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unmarked:

An R package for the Analysis of Wildlife Occurrence and Abundance Data

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Abstract

Ecological research uses data collection techniques that are prone to substantial and unique types of measurement error to address scientific questions about species abundance and distribution. These data collection schemes include a number of survey methods in which unmarked individuals are counted, or determined to be present, at spatially-referenced sites. Examples include site occupancy sampling, repeated counts, distance sampling, removal sampling, and double observer sampling. To appropriately analyze these data, hierarchical models have been developed to separately model explanatory variables of both a latent abundance or occurrence process and a conditional detection process. Because these models have a straightforward interpretation paralleling mechanisms under which the data arose, they have recently gained immense popularity. The common hierarchical structure of these models is well-suited for a unified modeling interface. The R package unmarked provides such a unified modeling framework, including tools for data exploration, model fitting, model criticism, post-hoc analysis, and model comparison.

Keywords: ecological, wildlife, hierarchical, occupancy, occurrence, distance, point count.

1. Introduction

1.1. Imperfect detection and data collection in ecological research

A fundamental goal of ecological research is to understand how environmental variables influence spatial or temporal variation in species abundance or occurrence. Addressing these research questions is complicated by imperfect detection of individuals or species. Individuals may go undetected when present for a variety of reasons including their proximity to the observer, cryptic behavior, or camouflage. Thus, imperfect detection can introduce substantial measurement error and obscure underlying ecological relationships if ignored.

To accommodate imperfect detection of individuals or species, ecologists have developed specialized methods to survey wildlife populations such as site occupancy sampling, repeated counts, distance sampling, removal sampling, and double observer sampling (see Section 2 and (Williams et al. 2002) for definitions). This myriad of sampling methods are all unified by a common repeated-measures type of sampling design in which J observations are made at M spatial sample units during each of T seasons. As an example, Table 1.1 shows a typical repeated count dataset.

		Season 1			Season 2			
	Visit 1	Visit 2	Visit 3	Visit 1	Visit 2	Visit 3		
Site 1	3	3	2	0	0	0		
Site 2	0	0	0	0	0	0		
Site 3	4	6	5	5	5	3		
Site 4	0	0	0	0	0	2		

Table 1: An example of the data structure required by **unmarked**'s fitting functions. Counts of organisms were made at M=4 sites during T=2 seasons with J=3 visits per season.

In addition to the repeated-measures design, there are two important features to note about the data shown in Table 1.1. First, the counts are of individuals that cannot be uniquely recognized, so although it may be possible to determine if three individuals were detected on two different visits (as was the case at Site 1 in Table 1.1), it is not possible to determine if they were the *same* three individuals. Second, this design is often referred to as a 'metapopulation design' because the population may be regarded as an aggregation of subpopulations and because the spatial sampling is an explicit component of the problem. This is important because it provides a basis for modeling variation among sites as a function of site-specific covariates, which is often of central interest, and because metapopulation processes such as local colonization and extinction can be directly estimated.

Both the metapopulation design and absence of individual recognition distinguish this study design from another large class of ecological sampling methods known as 'capture-recapture' methods. A vast number of models have been developed for capture-recapture data, and many of them are implemented in the software program MARK (White and Burnham 1999). The origins of **unmarked** (the package and its name) arose to fill the void of models for data from studies of unmarked individuals involving explicit spatial sampling.

1.2. Hierarchical Models for Metapopulation Designs

While many ecological studies produce data consistent with a metapopulation design, the development and implementation of models for inference under individual sampling protocols has proceeded in a piecemeal fashion, without a coherent analysis platform. Recently, however, a broad class of hierarchical models (see (Royle and Dorazio 2008) for a general treatment) has been developed that offers a unified framework for analysis by formally recognizing that observations are generated by a combination of (1) a *state* process determining abundance or species occurrence at each site and (2) a *detection* process that yields observations conditional on the state process. The model for the state process describes abundance or occurrence at

each site, but due to imperfect detection, these quantities are regarded actual abundance or occurrence as a latent variable. These models greatly improve upon traditional models that focused solely on modeling the observation process and used *ad hoc* methods of 'correcting' observed data for detection probability.

This paper introduces **unmarked**, an R package that provides a unified approach for fitting this broad class of hierarchical models developed for sampling biological populations. Inference in **unmarked** is based on the integrated likelihood wherein the latent state variable is marginalized out of the conditional likelihood.

1.3. Scope and features of unmarked

unmarked provides tools to assist researchers with every step of the analysis process, including data manipulation and exploration, model fitting, post-hoc analysis, model criticism, and model selection. unmarked provides a growing list of model-fitting functions designed for specific sampling methods. The fitting functions each find the maximum likelihood estimates of parameters from a particular model (Section 3.3) and return an object that can be easily manipulated. Methods exist for performing numerous post-hoc analyses such as requesting linear combinations of parameters, back-transforming parameters to constrained scales, determining confidence intervals, and evaluating goodness of fit. The model specification syntax of the fitting functions was designed to resemble the syntax of R's common fitting functions such as 1m for fitting linear models.

Although there is existing software for fitting some of these models (e.g. (Hines and MacKenzie 2002)), there are a number of advantages to a unified framework within R. Many researchers are already familiar with R and use its powerful data manipulation and plotting capabilities. Sometimes many species are analyzed in tandem, so that a common method of aggregating and post-processing of results is needed, a task easily accomplished in R. Unlike other available software, **unmarked** makes it easy to map habitat-specific abundance and species distributions when combined with R's GIS capabilities. Another important advantage of **unmarked**'s approach is that researchers can easily simulate and analyze data within the same computational environment. This work flow permits simulation studies for power analysis calculations or the effectiveness of future sampling designs. All of this is made much simpler by analyzing the data within R, and using a single environment to complete all phases of the analysis is much less error-prone than switching between applications.

In this paper, Section 2 gives a brief summary of many of the models **unmarked** is capable of fitting. Section 3 describes general **unmarked** usage aided by a running data example.

