distance band is increasing. This is one reason we have a lower estimate of density constant N in each distance band from the centroid but obyiously the area of each move away from the centroid of the trap array. That is,  $x_i \sim U$ niform(0, B) implies constant over space. In particular, this model implies that density decreases as we tant reason: imposing a uniform prior distribution on x implies that density is not The model developed in the previous section is not a very good model for one impor-

Fortunately, we are not restricted to use of this specific distribution for the indito increase B, we would see density continue to decrease.

centroid). This can be verified empirically as follows: in each additional distance band increases linearly with radius (i.e., distance from is a sort of triangular distribution in density induced because the incremental area  $Pr(x) = Pr(X < x) = (x > 1)/(\pi x^2)/(\pi x^2)$ . Then, f(x) = dF/dx = 2(x > X)which tells us we should choose  $[x] = 2x/B^2$ . This can be derived by noting that farther away from the centroid, so that density is constant. In fact, theory exists be proportional to the amount of area in each successive distance band as you move do is impose a prior on distance from the centroid, x, such that abundance should ing density as distance from the centroid increases. Conceptually, what we want to whether we can choose a better distribution for B = one that doesn't imply a decreasvidual covariate. Clearly, it is a bad choice and therefore, we should think about

[8 (D) investor sat (\$\forallag (0\) > abline (0\) (BUAT = Ylilidsdorq,001, [T > b]b) faid < ([T > p]p) laid < > q < - zdxc (n + n + A + A)(I'I-'0000t) Jrunz -> A < (t't=00000t) frunz -> n <

there is not a built-in way to do so. However, we can implement a discrete ver-It would be useful if we could describe this distribution directly in BUGS but OZEOd

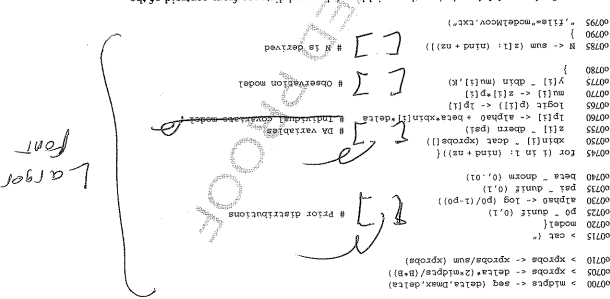
this (noting that we have already loaded and processed the Fort Drum bear data) are pure once and then pass them to BUGS as data. The R commands for doing all of the interval probabilities are, approximately,  $p_i = \delta(2m_i/B^2)$ , which we can com-Similar are  $m_1 = g_1 + g_2 + g_3 + g_4 + g_5 + g_6 + g_6$ probabilities proportional to 2 \* x. In particular, if we denote the cut-points by sion of the pdf.6 To do this, we break binto L distance classes of width 8, with

given in the following R/BUGS script:

<sup>6</sup>We might also be able to use what is referred to in WinBUGS jargon as the "zeros trick" (see Advanced 0£0qa > xpin <- xcent%kdelta + 1 \$6900 5. delta <-.2 06900

This is just length  $\times$  width, the area of small rectangles approximating the integral. BUGS tricks in the manual) although we haven't pursued this approach.

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previous analyses of this chapter, we do this: the model, keeping in mind that the data objects required below have been defined in distance in the expression for Lp [1]; xbin[1] has to be multiplied by 8. To fit xbin is then the integer category in units of 8 from 0. Thus, to convert back to version of the pdf of x can be used, as described previously. The new variable labeled trap array) has been rounded or binned (placed into a distance bin) so that the discrete In the model description, the variable x (observed distance from centroid of the

```
n.iter = IPOOO, n.burnin = 1000, n.thin = 2)
                                                                   06800
           working.directory = getwd (), debug = FALSE, n.chains = 3,
                                                                   62800
00820
                                                                   $1800
                                   > parama2 <- c('p0', 'pai', 'N', 'beta')
                                                                   01800
                                         delta #delta)
                                                                   20800
 > data2 <- list (y = y, ng = na nind = nind, K = K, xbin = xbin, xprobs = xprobs,
                                                                   00800
```

array) on upto 20. The results are given in Table 4.7. values of B from B = 12 (restricting values of x to be in close proximity to the trap appears invariant to B as long as B is sufficiently large. We fit the model for a set of find empirically, that while N is highly sensitive to the prescribed value of B, density the area under consideration, there should be more individuals in it. Fortunately, we W we want by changing B! However, it is intuitively reasonable that, as we increase our choice of B. This might be disconcerting to some—we can get whatever value of size that applies to the particular value of B and, as such, we will see that N scales with population of Windividuals resides. The parameter N of the model is the population By specification of B, this model induces a clear definition of area in which the

appear insensitive to choice of B once we reach about B = 17 or so. The estimated We see that the posterior mean and SD of density (individuals per square km)

