Fitting occupancy models with sp0ccupancy

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October 07, 2021

Contents

1	Introduction 1.1 Example data set: Foliage-gleaning birds at Hubbard Brook	1 1
2	Single species occupancy models	2
_	2.1 Basic model description	2
	2.2 Fitting single species occupancy models with PGOcc	3
	2.3 Convergence diagnostics	6
	2.4 Posterior predictive checks	9
	2.5 Model selection using WAIC	11
	2.6 Prediction	12
	2.0 Trediction	12
3	8	14
	3.1 Basic model description	14
	3.2 Fitting single species spatial occupancy models with spPGOcc	15
	3.3 Convergence diagnostics	19
	3.4 Posterior predictive checks	21
		22
	3.6 Prediction	22
4	Multispecies occupancy models	23
	4.1 Basic model description	23
	4.2 Fitting multispecies occupancy models with msPGOcc	23
	4.3 Convergence diagnostics	29
	4.4 Posterior predictive checks	30
	4.5 Model selection using WAIC	31
	4.6 Prediction	31
5	Multispecies spatial occupancy models	32
		32
	5.2 Fitting multispecies spatial occupancy models with spMsPGOcc	33
	5.3 Convergence diagnostics	41
		42
	•	43
		43
c	Single anguing intermeted accurage weedels	4.4
6	8	44
	The state of the s	44
	0 0	45
		46
	6.4 Convergence diagnostics	51
	6.5 Posterior predictive checks	51

	6.6	Model selection using WAIC	52
	6.7	Prediction	52
7	Sing	gle species spatial integrated occupancy models	53
	7.1	Basic model description	53
	7.2	Simulating data using simIntOcc	53
	7.3	Fitting single speces spatial integrated occupancy models using spIntPGOcc	54
		Convergence diagnostics	
	7.5	Posterior predictive checks	59
	7.6	Model selection using WAIC	60
		Prediction	
\mathbf{Re}	efere	nces	62

1 Introduction

This vignette provides worked examples and explanations on fitting single species and multispecies occupancy models available in the sp0ccupancy R package. We will provide step by step examples on how to fit the following models:

- 1. Occupancy model using PGOcc.
- 2. Spatial occupancy model using spPGOcc.
- 3. Multispecies occupancy model using msPGOcc.
- 4. Spatial multispecies occupancy model using spMsPGOcc.
- 5. Integrated occupancy model using intPGOcc.
- 6. Spatial integrated occupancy model using spIntPGOcc.

In this vignette, we will provide a brief description of each model, with full statistical details provided in a separate MCMC sampler vignette. We will also show how sp0ccupancy provides functions for posterior predictive checks as a Goodness of Fit assessment, model comparison and assessment using the Widely Applicable Information Criterion (WAIC), and out of sample predictions using standard R helper functions (i.e., predict). sp0ccupancy also provides functionality for performing k-fold cross-validation, which is described in a separate vignette.

To get started, we load the spOccupancy package, as well as the coda package, which we will use for some MCMC diagnostics. We will also use the stars and ggplot2 packages to create some very basic plots of our results.

```
library(sp0ccupancy)
library(coda)
library(stars)
library(ggplot2)
```

1.1 Example data set: Foliage-gleaning birds at Hubbard Brook

As an example data set throughout this vignette, we will use data from twelve foliage-gleaning birds collected from point count surveys at Hubbard Brook Experimental Forest (HBEF) in New Hampshire, USA. Specific details on the data set are available on the Hubbard Brook website and Doser et al. (2021). The data are provided in the sp0ccupancy package and are loaded with data(hbef2015). Some brief information on the data collection protocol and the species included in the data set are found via help(hbef2015).

```
data(hbef2015)
str(hbef2015)
```

```
List of 4
           : num [1:12, 1:373, 1:3] 0 0 0 1 0 1 1 0 0 0 ...
 $ у
  ..- attr(*, "dimnames")=List of 3
  ....$ : chr [1:12] "AMRE" "BAWW" "BHVI" "BLBW" ...
  ....$ : chr [1:373] "1" "2" "3" "4" ...
  ....$ : chr [1:3] "1" "2" "3"
 $ occ.covs: num [1:373, 1:2] -0.889 -0.765 -0.413 -0.14 -0.13 ...
  ..- attr(*, "dimnames")=List of 2
  .. ..$ : NULL
  .. ..$ : chr [1:2] "Elevation" "Elevation.2"
 $ det.covs:List of 3
  ..$ day : num [1:373, 1:3] -1.62 -1.62 -1.62 -1.62 -1.62 ...
  ..$ tod : num [1:373, 1:3] -1.565 -1.378 -1.084 -0.79 -0.549 ...
  ..$ day.2: num [1:373, 1:3] 2.61 2.61 2.61 2.61 2.61 ...
 $ coords : num [1:373, 1:2] 280000 280000 280000 280001 280000 ...
  ..- attr(*, "dimnames")=List of 2
  ....$ : chr [1:373] "1" "2" "3" "4" ...
  ....$ : chr [1:2] "Easting" "Northing"
```

The object hbef2015 is a list comprised of the detection-nondetection data (y), covariates on the occurrence portion of the model (occ.covs), covariates on the detection portion of the model (det.covs), and the spatial coordinates of each site (coords) for use in spatial occupancy models and plotting. This list is in the exact format required for input to sp0ccupancy model functions. hbef2015 contains data on 12 species in the three-dimensional array y, where the dimensions of y correspond to species (12), sites (373), and replicates (3). Here we will use data on the charming Ovenbird (OVEN; Seiurus aurocapilla) to display single species models, so we next subset the hbef2015 list to only include data from OVEN in a new object ovenHBEF.

```
sp.names <- attr(hbef2015$y, "dimnames")[[1]]
ovenHBEF <- hbef2015
ovenHBEF$y <- ovenHBEF$y[sp.names == "OVEN", , ]
table(ovenHBEF$y)

0 1</pre>
```

We see OVEN is detected at around half of all site-replicate combinations.

2 Single species occupancy models

2.1 Basic model description

569 539

Let z_j be the true presence (1) or absence (0) of a species at site j, with j = 1, ..., J. We assume this latent occurrence process arises from a Bernoulli process following

$$z_j \sim \text{Bernoulli}(\psi_j),$$

 $\text{logit}(\psi_j) = \mathbf{x}'_j \cdot \boldsymbol{\beta},$ (1)

where ψ_j is the probability of occurrence at site j, which is a function of site-specific covariates X and a vector of regression coefficients (β) .