2. Models implemented in unmarked

The list of models implemented in **unmarked** continues to grow as new models are developed. Table 2 shows the models available as of version 0.9-0. Rather than describe each fitting function in detail, this section provides a summary of several of the most common sampling techniques and how **unmarked** can be used to model the resulting data.

Multi-season Abundance	Multi-season Occupancy	Repeated Multinomial Counts	Multinomial Counts	Distance-sampling	Abundance from repeated counts	Abundance from presence-absence	Single-season Occupancy	Model
pcountOpen	colext	gmultmix	multinomPois	distsamp	pcount	occuRN	occu	Fitting function
unmarkedPCO	unmarkedMultFrame	${ m unmarkedFrameGMM}$	${\bf unmarked Frame MPois}$	${\rm unmarkedFrameDS}$	unmarked Frame PC ount	${ m unmarkedFrameOccu}$	${\tt unmarkedFrameOccu}$	Data
(Dail and Madsen 2011)	(MacKenzie et al. 2003)	(Chandler et al. In press)	(Royle 2004a)	(Royle $et \ al. \ 2004$)	(Royle 2004b)	(Royle and Nichols 2003)	(MacKenzie et al. 2002)	Citation

Table 2: Models currently handled by unmarked along with their associated fitting functions (Section 2) and data type (Section 3.1).

2.1. Site occupancy models

Single season site occupancy model

An important state variable in ecological research is species occurrence (or 'site occupancy'), say Z_i , a binary state variable such that $Z_i = 1$ if site i is occupied by a species and $Z_i = 0$ otherwise. Much interest is focused on estimating functions of species occurrence (e.g., number of occupied sites) or to identify factors that are associated with changes in the probability of a site being occupied, i.e., $\psi_i = \Pr(Z_i = 1)$. To estimate these parameters, researchers employ a sampling design, whereby surveyors visit a sample of M sites and record the binary response Y_{ij} of species detection (Y = 1) or non-detection (Y = 0) during $j = 1, \ldots, J_i$ visits to the ith site during a 'season' (MacKenzie et al. 2002). A key feature of species occurrence surveys is that false absences are possible, so that a species might go undetected $(Y_{ij} = 0)$ even if it is present $(Z_i = 1)$. A set of parameters p_{ij} accounts for this observation process, where p_{ij} is the probability of detecting the species at site i during the jth survey occasion given that site i is truly occupied. That is, $p_{ij} = \Pr(Y_{ij} = 1 | Z_i = 1)$.

The key assumptions made when modeling these data are that the occupancy state at a site remains constant throughout the season and repeated visits at a site are independent. The first assumption is commonly referred to as the 'population closure' assumption, meaning that no births, deaths, emigrations, or immigrations occur during the season. Season, therefore, will generally refer to a very short time frame, such as a few months during a breeding season, or even a few minutes if repeated visits are made in quick succession. Replicate samples (i.e., J > 1) are necessary to obtain information about the detection rate separate from the occupancy rate.

The following hierarchical model describes the joint distribution of the observations conditional on the latent occupancy state, and the marginal distribution of the latent occupancy state variable:

$$Z_i \sim \text{Bernoulli}(\psi_i) \quad \text{for } i = 1, 2, \dots, M$$

 $Y_{ij}|Z_i \sim \text{Bernoulli}(Z_i p_{ij}) \quad \text{for } j = 1, 2, \dots, J_i$

Removing the latent Z variables by marginalization yields the likelihood:

$$L(\psi, p|Y_{ij}) = \prod_{i}^{M} \left\{ \prod_{j}^{J} \left(p^{Y_{ij}} (1-p)^{1-Y_{ij}} \right) \psi + I(Y_{i.} = 0)(1-\psi) \right\}, \tag{1}$$

where I is the indicator function taking the value 1 if its argument is true, and 0 otherwise. The model and likelihood are easily generalizable to accommodate covariates on detection probability or occupancy. Variables that are related to the occupancy state are modeled as

$$logit(\psi_i) = \mathbf{x}_i' \beta$$
,

where \mathbf{x}_i is a vector of site-level covariates and β is a vector of their corresponding effect parameters. Similarly, the probability of detection can be modeled with

$$logit(p_{ij}) = \mathbf{v}'_{ij}\alpha,$$

where \mathbf{v}_{ij} is a vector of observation-level covariates and α is a vector of their corresponding effect parameters. Examples of site-level covariates include habitat characteristics such as vegetation height. Observation-level covariates could include time of day, date, wind speed, or other factors that might affect detection probability. The function occu implements this model.

Multi-season site occupancy model

Sometimes the study objective is to understand the dynamics of the occupancy state over time. To obtain such information researchers conduct repeated occupancy studies (see Section 2.1.1) at the same sample of sites over consecutive seasons (MacKenzie et al. 2003) and seek to estimate probabilities of colonization (γ_{it}) and extinction (ϵ_{it}), where colonization is the change of an unoccupied site to occupied and extinction if the change of an occupied site to unoccupied. If the occupancy status is assumed to evolve according to a Markov process, then a 2-state finite hidden Markov model describes these data. Let Y_{itj} denote the observed species occurrence status at visit j during season t to site i. Then

$$Z_{i1} \sim \text{Bernoulli}(\psi_i)$$

$$Z_{it} \sim \begin{cases} \text{Bernoulli}(\gamma_{i(t-1)}) & \text{if } Z_{i(t-1)} = 0 \\ \text{Bernoulli}(1 - \epsilon_{i(t-1)}) & \text{if } Z_{i(t-1)} = 1 \end{cases},$$

$$\text{for } t = 2, 3, \dots, T$$

$$Y_{itj}|Z_{it} \sim \text{Bernoulli}(Z_{it}p_{itj})$$

This model generalizes the single season site occupancy model by relaxing the population closure assumption. To define the likelihood of this model, let $\phi_0 = (\psi, 1 - \psi)$ and

$$\theta_{\mathbf{Y_{it}}} = \begin{pmatrix} \prod_{j}^{J} pI(Y_{ijt} = 0)(1 - p_{ijt}) \\ I(\sum_{j}^{J} Y_{ijt} = 0). \end{pmatrix}$$

Furthermore, let the matrix of one-step Markov transition probabilities be

$$\Phi_t = \begin{pmatrix} 1 - \epsilon & \epsilon \\ 1 - \gamma & \gamma \end{pmatrix}$$

The likelihood is then

$$L(\psi, \epsilon, \gamma, p | Y_{ijt}) = \prod_{i}^{R} \left\{ \phi_0 \left\{ \prod_{t}^{T-1} \Phi_t \theta_{Y_{it}} \right\} \theta_{Y_{iT}} \right\}, \tag{2}$$

which can be maximized by the **colext** function in **unmarked**. Again, each of the four model parameters ψ , γ , ϵ , and p can be modeled as functions of covariates on the logit scale.