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### CHAPTER 4 Closed Population Models

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Table 4.7 Analysis of Fort Drum bear hair snare data using the individual covariate model, for different values of B, the upper limit of the uniform distribution of 'distance from centroid of the trap array'. "Density" is the posterior mean of density.

71000		Ino D	Cent.
440.0		0.250	l so
640.0	1929P ,,,1864;	0.250	, er
640.0		0.2 <del>4</del> 9	18
0.044	"Higgs scrap"	0.249	Z1
140.0		0.244	91
860.0	The state of the s	0.230	12
CS tolteteon	(ve	Density (post mea	
			mean of density,

that definity is invariant to the choice of area, which seems like a desirable feature. center and the centroid of the trap array. Under this definition of the system, we see space and that probability of detection depends on the distance between home range direct information; that individual home range centers are randomly distributed in centroid" that is a reasonable and standard null model—it posits, in the absence of other hand, the individual covariate model uses an explicit model for "distance from get exactly the same estimate of N no matter what we declare the area to be. On the model, based on a buffered "trap area," are justifiable by any theory. In fact, we would no reason at all to think that the estimates produced under either closed population which the population resides is not defined under either model. There is therefore of models Mo and Mh are distinctly not spatially explicit models—the area within really for comparing or contrasting these various estimates. In particular, application that reported from model  $M_0$  using the buffered area (S=0.18). There is no basis hand our estimate of D=0.25 here (based on the posterior mean) is higher than estimates were in the range of 032-043 and see Section 4.4 above. On the other make it up). Using MLEs of N in conjunction with buffer strips (see Table 1.1) our Mh for which no relevant "area" quantity is explicit in the model (and so we had to density of 0.25 per km2 is actually quite a bit lower than we reported using model

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### 4.5.4 Invariance of density to B

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Under model  $M_x$ , and under models that we consider in later chapters, a general property of the estimators is that while N increases with the prescribed area of the model (defined by B in this model), we expect that density estimators should be invariant to this area. In the model used above, we note that Area(B) =  $\pi$  B<sup>2</sup> and E(N(B)) =  $\lambda$ Area(B) and thus E(Density(B)) =  $\lambda$ , i.e., constant. This should be interpreted as the prior density. Absent data, then realizations under the model, will have density  $\lambda$  regardless of what B is prescribed to be. As we verified empirically have density  $\lambda$  regardless of what B is prescribed to be. As we verified empirically

above, posterior summaries of density are also invariant to B as long as the prescribed

area is sufficiently large.

#### 4.6 Distance Sampling: a Primitive SCR Model 411

# 4.5.5 Toward fully spatial capture-recapture models

define that) are biased because the observations are truncated—we can only observe s, is estimated. Finally, it ignores that estimates of stateund the "edge" (however we practice. Moreover, this approach also ignored the variable precision with which each and expansive relative to typical home range sizes which might not be reasonable in estimate of home range center makes sense but more so when the trapping grid is dense to the home range center sp. Intuitively, taking the average encounter location as an ter "data" being the average observed encounter location, and equated that summary model trap-specific effects (e.g., trap effort or type). Moreover, we applied this model and the locations of each individual encounter, so that we cannot use this model to does not make full use of the spatial information in the data set, i.e., the trap locations of a precise relationship between M and area), Is not ideal for all purposes because it inherent in almost all capture-recapture studies (induced heterogeneity and absence While the use of an individual covariate model resolves two important problems

However, there is hope to extend this model in order to resolve these remaining

locations interior to the array.

for si. This is actually easier, and less ad hoc in a number of respects, and you should adapted easily to deal with s; as a latent variable, simply by adding a prior distribution variables with an explicit model. We note that the model fitted previously could be and the observation locations (i.e., trap-specific encounters) are linked to those latent here. In that chapter we build a model in which si are regarded as latent variables covariate model that definitively resolves the adhoc nature of the approach we took deficiencies. In the next chapter we provide a further extension of this individual