We do not directly observe z_j and rather we observe an imperfect representation of the latent occurrence process. Let $y_{j,k}$ be the observed detection (1) or nondetection (0) of a species of interest at site j during

replicate k for each of $k = 1, ..., K_j$ replicates. We envision the detection-nondetection data as arising from a Bernoulli process conditional on the true latent occurrence process:

$$y_{j,k} \sim \text{Bernoulli}(p_{j,k} \cdot z_j),$$

 $\text{logit}(p_{j,k}) = \mathbf{v}'_{j,k} \cdot \boldsymbol{\alpha},$ (2)

where $p_{j,k}$ is the probability of detecting a species at site j during replicate k (given it is present at site j), which is a function of site and replicate specific covariates V and a vector of regression coefficients (α).

To complete the Bayesian specification of the model, we assign multivariate normal priors for the occurrence (β) and detection (α) regression coefficients. To yield an efficient implementation of the occupancy model using a logit link function, we use Pólya-Gamma data augmentation (Polson, Scott, and Windle 2013), which is described in depth in a separate MCMC sampler vignette (Polya-Gamma is where the PG comes from in all sp0ccupancy model fitting functions).

2.2 Fitting single species occupancy models with PGOcc

The PGOcc function fits single species occupancy models using Pólya-Gamma latent variables, which makes it more efficient than standard implementations of occupancy models using a logit link function (Clark and Altwegg 2019; Polson, Scott, and Windle 2013). PGOcc has the following arguments:

```
PGOcc(occ.formula, det.formula, data, starting, priors, n.samples,
    n.omp.threads = 1, verbose = TRUE, n.report = 100,
    n.burn = round(.10 * n.samples), n.thin = 1,
    k.fold, k.fold.threads = 1, k.fold.seed, ...)
```

The last three arguments (k.fold, k.fold.threads, and k.fold.seed) provide functionality for performing k-fold cross-validation. These three arguments are in all spOccupancy model fitting functions. In this vignette we won't perform cross-validation and will leave these arguments alone (see the separate cross-validation vignette for details on performing k-fold cross-validation).

The first two arguments, occ.formula and det.formula, use standard R model syntax to denote the covariates included in the occurrence and detection portions of the model, respectively. Only the right hand side of the formulas are included. Random intercepts can be included in both the occurrence and detection portions of the occupancy model using lme4 syntax (Bates et al. 2015). The names of variables given in the formulas should correspond to those found in data, which is a list consisting of the following tags: y (detection-nondetection data), occ.covs (occurrence covariates), det.covs (detection covariates). y should be stored as a sites x replicate matrix, occ.covs as a matrix or data frame with site-specific covariate values, and det.covs as a list with each list element corresponding to a covariate to include in the detection portion of the model. Covariates on detection can vary by site and/or survey, and so these covariates may be specified as a site by survey matrix for survey-level covariates or as a one-dimensional vector for survey level covariates. The ovenHBEF list is already in the required format. Here we will model OVEN occurrence as a function of linear and quadratic elevation and will include three observational covariates (linear and quadratic day of survey, time of day of survey) on the detection portion of the model. We specify the formulas below

```
oven.occ.formula <- ~ Elevation + Elevation.2
oven.det.formula <- ~ day + tod + day.2
# Check out the format of ovenHBEF
str(ovenHBEF)

List of 4
$ y : num [1:373, 1:3] 1 1 0 1 0 0 1 1 1 1 ...
..- attr(*, "dimnames")=List of 2
....$ : chr [1:373] "1" "2" "3" "4" ...</pre>
```

```
....$ : chr [1:3] "1" "2" "3"
$ occ.covs: num [1:373, 1:2] -0.889 -0.765 -0.413 -0.14 -0.13 ...
..- attr(*, "dimnames")=List of 2
....$ : NULL
....$ : chr [1:2] "Elevation" "Elevation.2"
$ det.covs:List of 3
..$ day : num [1:373, 1:3] -1.62 -1.62 -1.62 -1.62 -1.62 ...
..$ tod : num [1:373, 1:3] -1.565 -1.378 -1.084 -0.79 -0.549 ...
..$ day.2: num [1:373, 1:3] 2.61 2.61 2.61 2.61 2.61 ...
$ coords : num [1:373, 1:2] 280000 280000 280000 280001 280000 ...
..- attr(*, "dimnames")=List of 2
....$ : chr [1:373] "1" "2" "3" "4" ...
....$ : chr [1:2] "Easting" "Northing"
```

Note that the covariates in hbef2015 are already standardized to have mean 0 and standard deviation 1. If the data were not already standardized, we could do the standardization directly in the formula specification (e.g., ~ scale(Elevation) + scale(Elevation)^2).

Next, we specify the starting values for the MCMC sampler in starting. PGOcc (and all other spOccupancy model fitting functions) will set starting values by default, but here we will do this explicitly. Starting values are specified in a list with the following tags: z (latent occurrence values), alpha (detection regression coefficients), and beta (occurrence regression coefficients). Below we set all initial values of the regression coefficients to 0, and set starting values for z based on the detection-nondetection data matrix.

We next specify the priors for the occurrence and detection regression coefficients. The Pólya-Gamma data augmentation algorithm employed by sp0ccupancy assumes normal priors for both the detection and occurrence regression coefficients. These priors are specified in a list with tags beta.normal for occurrence and alpha.normal for detection parameters. Each list element is then itself a list, with the first element of the list consisting of the hypermeans for each coefficient to be estimated and the second element of the list consisting of the hypervariances for each coefficient. By default, sp0ccupancy will set the hypermeans to 0 and the hypervariances to 2.72, which corresponds to a relatively flat prior on the probability scale (0, 1) (Broms, Hooten, and Fitzpatrick 2016). We will use these default priors here, but we specify them explicitly below for clarity

Our last step is to specify the number of samples to run the MCMC (n.samples), the amount of burn-in (n.burn), and how often we want to thin the posterior samples (n.thin). For a simple single species occupancy model, we shouldn't need too many samples and will only need a small amount of burn-in and minimal thinning.

```
n.samples <- 5000
n.burn <- 3000
n.thin <- 2
```