In addition to the two site-occupancy models discussed above, **unmarked** contains the function **occuRN** that to fit the model of Royle and Nichols (2003) that estimates abundance from detection, non-detection data.

2.2. Abundance models

Repeated count data

Occupancy is a crude summary of population structure and dynamics and, in practice, considerable effort is focused on obtaining information about abundance. A common sampling design for obtaining abundance information, analogous to the basic occupancy design described above, is to repeatedly visit a sample of M sites J times and record the number of unique individuals observed at each site. Similar assumptions are made as with occurrence data: (1) abundance at a site remains constant during a season and (2) counts at a site are independent. Royle (2004b) presented the following hierarchical model for repeated count data. Let N_i be the unobserved total number of individuals using a site and define Y_{ij} as the number of individuals observed during the jth visit. Then,

$$N_i \sim f(\lambda_i, \theta)$$
 for $i = 1, 2, \dots, M$
 $Y_{ij} | N_i \sim \text{Binomial}(N_i, p_{ij})$ for $j = 1, 2, \dots, J_i$,

where λ_i is the abundance rate at site i and p_{ij} is the detection probability during the jth visit to site i. f is a discrete distribution with support restricted to $N_i \geq 0$ and θ are extra parameters of f other than the location parameter, λ_i . unmarked currently supports f as Poisson or negative binomial. In the Poisson case, there is no θ . In the negative binomial case, θ is a dispersion parameter, which is useful when overdispersion is suspected.

As with the occupancy model, covariates may be included at either the state (here, abundance) or detection levels, but abundance is modeled through a log link to enforce it's positivity constraint.

$$\log(\lambda_i) = \mathbf{x}_i' \beta,$$

where \mathbf{x}_i is a vector of site-level covariates and β is a vector of their corresponding effect parameters. Similarly, the probability of detection can be modeled with

$$logit(p_{ij}) = \mathbf{v}'_{ij}\alpha,$$

where \mathbf{v}_{ij} is a vector of observation-level covariates and α is a vector of their corresponding effect parameters.

The N_i variables are latent and so analysis is easily carried out based on the integrated likelihood obtained by marginalizing each N_i from the conditional likelihood Royle *et al.* (2004):

$$L(Y_{ij}|p,\lambda) = \prod_{i}^{M} \left\{ \sum_{N_{i}=max(\mathbf{Y_{i}})}^{\infty} \left(\prod_{j}^{J} \frac{N_{i!}}{(N_{i}-Y_{ij})!} p^{Y_{ij}} (1-p)^{N_{i}-Y_{ij}} \right) \frac{e^{-\lambda} \lambda^{N}}{N!} \right\}.$$
(3)

This likelihood can be maximized using the pcount function.

As the single-season occupancy model was generalized for multi-season data, the counterpart of this model was proposed by that relaxes the population closure assumption and may be fit using the pcountOpen.

General multinomial-Poisson mixture model

Here we discuss a more general class of models that can be customized by the user for a variety of sampling methods, the multinomial-Poisson model (Royle 2004a). The general form of this model is

$$N_i \sim \text{Poisson}(\lambda_i)$$
 for $i = 1, 2, ..., M$

$$\begin{pmatrix} \mathbf{Y}_i \\ N_i - \sum_{j=1}^J Y_{ij} \end{pmatrix} \middle| N_i \sim \text{Multinomial} \left(N_i, \begin{pmatrix} \boldsymbol{\pi}_i \\ \boldsymbol{\pi}_i^* \end{pmatrix} \right)$$

where N_i is the latent abundance at site i as with the repeated count model, and $\pi_i = (\pi_{i1}, \pi_{i2}, \dots, \pi_{iJ})'$ is the vector of cell probabilities of observing responses in the J possible categories, and \mathbf{Y}_i is the vector of counts that were actually observed. In general, π_i is determined by the specific sampling method and $\sum_j \pi_{ij} \leq 1$ because detection is imperfect and $\pi_i^* = 1 - \sum_j \pi_{ij}$ is the probability of the species escaping detection at site i.

It is worth noting here that the replication here is of a different nature — not over time necessarily — but still effectively replication as far as the design goes. That is you have a 'sites by K' matrix of counts its just that, with a multinomial protocol., the replicate counts are DEPENDENT instead of independent - and this is modeled with the multinomial observation model. Not sure where to say this blurb but probably should be in the first paragraph of this section.

To illustrate the likelihood under a multinomial observation model, suppose that a sampling method produces a multinomial observation with 3 observable frequencies at each site. As before, N_i are latent variables, and so inference is based on the integrated likelihood of \mathbf{y}_i which is:

$$L(\mathbf{Y}_{i}|\mathbf{p},\lambda) = \prod_{i=1}^{N} \left\{ \sum_{M_{i}=0}^{\infty} \left(\frac{N_{i}!}{y_{i1}! y_{i2}! y_{i3}! y_{i0}!} \pi_{1}^{y_{i1}} \pi_{2}^{y_{i2}} \pi_{3}^{y_{i3}} \pi^{*N_{i}-y_{i.}} \right) \frac{e^{-\lambda} \lambda^{N}}{N!} \right\}$$
(4)

For the specific case of a multinomial/Poisson mixture, the likelihood simplifies analytically (to a product-Poisson likelihood). The function multinomPois can be used to maximize this likelihood.