### 4.6 Distance sampling: a primitive SCR model

it describes, explicitly, the spatial organization of individual locations (although this routinely for decades and, formally, it is a spatially explicit model in the sense that populations. This first and most basic spatial capture-recapture model has been used other individuals) repeatedly and so distance sampling can be applied to unmarked the need to explicitly identify individuals (except they need to be distinguished from vations are made at only K = I sampling occasion. Distance sampling eliminates fact, distance sampling is precisely an individual-covariate model, except that obserbetween an individual's location say u and the observation location or transect. In ulation model with an individual covariate. The covariate in this case, x, is the distance methodological context, the distance sampling model is a special case of a closed population models, distance sampling provides explicit estimates of density. In terms of 1980; Bucklandet alt, 2001; Buckland, 2004) because, unlike ordinary closed popis one of the most popular methods for estimating animal abundance (Burnham et al., assumption is that detection probability is a function of distance. Distance sampling surements of distance from an observer to individual animals (or groups). The basic posse Distance sampling is a class of methods for estimating animal density from mea-

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is not always stated explicitly) and, as a result, somewhat general models of how individuals are distributed in space can be specified (Hedley et al., 1999; Royle et al., 2004; Johnson, 2010; Niemi and Fernández, 2010; Sillett et al., 2012).

As with other models we've encountered in this chapter, the distance sampling model, under data augmentation, includes a set of M zero-inflation variables  $z_i$  and a binomial observation model expressed conditional on  $z_i$  (binomial for z=1, and fixed zeros for z=0). In distance sampling we pay for having only a single sample occasion (i.e., K=1) by requiring constraints on the model of detection probability, normally imposed as the assumption that detection probability is 1.0 when distance equals 0. A standard model for detection probability is the "half-normal" model:

$$\binom{7}{2}x\log - 3\log = \frac{1}{2}d$$

for  $\alpha_1 > 0$ , where  $x_1$  denotes the distance at which the *i*th individual is detected relative to some reference location where perfect detectability (p = 1) is assumed. This encounter probability model is more often written with  $\alpha_1 = 1/2\sigma^2$ . If K > 1 then an intercept in this model, say  $\alpha_0$ , is identifiable and such models are usually called "capture-recapture distance sampling" (Alpizar-Jara and Pollock, 1996; Borchers et al., 1998).

As with previous examples, we require a distribution for the individual covariate poses

zi. The customary choice is

PANEL 4.2

$$(a,0)$$
mrotin $U \sim \sqrt{x}$ 

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). Specification of this distance sampling model in the BUGS language is shown in Panel 4.2, taken from Royle and Dorazio (2008).

# D = N/total area of transe D <- N/striparea ([(zn+bnin):1]s)mus -> N # N is a derived parameter ([i]um)nraedb ~ [i]y # Observation model [i]q\*[i]z -> [i]um (iaq)nrədb [i]s x [i]x x [i]x - x (iaqu) (iaqu) x [ii]q - x (iaqu) x (iaqu) - x (iaqu) ([i]x\*[i]x)\*lsdqls -# Defection inuction # B=strip width # DV variables }((za+baia):t al t) tol (1,0)linub " isq (01 0) lings ~ isaqis # Prior distributions

Distance sampling model in **BUGS** for a line transect situation, using a half-normal detection function.

4.6 Distance Sampling: a Primitive SCR Model

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Thus we can write the general distance sampling model as location instead of distance between individual and observation point (or transect). pling model can be equivalently specified by putting a prior distribution on individual As with the individual covariate model in the previous section, the distance sam-

along with

 $(1o'||0\mathbf{x} - |\mathbf{n}||) y = !d$ 

(S) mnolinU  $\sim 1$ u

distance, we provided the analogous argument in the previous section. Is in the surface of the form of the following the surface of the following surface of the foll a transect of length L is used and x is distance to the transect, then  $F(x) = \Pr(X \le 1)$ in space, then the distribution of Euclidean distance is also uniform. In particular, if location. Basic math can be used to argue that if individuals have a uniform distribution for the sample of n individuals. In practice it is easier to record distance instead of where  $x_0$  is a fixed point (or line) and  $u_1$  is the individual's location, which is observed

described, conceptually, by a hierarchical model of the form: terization of SCR models (Chapter 0.6), we suggested that every SCR model can be models can be described (Royle et alph 2011a). In the context of our general characa reduced information summary of location, u. Some intermediate forms of SCR/DS ous section, where the encounter probability is related directly to distance, which is spatial capture-recapture models, such as those derived from model Mx of the previ-The preceding paragraph makes it clear that digiance sampling is a special case of