We are now set to run the occupancy model. Single species occupancy models are fast, and so we set n.omp.threads = 1 to indicate we won't use multiple threads to run the model. For more time consuming

models, we can set n.omp.threads to a number greater than 1 and smaller than the number of threads on the computer you are using. Note this argument will only use multiple threads if spOccupancy was compiled for OpenMP support. The verbose argument is a logical value indicating whether or not MCMC sampler progress is reported to the screen. If verbose = TRUE, sampler progress is reported after the specified number of iterations in the n.report argument. We set verbose = TRUE and n.report = 1000 to report progress after every 1000th MCMC iteration.

```
out <- PGOcc(occ.formula = oven.occ.formula,</pre>
           det.formula = oven.det.formula,
           data = ovenHBEF,
           starting = oven.starting,
           n.samples = n.samples,
           priors = oven.priors,
           n.omp.threads = 1,
           verbose = TRUE,
           n.report = 1000,
           n.burn = n.burn,
           n.thin = n.thin)
   Preparing the data
   Model description
_____
Occupancy model with Polya-Gamma latent
variable fit with 373 sites.
Number of MCMC samples: 5000
Burn-in: 3000
Thinning Rate: 2
Total Posterior Samples: 1000
Source compiled with OpenMP support and model fit using 1 thread(s).
Sampling ...
Sampled: 1000 of 5000, 20.00%
Sampled: 2000 of 5000, 40.00%
_____
Sampled: 3000 of 5000, 60.00%
Sampled: 4000 of 5000, 80.00%
Sampled: 5000 of 5000, 100.00%
```

```
[1] "beta.samples" "alpha.samples" "z.samples" "psi.samples" [5] "y.rep.samples" "X" "X.p" "y" [9] "n.samples" "call" "n.post" "n.thin" [13] "n.burn" "pRE" "psiRE" "run.time"
```

names(out)

PGOcc returns a list of class PGOcc with a suite of different objects, many of them being coda::mcmc objects of posterior samples. Notice the "Preparing the data" printed section doesn't have any information shown in

it. spOccupancy model fitting functions will present messages when preparing the data for the model in this section, or will print out the default priors or starting values used when they are not specified in the function call. Here we specified everything explicitly so no information was reported.

For a nice summary of the regression parameters we can use summary on the resulting PGOcc object.

```
summary(out)
```

Call:

```
PGOcc(occ.formula = oven.occ.formula, det.formula = oven.det.formula,
  data = ovenHBEF, starting = oven.starting, priors = oven.priors,
  n.samples = n.samples, n.omp.threads = 1, verbose = TRUE,
  n.report = 1000, n.burn = n.burn, n.thin = n.thin)
```

Chain Information:

Total samples: 5000

Burn-in: 3000

Thin: 2

Total Posterior Samples: 1000

Occurrence:

```
2.5% 25% 50% 75% 97.5% (Intercept) 2.0113 2.3454 2.5489 2.8095 3.2653 Elevation -1.9424 -1.3525 -1.2050 -1.0772 -0.8654 Elevation.2 -1.0730 -0.8508 -0.7442 -0.6335 -0.3116
```

Detection:

```
2.5% 25% 50% 75% 97.5% (Intercept) 0.3631 0.5527 0.6331 0.7277 0.8748 day -0.1588 -0.0755 -0.0346 0.0168 0.0961 tod -0.2550 -0.1727 -0.1280 -0.0789 0.0024 day.2 -0.3118 -0.2213 -0.1655 -0.1133 0.0095
```

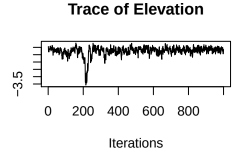
Note that all coefficients are printed on the logit scale. We see OVEN is fairly prominent in the forest given the large intercept value, and the negative linear and quadratic terms for Elevation suggest occurrence probability peaks at mid-elevations.

2.3 Convergence diagnostics

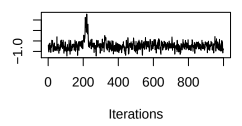
The posterior samples in the PGOcc object are coda::mcmc objects, which we can quickly assess for convergence visually using trace plots.

```
plot(out$beta.samples, density = FALSE)
```

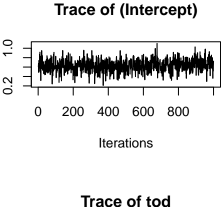
Trace of (Intercept) 9. 0 200 400 600 800 Iterations

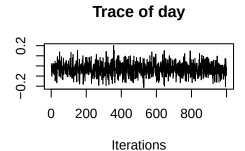


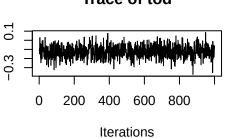


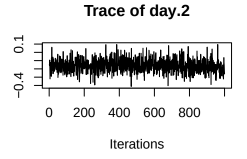


plot(out\$alpha.samples, density = FALSE)









For a complete analysis (i.e., in a peer-reviewed manuscript), we will likely want to more formally check for convergence, perhaps using the Gelman-Rubin R-hat diagnostic (Brooks and Gelman 1998). This requires running multiple chains at largely different starting values for the regression parameters. For a single species

non-spatial occupancy model, we can accomplish this by running multiple chains sequentially (since they run really fast) with different starting values, then combining the output into a coda::mcmc.list object for use with the coda::gelman.diag function. Notice below we set verbose = FALSE to suppress the messages printed by PGOcc.

```
oven.starting <- list(alpha = rep(2, p.det),</pre>
                      beta = rep(2, p.occ),
                       z = apply(ovenHBEF$y, 1, max, na.rm = TRUE))
out.2 <- PGOcc(occ.formula = oven.occ.formula,</pre>
               det.formula = oven.det.formula,
               data = ovenHBEF,
               starting = oven.starting,
               n.samples = n.samples,
               priors = oven.priors,
               n.omp.threads = 1,
               verbose = FALSE,
               n.report = 1000,
               n.burn = n.burn,
               n.thin = n.thin)
oven.starting <- list(alpha = rep(-2, p.det),
                      beta = rep(-2, p.occ),
                       z = apply(ovenHBEF$y, 1, max, na.rm = TRUE))
out.3 <- PGOcc(occ.formula = oven.occ.formula,</pre>
               det.formula = oven.det.formula,
               data = ovenHBEF,
               starting = oven.starting,
               n.samples = n.samples,
               priors = oven.priors,
               n.omp.threads = 1,
               verbose = FALSE,
               n.report = 1000,
               n.burn = n.burn,
               n.thin = n.thin)
# beta convergence
gelman.diag(mcmc.list(out$beta.samples, out.2$beta.samples,
                       out.3$beta.samples))
```

Potential scale reduction factors:

```
Point est. Upper C.I.
(Intercept) 1.02 1.06
Elevation 1.02 1.04
Elevation.2 1.00 1.01
```

Multivariate psrf

1.02

Potential scale reduction factors:

Point est. Upper C.I.

```
(Intercept) 1 1.01
day 1 1.00
tod 1 1.01
day.2 1 1.01
```

Multivariate psrf

1.01

All R-hat values are less than 1.1, indicating the chains have converged and we are in good shape to proceed.