For multinomial-Poisson sampling methods, the actual observations are an underlying categorical detection variable with $M \leq J$ levels so that the J-dimensional \mathbf{Y}_i is derived from the M-dimensional raw counts in some sampling method-specific manner. Thus, it is necessary to model the detection at the raw observation level, denoted p_{ik} for k = 1, 2, ..., M at site i. Then we derive the multinomial cell probabilities π_i through the sampling technique-specific relationship $\pi_{ij} = g(p_{ik})$ where p_{ik} is the underlying probability of detection and g is some sampling method-specific function.

Thus, the only two requirements to adapt **unmarked**'s general multinomial Poisson modeling to a new sampling method is to specify g and a binary 0-1 matrix that describes the mapping of elements of $\mathbf{p}_i = (p_{i1}, \dots, p_{iR})'$ to elements of π_i . This mapping matrix, referred to in **unmarked** as obsToY, is necessary to consistently clean missing values from the data and relate observation-level covariates with the responses. The (j,k)th element of obsToY is 1 only if p_{ik} is involved in the computation of π_{ij} . The detection function g is called piFun in **unmarked**.

Covariates may be included in either the state (here, abundance) or detection models, through \mathbf{p}_i (not π_i).

$$\log(\lambda_i) = \mathbf{x}_i' \beta,$$

where \mathbf{x}_i is a vector of site-level covariates and β is a vector of their corresponding effect parameters. Similarly, the probability of detection can be modeled with

$$logit(p_{ij}) = \mathbf{v}'_{ij}\alpha,$$

where \mathbf{v}_{ij} is a vector of observation-level covariates and α is a vector of their corresponding effect parameters.

We now describe two common sampling methods that can be modeled with the multinomial-Poisson model: removal sampling and double observer sampling. These two methods are included in **unmarked**, but additional methods may easily be specified by defining a user-specified piFun function and obsToY matrix.

Removal sampling Popular in fisheries, removal sampling is commonly implemented by visiting a sample of M sites J times each and trapping or otherwise capturing and then removing individuals at each visit. Thus, Y_{ij} is the number of individuals captured at the jth visit for j = 1, 2, ..., J. We can specify g for removal sampling as follows. The probability of an individual at site i being captured on the first visit is $\pi_{i1} = p_{i1}$. The probability of capture on the jth visit is

$$\pi_{ij} = \prod_{k=1}^{j-1} (1 - p_{ik}) p_{ij},$$

for j = 2, ..., J and the probability of not being sampled is

$$\pi_{i,J+1} = \prod_{j=1}^{J} (1 - p_{ij})$$

Or, equivalently,

$$\pi_i = \begin{pmatrix} p_{i1} \\ (1 - p_{i1})p_{i2} \\ \vdots \\ \prod_{j=1}^J (1 - p_{ij})p_{iJ} \end{pmatrix}$$

Thus, the mapping matrix is an $J \times J$ matrix with ones in the upper triangle,

$$\begin{pmatrix} 1 & 1 & \dots & 1 \\ 0 & 1 & \dots & 1 \\ \vdots & & \ddots & \vdots \\ 0 & 0 & \dots & 1 \end{pmatrix}$$

Double observer sampling Double observer sampling involves collecting data by a team of two surveyors simultaneously visiting a site. Each observer records a list of detected animals and at the end of the survey, the two observers attempt to reconcile their counts. If individuals are not uniquely marked, this may be a difficult task in practice; however, assuming that individuals can be distinguised, the data at each site are a vector of length three \mathbf{Y}_i , corresponding to the numbers of individuals seen by only observer one, only observer two, and both observers. Thus, for double observer sampling, g is defined as follows.

$$\pi_i = \begin{pmatrix} p_{i1}(1 - p_{i2}) \\ (1 - p_{i1})p_{i2} \\ p_{i1}p_{i2} \\ (1 - p_{i1})(1 - p_{i2}). \end{pmatrix}$$

The obsToY mapping matrix for double observer sampling is the following 3×2 matrix.

$$\begin{pmatrix} 1 & 0 \\ 0 & 1 \\ 1 & 1 \end{pmatrix}$$

3. unmarked usage

unmarked provides data structures, fitting syntax, and post-processing that form a cohesive framework for site-based ecological data analysis. In order to achieve these goals, unmarked uses the S4 class system (Chambers 2008). As R's most modern system of class-based programming, S4 allows customization of functions, referred to as methods, to specific object classes and superclasses. For example, when the generic predict method is called with any unmarked model fit object as an argument, the actual predict implementation depends on the specific model that was fit. Use of class-based programming can provide more reliable and maintainable software while also making the program more user-friendly (Chambers 2008).

3.1. Preparing data

unmarked uses a custom S4 data structure called the unmarkedFrame to store all data and metadata related to a sampling survey. Although this at first appears to add an extra layer of work for the user, there are several reasons for this design choice. The multilevel structure of the models means that standard rectangular data structures such as data.frames or matrices are not suitable for storing the data. For example, covariates might have been measured separately at the site level and at the visit level. Furthermore, the length of the response vector \mathbf{Y}_i at site i might differ from the number of observations at the site as in the multinomial Poisson model. In some cases, metadata of arbitrary dimensions may need to be associated with the data. For example, in distance sampling it is necessary to store the units of measure and the survey design type. Aside from these technical reasons, Gentleman (2009) pointed out that the use of such portable custom data objects can simplify future reference to previous analyses, an often neglected aspect of research. Repeated fitting calls using the same set of data require less code repetition if all data are contained in a single object. Finally, calls to fitting functions have a cleaner appearance with a more obvious purpose when the call is not buried in data arguments. The parent data class is called an unmarkedFrame and each

unmarked fitting function has its own data type that extends the unmarkedFrame. Thus, the first step when using unmarked is to import data into the proper type of unmarkedFrame. To ease this step, unmarked includes several helper functions to automatically convert data into an unmarkedFrame: csvToUMF which imports data directly from a comma-separated value text file, formatWide and formatLong which convert data from data frames, and the family of unmarkedFrame constructor functions.