 $\cdot [s][s|u][u|\sqrt{s}]$ 

model of the form the model components for the observed data u.8 Thus, we are left with a hierarchical Distance sampling ignores the part of the model pertaining to s, and deals only with

[n][n]

u and condition on sawhich is not observed: In contrast, as we will see in the next chapters, many SCR models (Chapter 5) ignore

[s][s|A]

distance sampling data on species such as Jaguars or tigers! Clearly, distance sampling and accounts for spatial heterogeneity in detection? For one, imagine trying to collect So why bother with SCR models when distance sampling yields density estimates and whetheror not the latent variables are observable (in distance sampling they are). models! The main differences have to do with interpretation of model components Since [u] said [s] are both assumed to be uniformly distributed, these are equivalent

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sible. For tigers, it is much easier, efficient, and safer to employ camera traps or track requires that one can collect large quantities of distance data, which is not always pos-

<sup>8</sup>Equivalently, we could also say that [u] in the distance sampling model is  $[u] = \int [u|s][s]ds$ .

plates and then apply SCR models. Furthermore, as we will see in Chapter 15, SCR models can make use of distance data, allowing us to study distribution, movement, and density. Thus, SCR models are more general and versatile than distance sampling models twhich clearly are a special ease, and can accommodate data from virtually all animal survey designs.

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### 4.6.1 Example: Sonoran desert tortoise study

We illustrate the application of distance sampling models using data on the Sonoran desert tortoise (Gopherus agassizii), shown in Figure 4.6, collected along transects in southern Arizona (see Zylstra et al. (2010) for details). The data are from 120 square transects having four 250-m sides, although we ignore this detail in our analysis here and regard them as 1 km transects, and we pooled the detection data from all 120 transects. The histogram of encounter distances from the 65 encountered individuals is shown in Figure 4.7.

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Commands for reading in and organizing the data for analysis using WinbUGS are given in the help file Ployloided with the scrbook package. To compute density, the total sampled area of the transects strtparea is input as data, and computed as: 120 (transects) multiplied by the length (1000 m) and halfwidth (B = 40 m), then multiplied by 2, and divided by 10000 to convert to units of individuals per ha. We also provide commands for analyzing the data with unmaxked individuals per ha. We also provide commands for analyzing the data with unmaxked (Fiske and Chandler, 2011) using hierarchical distance sampling models (Royle et al.,

Posterior summaries for the tortoise data are given in Table 4.8. Estimated density posterior mean) is 0.54 individuals per ha and the estimated scale parameter of the distance function (posterior mean) is  $\sigma = 9.12$  m. The R-hat statistics of around 1.02 distance function (posterior mean) is  $\sigma = 9.12$  m. The R-hat statistics of around 1.02

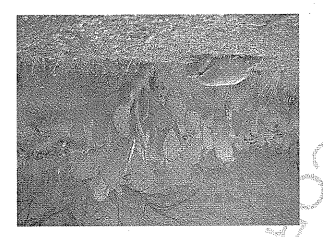
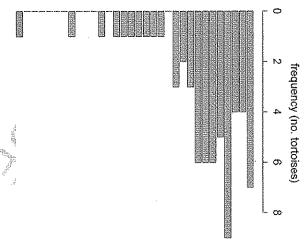


FIGURE 4.6

Desert tortoise in its native habitat (Photo credit: Erin Zylstra, Univ. of Arizona).

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qiarance (m) 11 13 16 17 19 21 23 26 27 29 31

FIGURE 4.7

of survey transect. Distance histogram of n=65 Sonoran desert fortoise detections from a total of 120 km

iterations and the first 1000 discarded, thinning by 2. Results were obtained using WinBUGS running 3 chains, each with 3000 Table 4.8 Posterior summaries from the tortoise distance sampling data.