2.4 Posterior predictive checks

Fit statistic: freeman-tukey

The function ppcOcc performs a posterior predictive check on all spOccupancy model objects as a Goodness of Fit (GoF) assessment. The fundamental idea of a posterior predictive check is as follows: our model should generate data that closely align with the observed data. If there are drastic differences in the true data from the model generated data, our model likely is not very useful (Hobbs and Hooten 2015). GoF assessments are more complicated using binary data, like detection-nondetection used in occupancy models, as standard approaches are not valid assessments for binary data (Broms, Hooten, and Fitzpatrick 2016; McCullagh 2019). Thus, any approach to assess model fit for detection-nondetection data must bin the raw values in some manner, and then perform a model fit assessment on the binned values. There are numerous ways we could envision binning the raw detection-nondetection values (Kéry and Royle 2015).

The resulting PGOcc model object is sent as input to the ppcOcc function, along with a fit statistic (fit.stat) and numeric value indicating how to group the data (group). Currently supported fit statistics include the Freeman-Tukey statistic and the Chi-Square statistic (freeman-tukey or chi-square, respectively, Kéry and Royle (2015)). Currently, ppcOcc allows the user to group the data by row (site; group = 1) or column (replicate; group = 2). ppcOcc will then return a set of posterior samples for the fit statistic (or discrepancy measure) using the observed data (fit.y) and model generated data set (fit.y.rep), summed across all data points. These values can be used with the summary function to generate a Bayesian p-value. Bayesian p-values are sensitive to individual values, so we should also explore the discrepancy measures for each "grouped" data point. ppcOcc returns a matrix of posterior quantiles for the fit statistic for both the observed (fit.y.group.quants) and model generated data (fit.y.rep.group.quants) for each "grouped" data point.

We next perform a posterior predictive check using the Freeman-Tukey statistic grouping the data by sites. We summarize the posterior predictive check with the summary function, which reports a Bayesian p-value. A Bayesian p-value that hovers around 0.5 indicates adequate model fit, while values less than 0.1 or greater than 0.9 suggest our model does not fit the data well (Hobbs and Hooten 2015).

```
ppc.out <- ppcOcc(out, fit.stat = 'freeman-tukey', group = 1)
summary(ppc.out)

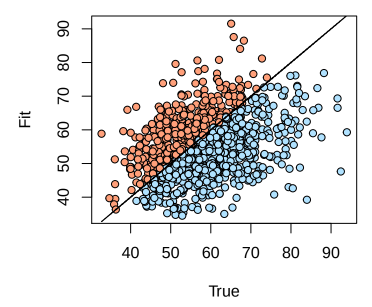
Call:
ppcOcc(object = out, fit.stat = "freeman-tukey", group = 1)

Chain Information:
Total samples: 5000
Burn-in: 3000
Thin: 2
Total Posterior Samples: 1000

Bayesian p-value: 0.379</pre>
```

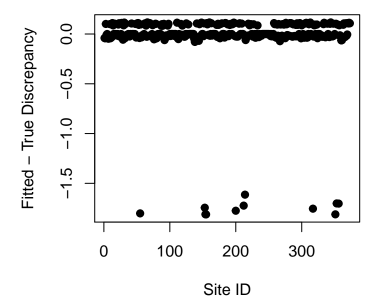
The Bayesian p-value is the proportion of posterior samples of the fit statistic of the model generated data that are greater than the corresponding fit statistic of the true data, summed across all "grouped" data points.

We can create a visual representation of the Bayesian p-value as follows, which is highly motivated by Kéry and Royle (2015).



Our Bayesian p-value indicates no lack of fit for the OVEN model. However, relying solely on the Bayesian p-value as an assessment of model fit is not always a great option, as individual data points can have an overbearing influence on the resulting summary value. Instead of summing across all data points for a single discrepancy measure, ppcOcc also allows us to explore discrepancy measures on a "grouped" point by point basis. The resulting ppcOcc object will contain the objects fit.y.group.quants and fit.y.rep.group.quants, which contain quantiles of the posterior distributions for the discrepancy measures of each grouped data point. Below we plot the difference in the discrepancy measure between the fitted and true data across each of the sites.

```
diff.fit <- ppc.out$fit.y.rep.group.quants[3, ] - ppc.out$fit.y.group.quants[3, ]
plot(diff.fit, pch = 19, xlab = 'Site ID', ylab = 'Fitted - True Discrepancy')</pre>
```



We see there are a few sites where the true discrepancy is much larger than the discrepancy under the fitted data. Here we will ignore this, but in a real analysis we would explore these sites further to see what could explain this pattern (e.g., are the sites close together in space?).

2.5 Model selection using WAIC

Posterior predictive checks allow us to assess how well our model fits the data, but they are not very useful if we want to compare multiple competing models and ultimately select a final model based on some criterion. Bayesian model selection is very much a constantly changing field, especially in the ecological and environmental sciences. See Hooten and Hobbs (2015) for an accessible overview of Bayesian model selection for ecologists.

For Bayesian hierarchical models like occupancy models, the most common Bayesian model selection criterion, DIC, is not applicable (Hooten and Hobbs 2015). Instead, we can use the Widely Applicable Information Criterion (Watanabe 2010) to compare a set of models and select the best performing model according to the WAIC for final analysis. When focused primarily on predictive performance, a k-fold cross-validation approach is another attractive (but more computationally intensive) alternative to compare a series of models, especially since WAIC may not be a reliable metric for all models (Link, Sauer, and Niven 2020). In spOccupancy, k-fold cross-validation is accomplished using the arguments k.fold, k.fold.threads, and k.fold.seed in the model fitting function. We detail how to perform k-fold cross-validation using spOccupancy model objects in a separate vignette.