An unmarkedFrame object contains components referred to as slots, which hold the data and metadata. All unmarkedFrame objects contain a slot for the observation matrix y, a data.frame of site-level covariates siteCovs, and a data.frame of observation-level covariates obsCovs. The y matrix is the only required unmarkedFrame slot. Each row of y contains either the observed counts or presence-absence data at each of the R sites. siteCovs is an M-row data.frame with a column for each site-level covariate. obsCovs is an M-row data.frame with a column for each observation-level covariate. Thus each row of obsCovs corresponds to a particular observation, with the order corresponding to site varying slower and observation within site varying faster. Both siteCovs and obsCovs can contain NA values corresponding to unbalanced or missing data. If any terms in the model specification are missing for that observation, unmarked automatically removes observations. unmarked provides constructor functions to make creating unmarkedFrames straightforward. For each specific data type, specific types of unmarkedFrames extend the basic unmarkedFrame to handle model-specific nuances.

Importing repeated count data

Here is an example of creating an unmarkedFrame for repeated count data (Section 2.2.1). First, load the included data set of Mallard (*Anas platyrhynchos*) point counts described in Kéry *et al.* (2005).

> library(unmarked)
> data(mallard)

Loading the mallard data makes three objects available within the R workspace. The matrix mallard occurrence of mallards counted at each of M=239 sites on J=3 visits. It is formatted such that each row is a site and each column corresponds to a visit. The site-level covariates are column vectors in the mallard site data frame, which has M rows. The observation-level covariates are a list object with separate $M \times J$ matrices for each observation-level covariate. The unmarked Frame constructors can accept obsCovs in this list format or as a data frame in the format described above.

The following call to unmarkedFramePCount organizes the observations and covariates into an object that can be passed to the data argument of the fitting function pcount.

```
> mallardUMF <- unmarkedFramePCount(y = mallard.y, siteCovs = mallard.site,
+ obsCovs = mallard.obs)</pre>
```

Printing unmarkedFrames shows them as data.frames to make it easier to verify that the data were converted correctly. Here only the first five sites (rows) and the first two visits are shown.

> mallardUMF[1:5, 1:2]

Data frame representation of unmarkedFrame object.

```
y.1 y.2 elev length forest ivel.1 ivel.2 date.1 date.2
1 0 0 -1.173 0.801 -1.156 -0.506 -0.506 -1.761 0.310
2 0 0 -1.127 0.115 -0.501 -0.934 -0.991 -2.904 -1.047
3 3 2 -0.198 -0.479 -0.101 -1.136 -1.339 -1.690 -0.476
4 0 0 -0.105 0.315 0.008 -0.819 -0.927 -2.190 -0.690
5 3 0 -1.034 -1.102 -1.193 0.638 0.880 -1.833 0.167
```

The first three columns contain the count data. The site-level covariates are elevation (elev), transect length (length), and the proportion of forest covering the site (forest). The two observation-level covariates are a measure of survey effort (ivel) and the date of the survey (date). Note that these variables are standardized to mean of 0 and unit variance. We can also check data contents with a quick summary.

> summary(mallardUMF)

unmarkedFrame Object

239 sites

Maximum number of observations per site: 3 Mean number of observations per site: 2.76 Sites with at least one detection: 40

Tabulation of y observations:

```
0
        1
              2
                    3
                               7
                                    10
                                          12 <NA>
576
                   9
                         6
                                           1
       54
            11
                               1
                                    1
                                               58
```

Site-level covariates:

elev	length	forest		
Min. :-1.436000	Min. $:-4.95e+00$	Min. :-1.265000		
1st Qu.:-0.956500	1st Qu.:-5.63e-01	1st Qu.:-0.956000		
Median :-0.198000	Median : 4.50e-02	Median :-0.065000		
Mean :-0.000046	Mean :-2.93e-05	Mean : 0.000067		
3rd Qu.: 0.994000	3rd Qu.: 6.26e-01	3rd Qu.: 0.790000		
Max. : 2.434000	Max. : 2.25e+00	Max. : 2.299000		

Observation-level covariates:

ivel			date			
Min.	:-	-1.75e+00	Min.	:-	-2.90e+00	
1st Qu.	:-	-6.66e-01	1st Qu.	:-	-1.12e+00	
Median	:-	-1.39e-01	Median	:-	-1.19e-01	
Mean	:	1.50e-05	Mean	:	7.26e-05	
3rd Qu.	:	5.49e-01	3rd Qu.	:	1.31e+00	
Max.	:	5.98e+00	Max.	:	3.81e+00	
NA's	:	5.20e+01	NA's	:	4.20e+01	

The summary reveals that only 40 sites have at least one detection. The tabulation of y observations provides additional evidence of sparse counts, with no mallards being detected during 576 of the surveys. The 58 NA values correspond to missing data.

Importing removal sampling data

To illustrate the slightly different syntax for removal sampling data example, we will import data from a removal survey of ovenbirds (Seiurus aurocapillus) described by Royle (2004a). The data consist of a list named ovendata.list with a matrix data containing the removal counts for 4 visits and a data.frame called covariates containing site-level covariates information.