00f 20.f 00.263 00.313 00.314 f7.43 70.313 00f 20.f 38.0 48.0 64.0 30.0 48.0 89 20.f 37.0 f8.0 94.0 70.0 f8.0	*****
001	
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	٨
051 Z0.1 7T.01 70.9 TT.7 TT.0 21.9	3
0£f S0.f f0.0 f0.0 00.0 00.0 f0.0	17

 $\psi=1$  and so the degree of data augmentation appears sufficient. mass of the data augmentation parameter  $\psi$  is located away from the upper bound suggest that slightly longer MCMC simulations might be called for. The posterior

# 4.7 Summary and outlook

generalized linear models. Indeed, the only real distinction is that in capture-recapture po400 Traditional closed population capture-recapture models are closely related to binomial

zero-inflated models is practically convenient, especially using the  ${f BUGS}$  variants. GLMs and GLMMs instead of ordinary GLMs or GLMMs. The analysis of such the unknown-N problem using data augmentation then we are left with zero-inflated vations can be modeled using a zero-inflated binomial distribution. When we deal with using data augmentation, which arises under a Uniform (0, M) prior for N, the obserencounter frequency data on observed individuals. If instead we analyze the models binomial (from which the so-called conditional likelihood derives) since we only have that the observations don't have a standard binomial distribution but, rather, a truncated analysis of capture-recapture models. The classical approach to inference recognizes thetical "complete" data set is unknown. This requires special consideration in the models, the population size parameter N corresponding also to the size of a hypo

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because it does not make full use of the spatial encounter histories, which provide prior on the individual covariate). The model is still not completely general, however, provides a direct estimate of density because area is a feature of the model (via the due to the spatial juxtaposition of individuals with the array of traps, and it also use of closed population, models: it allows for heterogeneity in encounter probability the centroid of the trap array). The model addressed some important problems in the estimator to the choice of population size area (induced by maximum distance from uniformly distributed in space. This assumption provides for invariance of the density covariate which was derived from an assumption that individual locations are, a priori, likelihood using data augmentation, and placed a prior distribution on the individual distance to the centroid of the trapping array as a covariate. We analyzed the full encounter location as an individual covariate. We did this in a novel way, by using observed locations of individuals—we can think of using some function of mean spatial capture-recapture models arise/by defining individual covariates based on of this book are closely related to individual covariate models (model  $M_x$ ). Naturally, Spatial capture-recapture models that we will consider in the rest of the chapters

model more closely resembles the SCR model we introduce in the next chapter. distribution on location, s, of each individual, then the form of the distance sampling model on distance in the classical distance sampling model, we were to place the prior and analyzing spatial capture-recapture models. In fact if, instead of placing an explicit individual covariate models more broadly provides a solid basis for understanding model—but one without replicate samples. Understanding distance sampling and sampling. The model underlying distance sampling is precisely a special kind of SCR A specific individual covariate model that is in widespread use is classical distance

direct information about the locations and density of individuals.

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fully introduced in the following chapter. chapter facilitates understanding of spatial capture-recapture models, which are tial capture-recapture. Providing background on capture-recapture in general, the last section of this chapter highlights the parallels of distance sampling and spacapture location, are a step towards fully spatial capture-recapture models. The the covariate is some description of an animal's location in space, such as average and discuss their shortcomings. We show that individual covariate models, where area in order to estimate density. We present some common approaches to do so not linked to a specific area and so ad hoc approaches are necessary to define an variation in p. The drawback of non-spatial estimates of abundance is that they are code for non-spatial capture-recapture models incorporating different sources of of the book. To demonstrate data augmentation, we present examples and BUGS ual covariates, and thus we use data augmentation frequently throughout the rest capture-recapture model facilitates Bayesian analysis and the inclusion of individare part of the population but were never observed. This reformulation of the encounter histories, and estimating how many of these hypothetical individuals involves adding a large number of all-zero encounter histories to the n observed we are interested in estimating N, and we can do so using data augmentation, which on each occasion) are Bernoulli trials with detection probability p. Usually, though, ple logistic regressions, where the observations (0 = not captured or 1 = captured)Bayesian framework. If population size (M) is known, these models resemble simcapture-recapture models for estimating abundance, emphasizing analysis in a Abstract: In this chapter we introduce traditional (non-spatial) closed-population

Keywords: Behavioral response, Data augmentation, Density estimation, Detection probability, Distance sampling, Effective area sampled, Heterogeneity model, Individual covariate model, MMDM, Model Mh, Non-spatial capture-recapture

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