The WAIC is calculated for all spOccupancy model objects using the function waicOcc. We calculate the WAIC as

$$WAIC = -2 \times (elpd - pD),$$

where elpd is the expected log pointwise predictive density and PD is the effective number of parameters. We calculate elpd by calculating the likelihood for each posterior sample, taking the mean of these likelihoods,

taking the log of the mean of the likelihoods, and summing these values across all sites. We calculate the effective number of parameters by calculating the variance of the log likelihood for each site taken over all posterior samples, and then summing these values across all sites. See Appendix S1 from Broms, Hooten, and Fitzpatrick (2016) for more details.

We calculate the WAIC using waicOcc for our OVEN model below.

```
waicOcc(out)
```

```
elpd pD WAIC
-685.275085 7.014193 1384.578556
```

Next we rerun the OVEN model, but this time we assume occurrence is constant across the HBEF, and subsequently compare the WAIC value to the full model

```
p.occ <- 1
oven.starting <- list(alpha = rep(0, p.det),</pre>
                       beta = rep(0, p.occ),
                       z = apply(ovenHBEF$y, 1, max, na.rm = TRUE))
oven.priors <- list(alpha.normal = list(mean = rep(0, p.det),</pre>
                                          var = rep(2.72, p.det)),
                     beta.normal = list(mean = rep(0, p.occ),
                                         var = rep(2.72, p.occ)))
out.small <- PGOcc(occ.formula = ~ 1,</pre>
           det.formula = oven.det.formula,
           data = ovenHBEF,
           starting = oven.starting,
           n.samples = n.samples,
           priors = oven.priors,
           n.omp.threads = 1,
           verbose = FALSE,
           n.burn = n.burn,
           n.thin = n.thin)
waicOcc(out.small)
```

```
elpd pD WAIC
-729.225526 4.894526 1468.240104
```

Smaller values of WAIC indicate models with better performance. We see the WAIC for the model with elevation is smaller than the intercept only model, indicating elevation is an important predictor for OVEN occurrence in HBEF.

2.6 Prediction

All resulting model objects from sp0ccupancy model functions can be used with predict to generate a series of posterior predictive samples at non-sampled locations, given the values of all covariates used in the model fitting process. The object hbefElev (provided in the sp0ccupancy package) contains elevation values at a 30x30m resolution from the National Elevation Dataset across the entire HBEF. The values are standardized using the mean and standard deviation of the elevation values used to fit the model. We load the data below

```
data(hbefElev)
str(hbefElev)
```

```
'data.frame': 46091 obs. of 3 variables:

$ val : num 2.07 2.08 2.09 2.1 2.12 ...

$ Easting : num 276269 276291 276314 276336 276358 ...

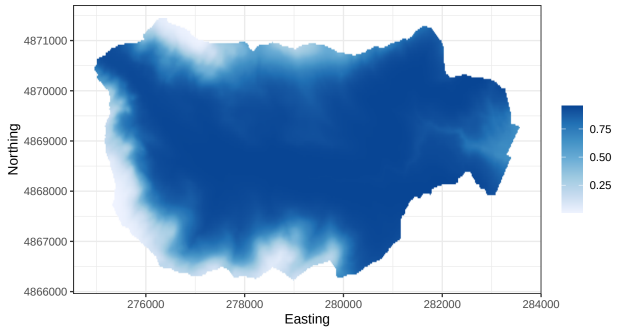
$ Northing: num 4871425 4871425 4871425 4871425 ...
```

The column val contains the standardized elevation values, while Easting and Northing contain the spatial coordinates that we will use for plotting. We can obtain posterior predictive samples for the occurrence probabilities at these sites by using the predict function and our PGOcc model object.

```
X.0 <- cbind(1, hbefElev$val, hbefElev$val^2)
out.pred <- predict(out, X.0)</pre>
```

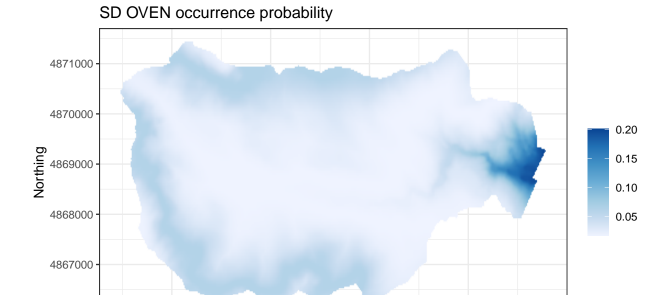
For PGOcc objects, the predict function takes two arguments: (1) the PGOcc model object; and (2) a matrix or data frame consisting of the design matrix for the prediction locations (including an intercept). The resulting object consists of posterior predictive samples for the latent occurrence probabilities (psi.O.samples) and latent occurrence values (z.O.samples). The beauty of the Bayesian paradigm is that these predictions all have fully propagated uncertainty. We can use these values to create plots of the predicted mean occurrence values, as well as their standard deviation.

Mean OVEN occurrence probability



```
ggplot() +
  geom_stars(data = dat.stars, aes(x = x, y = y, fill = sd.psi)) +
  scale_fill_distiller(palette = 'Blues', direction = 1, na.value = 'transparent') +
  labs(x = 'Easting', y = 'Northing', fill = '',
```

title = 'SD OVEN occurrence probability') +
theme_bw()



3 Single species spatial occupancy models

278000

3.1 Basic model description

276000

4866000

When working across large spatial domains, accounting for residual spatial autocorrelation in species distributions can often improve predictive performance, leading to more accurate species distribution maps (Guélat and Kéry 2018; Lany et al. 2020). We extend the basic single species occupancy model to incorporate a spatial Gaussian Process that accounts for unexplained spatial variation in species occurrence across a region of interest. The species-specific occurrence probability at site j, ψ_j , now takes the form

Easting

280000

$$logit(\psi_j) = \mathbf{x}_j' \cdot \boldsymbol{\beta} + \mathbf{w}_j, \tag{3}$$

282000

284000

where w_j is a realization from a zero-mean spatial Gaussian Process, i.e.,

$$\mathbf{w} \sim N(\mathbf{0}, \mathbf{\Sigma}(\mathbf{s}, \mathbf{s'}, \boldsymbol{\theta})).$$
 (4)

We define $\Sigma(s, s', \theta)$ as a $J \times J$ covariance matrix that is a function of the distances between any pair of site coordinates s and s' and a set of parameters (θ) that govern the spatial process. The vector θ is equal to $\theta = \{\sigma^2, \phi, \nu\}$, where σ^2 is a spatial variance parameter, ϕ is a spatial decay parameter, and ν is a spatial smoothness parameter. ν is only specified when using a Matern correlation function.