> data(ovendata)

THIS LOADS XYZ... Thus, it is evident that ovenbirds were detected at three of the first ten sites and no new birds were detected after the first removal occasion. The only additional specification required when importing removal data is to specify the particular type of multinomial-Poisson data as removal via the type argument.

```
> ovenFrame <- unmarkedFrameMPois(ovendata.list$data, siteCovs = ovendata.list$covariates,
+ type = "removal")
> head(ovenFrame, 5)
```

Data frame representation of unmarkedFrame object.

```
y.1 y.2 y.3 y.4
                    site
                           ufp trba
            0
                0 CAT003 23.43 62.5
        0
2
    1
        0
            0
                0 CAT004 15.62 70.0
                0 CAT011 35.54 90.0
4
                0 CAT013 25.78 60.0
    0
        0
                0 CAT017 25.00 122.5
```

Preparing multi-season occupancy data

The data structures are more complex when surveys occurred during more than one season. In this case, the covariates can occur at the site-, season-, and observation-levels. The following toy examples demonstrates how to prepare data for the colext fitting function if there were only four sites, two visits per season, and three seasons.

```
> M <- 4
> J <- 2
> T <- 3
> y <- matrix(0:1, nrow = M, ncol = J * T)
> rownames(y) <- paste("site", 1:M, sep = "")
> cn <- paste("season", rep(1:2, each = 3), sep = "")
> cn <- paste(cn, paste("v", rep(1:3, 2), sep = ""), sep = "")
> colnames(y) <- cn
> y
```

	season1v1	season1v2	season1v3	season2v1	season2v2	season2v3
site1	0	0	0	0	0	0
site2	1	1	1	1	1	1
site3	0	0	0	0	0	0
site4	1	1	1	1	1	1

```
> umf <- unmarkedMultFrame(y = y, numPrimary = T)</pre>
```

The function unmarkedMultFrame accepts siteCovs and obsCovs like all other unmarked-Frame classes, and it also has an argument yearlySiteCovs that can accept a list of $M \times T$ data.frames containing season-level covariates. The numPrimary argument specifies the number of seasons.

3.2. Manipulating unmarkedFrames

The various components of unmarkedFrames can be extracted and subsetted in a manner similar to the methods used to manipulate standard R objects. Subsetting can be accomplished using the 'bracket' notation. For example, the first five rows of data and all three removal occasions can be extracted from the ovenbird removal study using

```
> ovenFrame[1:5, 1:3]
```

Data frame representation of unmarkedFrame object.

```
y.1 y.2 y.3
                 site
                        ufp
                             trba
1
    0
        0
            0 CAT003 23.43
                             62.5
            0 CAT004 15.62
2
3
            0 CAT011 35.54
                             90.0
        0
            0 CAT013 25.78 60.0
            O CAT017 25.00 122.5
5
    0
        0
```

In some cases, the covariate data may need to be manipulated after the unmarkedFrame has been created. The following code demonstrates how to extract the site-level covariates, standardize those that are continuous (columns 2 and 3), and reinsert them back into the unmarkedFrame.

```
> sc <- siteCovs(ovenFrame)
> sc[, 2:3] <- scale(sc[, 2:3])
> siteCovs(ovenFrame) <- sc</pre>
```

3.3. Fitting models

As introduced in Section 2, each type of survey design has a corresponding fitting function. For example, to fit a repeated count model, we call pcount. Table 2 provides a summary of all models that unmarked currently fits. With the exception of the 'open' population models (colext, gmultmix, and pcountOpen), all fitting functions use a double right-hand sided formula syntax representing the hierarchical model structure. Specifically, covariates

affecting the detection process are specified following the first tilde, and covariates of the state process follow the second tilde. No left-hand side of the formula is specified because the unmarkedFrame defines the response variable uniquely as the y slot.

Fitting a repeated count model

Continuing the Mallard example, the following call to pcount fits a binomaial-Poisson mixture model (Section 2.2.1). The following code specifies that detection probability p should be modeled by day of year, including a quadratic term. We also wish to model abundance using elevation and proportion of area forested. As described in Section 2.2.1, covariates of detection are on the logit-scale and covariates of abundance are on the log scale for the repeated count model.

```
> fm.mallard.1 <- pcount(~date + I(date^2) ~ elev + forest,
      data = mallardUMF, K = 50)
> fm.mallard.1
Call:
pcount(formula = "date + I(date^2) " elev + forest, data = mallardUMF,
    K = 50
Abundance:
                                z P(>|z|)
            Estimate
                        SF.
              -1.853 0.236 -7.87 3.62e-15
(Intercept)
elev
              -1.232 0.234 -5.27 1.36e-07
forest
              -0.756 0.165 -4.60 4.32e-06
Detection:
                         SE
            Estimate
                                   z P(>|z|)
              0.3071 0.2149 1.4290 0.15301
(Intercept)
             -0.4091 0.1482 -2.7612 0.00576
date
             -0.0077 0.0853 -0.0903 0.92801
I(date^2)
AIC: 521.2
```

This initial fit suggests that Mallard abundance decreases with increasing elevation and forest. It also looks like a linear model might suffice for the detection model, so we subsequently fit the linear detection model as follows:

```
> fm.mallard.2 <- pcount(~date ~ elev + forest, data = mallardUMF,
+ K = 50)
> fm.mallard.2

Call:
pcount(formula = ~date ~ elev + forest, data = mallardUMF, K = 50)
```

Abundance:

```
Estimate SE z P(>|z|)
(Intercept) -1.857 0.232 -7.99 1.37e-15
elev -1.236 0.230 -5.38 7.59e-08
forest -0.756 0.164 -4.60 4.23e-06
```

Detection:

```
Estimate SE z P(>|z|)
(Intercept) 0.298 0.188 1.58 0.113983
date -0.401 0.114 -3.51 0.000449
```

AIC: 519.2

This seems to be a better model according to both the Wald p-value and AIC. The result suggests that detection probability decreases during the course of a year.