The detection portion of the occupancy model remains unchanged from the non-spatial occupancy model and follows Equation (2). Single species spatial occupancy models, like all models in sp0ccupancy are fit using Pólya-Gamma data augmentation (see MCMC sampler vignette for details).

When the number of sites is moderately large, say 1000, the above described spatial Gaussian process model can be drastically slow as a result of needing to take the inverse of the spatial covariance matrix $\Sigma(s, s', \theta)$ at each MCMC iteration. Numerous approximation methods exist to reduce this computational cost (Heaton et al. 2019). One attractive approach is the Nearest Neighbor Gaussian Process (NNGP; Datta et al. (2016)). Instead of modeling the spatial process using a full Gaussian Process as shown in Equation (4), we replace the Gaussian Process prior specification with a NNGP, which leads to drastic increases in run time with nearly identicial inference and prediction as the full Gaussian Process specification. See Datta et al. (2016), Finley et al. (2019), and the MCMC sampler vignette for additional statistical details on NNGPs and their implementation in spatial occupancy models.

3.2 Fitting single species spatial occupancy models with spPGOcc

The function spPGOcc fits single species spatial occupancy models using Pólya-Gamma latent variables, where spatial autocorrelation is accounted for using a spatial Gaussian Process. spPGOcc fits saptial occupancy models using either a full Gaussian process or an NNGP. See Finley, Datta, and Banerjee (2020) for details on using NNGPs with Polya-Gamma latent variables.

We will fit the same occupancy model for OVEN that we fit previously using PGOcc, but we will now make the model spatially explicit by incorporating a spatial process with spPGOcc. First, let's take a look at the arguments for spPGOcc:

We will walk through each of the arguments to spPGOcc in the context of our Ovenbird example. The occurrence (occ.formula) and detection (det.formula) formulas, as well as the list of data (data), take the same form as we saw in PGOcc, with the exception that random intercepts can only be specified in det.formula. Notice the coords matrix in the ovenHBEF list of data. We did not use this for PGOcc but specifying the spatial coordinates in data is required for all spatially explicit models in spOccupancy.

```
oven.occ.formula <- ~ Elevation + Elevation.2
oven.det.formula <- ~ day + tod + day.2
str(ovenHBEF) # coords is required for spPGOcc.</pre>
```

```
List of 4
 $ y
           : num [1:373, 1:3] 1 1 0 1 0 0 1 1 1 1 ...
  ..- attr(*, "dimnames")=List of 2
  ....$ : chr [1:373] "1" "2" "3" "4" ...
  .. ..$ : chr [1:3] "1" "2" "3"
 $ occ.covs: num [1:373, 1:2] -0.889 -0.765 -0.413 -0.14 -0.13 ...
  ..- attr(*, "dimnames")=List of 2
  .. ..$ : NULL
  ....$ : chr [1:2] "Elevation" "Elevation.2"
 $ det.covs:List of 3
  ..$ day : num [1:373, 1:3] -1.62 -1.62 -1.62 -1.62 -1.62 ...
  ..$ tod : num [1:373, 1:3] -1.565 -1.378 -1.084 -0.79 -0.549 ...
  ..$ day.2: num [1:373, 1:3] 2.61 2.61 2.61 2.61 2.61 ...
 $ coords : num [1:373, 1:2] 280000 280000 280000 280001 280000 ...
  ..- attr(*, "dimnames")=List of 2
```

```
....$: chr [1:373] "1" "2" "3" "4" ...
....$: chr [1:2] "Easting" "Northing"
```

The starting values (starting) are again specified in a list. Valid tags for starting values now additionally include the parameters associated with the spatial random effects. These include: sigma.sq (spatial variance parameter), phi (spatial range parameter), w (the latent spatial random effects at each site), and nu (spatial smoothness parameter). nu is only specified if using a Matern covariance function (i.e., cov.model = 'matern'). sp0ccupancy supports four spatial covariance models (exponential, spherical, gaussian, and matern), which are specified in the cov.model argument. Here we will use an exponential covariance model. As a starting value for the spatial range parameter phi, we compute the mean distance between points in HBEF and then set it equal to 3 divided by this mean distance. When using an exponential covariance function, $\frac{3}{\phi}$ is the effective range, or the distance at which the residual spatial correlation between two sites is 0.05 (Banerjee, Carlin, and Gelfand 2003). Thus our initial guess for this effective range is the average distance betweeen sites across HBEF.

The next three arguments (n.batch, batch.length, and accept.rate) are all related to the Adaptive MCMC sampler we use to fit the model. Updates for the spatial range parameter (and smoothness parameter if cov.model = 'matern') require the use of a Metropolis Hastings algorithm. We implement an adaptive Metropois-Hastings algorithm discussed in Roberts and Rosenthal (2009). This algorithm adjusts the tuning values for each parameter that requires a Metropolis-Hastings update within the sampler itself. This process results in a more efficient sampler than if we were to fix the tuning parameters prior to fitting the model. The parameter accept.rate is the target acceptance rate for each parameter, and the algorithm will adjust the tuning parameters to hover around this value. The default value is 0.43, which we suggest leaving as is unless you have a good reason to change it. The tuning parameters are updated after a single "batch". We must specify the total n.batch batches, where each "batch" consists of batch.length MCMC samples. Thus, the total number of MCMC samples is n.batch * batch.length. Typically, we set batch.length = 25 and then play around with n.batch until convergence is reached. Here we set n.batch = 400 for a total of 10000 MCMC samples. We will additionally specify a burn-in period of 2000 samples and a thinning rate of 8. We also need to specify an initial value for the tuning parameters for the spatial decay and smoothness parameters (if applicable). These values are sent as input in the form of a list with tags phi and nu. The initial tuning value can be any value greater than 0, but we recommend starting the value out around 0.5. After some initial runs of the model, if you notice the final acceptance rate of a parameter is much larger or smaller than the target acceptance rate (accept.rate), you can then change the initial tuning value to get closer to the target rate. Here we set the initial tuning value for phi to 1 after some initial runs of the model.

```
batch.length <- 25
n.batch <- 400
n.burn <- 2000
n.thin <- 8
oven.tuning <- list(phi = 1)</pre>
```