Fitting a multinomial-Poisson model

Here we demonstrate fitting a multinomial-Poisson mixture model to removal sampling data. The Ovenbird data has no observation-level covariates, so detection probability is assumed constant across visits. It is not necessary to specify that removal sampling was used when fitting the model because this information is already stored in the ovenFrame data. We model abundance as a function of understory forest coverage (ufp) and average basal tree area (trba).

```
> fm.oven.1 <- multinomPois(~1 ~ ufp + trba, ovenFrame)
> fm.oven.1
```

Call:

```
multinomPois(formula = ~1 ~ ufp + trba, data = ovenFrame)
```

Abundance:

	Estimate	SE	z	P(> z)
(Intercept)	0.102	0.119	0.864	0.388
ufp	0.100	0.126	0.794	0.427
trba	-0.171	0.135	-1.262	0.207

Detection:

```
Estimate SE z P(>|z|)
0.288 0.233 1.24 0.217
```

AIC: 326.1

3.4. Examining fitted models

Objects returned by unmarked's fitting functions also make use of the S4 class system. All fitted model objects belong to the unmarkedFit parent class. Thus, common operations such

as extracting coefficient estimates, covariance matrices for estimates, and confidence intervals have been adapted to behave similar to R's base functions.

For example, we can extract estimated coefficients either from the entire model, or from the state or detection levels by specifying the type argument.

```
> coef(fm.mallard.2)
```

```
lam(Int) lam(elev) lam(forest) p(Int) p(date)
-1.8565 -1.2355 -0.7555 0.2978 -0.4006
> coef(fm.mallard.2, type = "state")
lam(Int) lam(elev) lam(forest)
-1.8565 -1.2355 -0.7555
```

To check which types are available for a model, use the names method.

```
> names(fm.mallard.2)
[1] "state" "det"
```

Similarly, the vcov function extracts the covariance matrix of the estimates, using the observed Fisher information by default.

vcov also accepts a type argument, as does the convenience method SE, which returns standard errors from the square root of the diagonal of the covariance matrix.

Nonparametric bootstrapping can also be used to estimate the covariance matrix. unmarked implements a two-stage bootstrap in which the sites are first drawn with replacement, and then within each site, the observations are drawn with replacement. First, bootstrap draws must be taken using the nonparboot. nonparboot returns a new version of the unmarkedFit object with additional bootstrap sampling information. Thus, this new fit must be stored, either in a new fit object or the same one, and then subsequently queried for bootstrap summaries. In the following, bootstrapping can be quite slow. For these examples, we illustrate with the removal sampling data simply because computations are much faster. However, bootstrapping is available for any of the models that unmarked fits.

```
> set.seed(1234)
> fm.oven.1 <- nonparboot(fm.oven.1, B = 10)
> SE(fm.oven.1, type = "state")
```

```
lambda(Int) lambda(ufp) lambda(trba)
     0.1185     0.1263     0.1353

> SE(fm.oven.1, type = "state", method = "nonparboot")
lambda(Int) lambda(ufp) lambda(trba)
     0.1640     0.1463     0.1288
```

It looks like bootstrapping and asymptotic standard errors are close. The summary now states the number of bootstrap samples.

```
> summary(fm.oven.1)
Call:
multinomPois(formula = ~1 ~ ufp + trba, data = ovenFrame)
Abundance (log-scale):
                       SE
                                z P(>|z|)
            Estimate
(Intercept)
              0.102 0.119 0.864
                                    0.388
               0.100 0.126 0.794
ufp
                                    0.427
trba
             -0.171 0.135 -1.262
                                    0.207
Detection (logit-scale):
            SE
 Estimate
                   z P(>|z|)
    0.288 0.233 1.24
                      0.217
AIC: 326.1
Number of sites: 70
optim convergence code: 0
optim iterations: 33
Bootstrap iterations: 10
```

Additional bootstrap samples can be drawn by calling nonparboot again.

```
> fm.oven.1 <- nonparboot(fm.oven.1, B = 10)</pre>
```

Confidence intervals can be requested for the coefficients at either stage of the model. By default, the asymptotic normal approximation is used.

Profile confidence intervals are also available upon request. This can take some time, however, because for each parameter, a nested optimization within a root-finding algorithm is being used to find the profile limit.

The profile confidence intervals and normal approximations are quite similar here.

Linear combinations of estimates

Often, meaningful hypotheses can be addressed by estimating linear combinations of coefficient estimates. Linear combinations of coefficient estimates can be requested with linearComb.

Continuing the Ovenbird example, the following code estimates the log-abundance rate for a site with ufp = 0.5 and trba = 0.

```
> (lc <- linearComb(fm.oven.1, type = "state", coefficients = c(1, + 0.5, 0)))
```

Linear combination(s) of Abundance estimate(s)

```
Estimate SE (Intercept) ufp trba 0.153 0.130 1 0.5 0
```

Multiple sets of coefficients may be supplied as a matrix. The following code requests the estimated log-abundance for sites with ufp = 0.5 and trba = 1.

Linear combination(s) of Abundance estimate(s)

```
Estimate SE (Intercept) ufp trba
1 0.153 0.130 1 0.5 0
2 0.203 0.166 1 1.0 0
```

Standard errors and confidence intervals are also available for linear combinations of parameters. By requesting nonparametric bootstrapped standard errors, **unmarked** uses the samples that were drawn earlier.

```
> SE(lc, method = "nonparboot")
[1] 0.1288 0.1317
```

Back-transforming linear combinations of coefficients

Estimates of linear combination back-transformed to the native scale are likely to be more interesting than the direct linear combinations. For example, the logistic transformation is applied to estimates of detection rates, resulting in a probability bound between 0 and 1. This is accomplished with the backTransform. Standard errors of back-transformed estimates are estimated using the delta method. Confidence intervals are estimated by back-transforming the confidence interval of the original linear combination.

```
> (btlc <- backTransform(lc))</pre>
```

Backtransformed linear combination(s) of Abundance estimate(s)

```
Estimate SE LinComb (Intercept) ufp trba
1 1.16 0.151 0.153 1 0.5 0
2 1.22 0.203 0.203 1 1.0 0
```

Transformation: exp

> SE(btlc)

[1] 0.1510 0.2032

> confint(btlc)

0.025 0.975 1 0.9034 1.502

2 0.8846 1.695

3.5. Model selection

unmarked performs AIC-based model selection for structured lists of unmarkedFit objects. To demonstrate, we fit a few more models to the Ovenbird removal data, including an interaction model, two models with single predictors, and a model with no predictors.

```
> fm.oven.2 <- update(fm.oven.1, formula = ~1 ~ ufp * trba)
> fm.oven.3 <- update(fm.oven.1, formula = ~1 ~ ufp)
> fm.oven.4 <- update(fm.oven.1, formula = ~1 ~ trba)
> fm.oven.5 <- update(fm.oven.1, formula = ~1 ~ 1)</pre>
```

Now, we can organize the fitted models with the fitList function and the use the modSel method to rank the models by AIC.

```
> fmList \leftarrow fitList(\lambda) = fm.oven.1, \lambda(ufp*trba) = fm.oven.2,
      'lam(ufp)p(.)' = fm.oven.3, 'lam(trba)p(.)' = fm.oven.4,
      lam(.)p(.) = fm.oven.5)
> modSel(fmList)
                 nPars
                          AIC delta AICwt cumltvWt
lam(trba)p(.)
                     3 324.77 0.00 0.35
                                              0.35
lam(ufp)p(.)
                     3 325.73 0.96 0.21
                                              0.56
lam(ufp+trba)p(.)
                     4 326.14 1.37 0.17
                                              0.73
lam(.)p(.)
                     2 326.28
                               1.51 0.16
                                              0.90
lam(ufp*trba)p(.)
                     5 327.17
                               2.40 0.10
                                              1.00
```

It looks like the best model includes only tree basal area as a predictor of abundance. We can examine this relationship using the **predict** method (Figure 1).

predict functions much like linearComb except that new data can be passed to it as a data.frame rather than a design matrix. When the first argument given to predict is a list of models created by fitList, predict computes model-averaged predictions, which may be useful in the presence of high model selection uncertainty.

3.6. Goodness of fit

To conduct goodness of fit tests, **unmarked** provides a generic parametric bootstrapping function. It simulates data from the fitted model and applies a user-defined function that returns a fit-statistic such as the sum of squared residuals. Beyond serving as a tool to evaluate goodness of fit, **parboot** can be used to characterize uncertainty in any derived quantity of interest. Here we compute a chi-squared statistic to evaluate goodness of fit.

```
> set.seed(1234)
> chisq <- function(fm) {
+    observed <- getY(fm@data)
+    expected <- fitted(fm)
+    sum((observed - expected)^2/expected)
+ }
> pb <- parboot(fm.oven.1, statistic = chisq, nsim = 10)
> plot(pb, main = "")
```

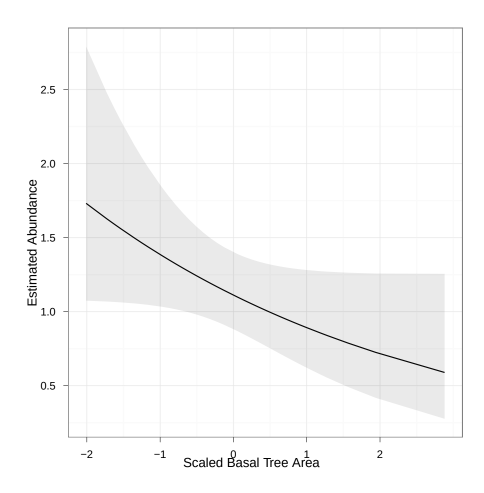
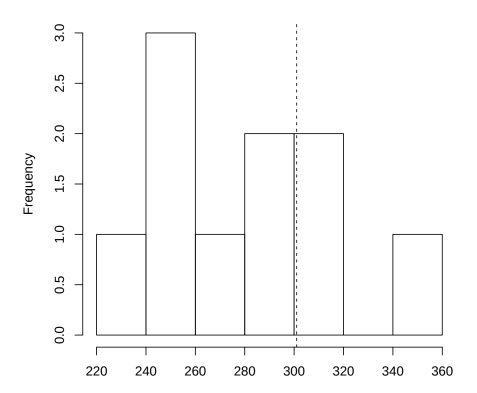


Figure 1: Examine estimated abundance for Ovenbird removal data. Band is 95% confidence interval.



The above call to plot with a parametric bootstrap object as the argument produces a useful graphic for assessing goodness of fit (Figure 2). The plot suggests that the model adequately explains these data.

4. Future directions for unmarked development

unmarked has become a stable and useful platform for the analysis of ecological data, but several areas of development could improve its utility. First, new models need to be added to cover the range of sampling techniques and population dynamics commonly encountered. Table 3 illustrates the current gaps that need to be filled. In most cases, models to fill these gaps have not been developed so more research is needed.

	Population Dynamics				
Sampling method	Closed	Open to movement	Open to demographic processes		
Occurrence sampling	occu	_	colext		
Repeated counts	pcount	_	pcountOpen		
Removal sampling,					
double observer sampling,	${\tt multinomPois}$	gmultmix	_		
and other multinomial designs					
Distance-sampling	distsamp	_	_		

Table 3: Model fitting functions classified by sampling method and population dynamics. Missing cells indicate models that have not been developed but are likely to be investigated in the future.

Second, each of the models in **unmarked** assumes independence among sites. However, ecologists often use sampling methods such as cluster sampling that induce spatial dependence. Typically, this is done for logistical convenience, but because few methods are available to account for spatial correlation and imperfect detection probability, the spatial dependence is often ignored. Rather than this being a weakness of the sampling design, we envision that this dependence can be used as information regarding the spatial distribution of individuals.

Third, many of the likelihoods are written in pure R, which can be slow for large problems. We are currently translating many of these functions into C++ with the help of the R package **Rcpp** (Eddelbuettel *et al.* 2011).

Finally, Bayesian estimation could be implemented for many of these models. An important advantage of Bayesian analysis over classical methods is that the latent abundance or occurrence variables can be treated as formal parameters. Thus posterior distributions could easily be calculated for derived parameters such as the proportion of sites occupied. Bayesian analysis also would provide a natural framework for incorporating additional sources of random variation. For example, one could model heterogeneity among sites not accounted for by covariates alone.

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