Priors are again specified in a list in the argument priors. We assume an inverse gamma prior for the spatial

variance parameter sigma.sq (tag is sigma.sq.ig), and uniform priors for the spatial decay parameter phi and smoothness parameter nu (if Matern), with the associated tags phi.unif and nu.unif. The hyperparameters of the inverse Gamma are passed as a vector of length two, with the first and second elements corresponding to the shape and scale, respectively. The lower and upper bounds of the uniform distribution are passed in as a two-element vector for the uniform priors.

The priors for the spatial parameters in a spatially-explicit model must be at least weakly informative for the model to converge (Banerjee, Carlin, and Gelfand 2003). For the inverse-Gammma prior on the spatial variance, we typically set the shape parameter to 2 and the scale parameter equal to our best guess of the spatial variance. Based on our previous work with these data, we expect the residual spatial variation to be minimal, and so we set the scale parameter below to 1. For the spatial decay parameter, we determine the bounds of the uniform distribution by computing the smallest distance between sites and the largest distance between sites. We then set the lower bound of the uniform to 3/max and the upper bound to 3/min, where min and max correspond to the predetermined distances between sites.

The argument n.omp.threads specifies the number of threads to use for parallelization, while verbose specifies whether or not to print the progress of the sampler. We highly recommend setting verbose = TRUE for all spatial models to ensure the adaptive MCMC is working as you want. The argument n.report specifies the interval to report the Metropolis sampler acceptance. Note that n.report is specified in terms of batches, not the overall number of samples. Below we set n.report = 100, which will result in information on the acceptance rate and tuning parameters every 100th batch.

```
n.omp.threads <- 1
verbose <- TRUE
n.report <- 100</pre>
```

The remaining parameters (NNGP, n.neighbors and search.type) relate to whether or not you want to fit the model with a Gaussian Process or NNGP. The argument NNGP is a logical value indicating whether to fit the model with an NNGP (TRUE) or a regular Gaussian Process (FALSE). For data sets that have more than 1000 locations, using an NNGP will have substantial increases in run time. Even for more modest size data sets (like the HBEF data set), using an NNGP will be quite a bit faster. Unless you are concerned about the NNGP approximation for some reason, we recommend setting NNGP = TRUE. The argument n.neighbors and search.type specify the number of neighbors used in the NNGP and the nearest neighbor search algorithm, respectively, to use for the NNGP model. Generally, the default values of these arguments will be adequate. Datta et al. (2016) showed that setting n.neighbors = 15 is usually sufficient, although for certain data sets a good approximation can be achieved with as small as five neighbors, which could substantially decrease run time. We generally recommend leaving search.type = "cb", as this results in a fast code book nearest neighbor search algorithm. However, details on when you may want to change this are described in Finley, Datta, and Banerjee (2020). We will run an NNGP model using the default value for search.type and setting n.neighbors = 5.

We now fit the model and summarize the results using summary.

```
batch.length = batch.length,
               priors = oven.priors,
               cov.model = cov.model,
               NNGP = TRUE,
               n.neighbors = 5,
               tuning = oven.tuning,
               n.report = n.report,
               n.burn = n.burn,
               n.thin = n.thin)
 -----
   Preparing the data
_____
   Building the neighbor list
Building the neighbors of neighbors list
_____
  Model description
NNGP Occupancy model with Polya-Gamma latent
variable fit with 373 sites.
Number of MCMC samples: 10000 (400 batches of length 25)
Burn-in: 2000
Thinning Rate: 8
Total Posterior Samples: 1000
Using the exponential spatial correlation model.
Using 5 nearest neighbors.
Source compiled with OpenMP support and model fit using 1 thread(s).
Adaptive Metropolis with target acceptance rate: 43.0
Sampling ...
Batch: 100 of 400, 25.00%
   parameter acceptance tuning
   phi 72.0 0.99005
  ._____
Batch: 200 of 400, 50.00%
   parameter acceptance tuning
   phi 76.0 0.81058
Batch: 300 of 400, 75.00%
   parameter acceptance tuning
   phi 12.0 0.63763
Batch: 400 of 400, 100.00%
class(out.sp)
```

[1] "spPGOcc"

```
names(out.sp)
                      "alpha.samples"
                                                         "psi.samples"
 [1] "beta.samples"
                                        "z.samples"
                      "theta.samples"
                                                         "tune"
 [5] "y.rep.samples"
                                        "w.samples"
 [9] "accept"
                      "coords"
                                        " X "
                                                         "X.p"
[13] "y"
                      "call"
                                        "n.samples"
                                                         "n.neighbors"
[17] "cov.model.indx"
                      "type"
                                        "n.post"
                                                         "n.thin"
[21] "n.burn"
                      "pRE"
                                        "run.time"
summary(out.sp)
Call:
spPGOcc(occ.formula = oven.occ.formula, det.formula = oven.det.formula,
   data = ovenHBEF, starting = oven.starting, priors = oven.priors,
   tuning = oven.tuning, cov.model = cov.model, NNGP = TRUE,
   n.neighbors = 5, n.batch = n.batch, batch.length = batch.length,
   n.report = n.report, n.burn = n.burn, n.thin = n.thin)
Chain Information:
Total samples: 10000
Burn-in: 2000
Thin: 8
Total Posterior Samples: 1000
Occurrence:
               2.5%
                        25%
                                50%
                                         75%
(Intercept) 2.2135 2.6537
                             2.9479 3.3085
                                            3.9852
           -2.2817 -1.6332 -1.4138 -1.2129 -0.9516
Elevation.2 -1.3043 -1.0014 -0.8572 -0.7224 -0.3936
Detection:
               2.5%
                        25%
                                50%
                                        75%
                                               97.5%
(Intercept) 0.3797 0.5441 0.6316
                                    0.7293 0.9145
day
            -0.1624 -0.0711 -0.0264 0.0132
tod
            -0.2652 -0.1753 -0.1273 -0.0750 0.0077
            -0.3275 -0.2129 -0.1572 -0.1002 -0.0050
day.2
Covariance:
           2.5%
                   25%
                          50%
                                 75% 97.5%
sigma.sq 0.2491 0.5435 0.9633 1.7365 3.7387
         0.0017 0.0051 0.0102 0.0192 0.0282
```

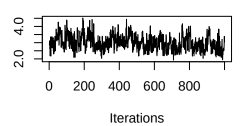
We see spPGOcc returns a list of class spPGOcc and consists of posterior samples for all parameters. Note that posterior samples for spatial parameters are stored in the list element theta.samples. The summary function reveals model results generally align with those found using the non-spatial model.

3.3 Convergence diagnostics

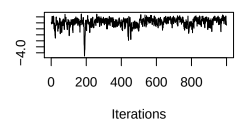
Convergence diagnostics, posterior predictive checks, model selection, and out-of-sample prediction all proceed analogously to what we saw with the non-spatial occupancy model using PGOcc.

```
plot(out.sp$beta.samples, density = FALSE)
```

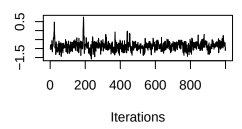
Trace of (Intercept)



Trace of Elevation

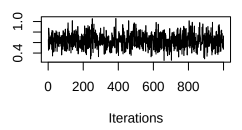


Trace of Elevation.2

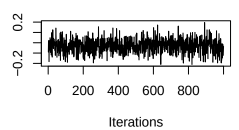


plot(out.sp\$alpha.samples, density = FALSE)

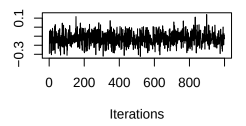
Trace of (Intercept)



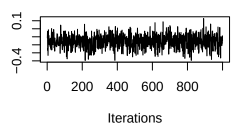
Trace of day



Trace of tod

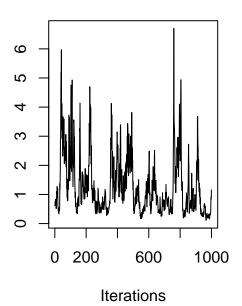


Trace of day.2

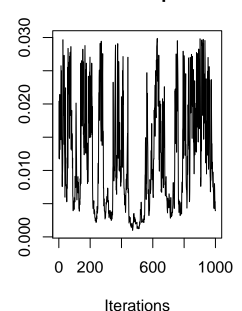


plot(out.sp\$theta.samples, density = FALSE)

Trace of sigma.sq



Trace of phi



We should run the chain for a bit longer to ensure convergence of the spatial parameters, but we'll resist doing so for now. Convergence can be more formally assessed using the Gelman-Rubin diagnostic as done for the nonspatial model.

3.4 Posterior predictive checks

For our posterior predictive check, we send the spPGOcc model object to the ppcOcc function, this time grouping by replicate (group = 2) instead of by site (group = 1).

```
ppc.sp.out <- ppcOcc(out.sp, fit.stat = 'freeman-tukey', group = 2)
summary(ppc.sp.out)</pre>
```

Call:

ppcOcc(object = out.sp, fit.stat = "freeman-tukey", group = 2)

Chain Information: Total samples: 10000

Burn-in: 2000 Thin: 8

Total Posterior Samples: 1000

Bayesian p-value: 0.88
Fit statistic: freeman-tukey

The Bayesian p-value is hovering right around 0.9 potentially suggesting a lack of fit, but we would want to run the model longer to ensure convergence before determining if there is lack of fit.

3.5 Model selection using WAIC

We next use the waicOcc function to compute the WAIC, which we can compare to the non-spatial model to assess the benefit of incorporating the spatial random effects.

```
waicOcc(out.sp)

elpd     pD     WAIC
-659.67490     30.84116 1381.03214

# Compare to non-spatial model
waicOcc(out)

elpd     pD     WAIC
-685.275085     7.014193 1384.578556
```

We see the WAIC value for the spatial model is slightly smaller to that of the nonspatial model, indicating that incorporation of the spatial random effects may yield improvement in predictive performance, although these would likely be quite small. This is not all that surprising, as we expect elevation to soak up most of the spatial variation in OVEN occurrence across the forest.

3.6 Prediction

Finally, we can perform out of sample prediction using the predict function just as before. Out of sample prediction for spatial models is more computationally intensive than non-spatial models, and so the predict function for spPGOcc class objects also has options for parallelization (n.omp.threads) and reporting sampler progress (verbose and n.report). Note that for spPGOcc, you also need to supply the coordinates of the out of sample prediction locations in addition to the covariate values. We do not execute the below code, but it can be used to compare prediction results from the nonspatial to the spatial model (which are extremely similar, except the standard deviations for the spatial predictions are larger).

```
coords.0 <- as.matrix(hbefElev[, c('Easting', 'Northing')])</pre>
out.sp.pred <- predict(out.sp, X.0, coords.0, verbose = FALSE)</pre>
plot.dat <- data.frame(x = hbefElev$Easting,</pre>
                        y = hbefElev$Northing,
                        mean.psi = apply(out.sp.pred$psi.0.samples, 2, mean),
                       sd.psi = apply(out.sp.pred$psi.0.samples, 2, sd))
dat.stars <- st_as_stars(plot.dat, dims = c('x', 'y'))</pre>
ggplot() +
  geom_stars(data = dat.stars, aes(x = x, y = y, fill = mean.psi)) +
  scale_fill_distiller(palette = 'Blues', direction = 1, na.value = 'transparent') +
  labs(x = 'Easting', y = 'Northing', fill = '',
       title = 'Mean OVEN occurrence probability') +
  theme bw()
ggplot() +
  geom_stars(data = dat.stars, aes(x = x, y = y, fill = sd.psi)) +
  scale_fill_distiller(palette = 'Blues', direction = 1, na.value = 'transparent') +
  labs(x = 'Easting', y = 'Northing', fill = '',
       title = 'SD OVEN occurrence probability') +
  theme_bw()
```

4 Multispecies occupancy models

4.1 Basic model description

Let $z_{i,j}$ be the true presence (1) or absence (0) of a species i at site j, with j = 1, ..., J and i = 1, ..., N. We assume the latent occurrence process arises from a Bernoulli process following

$$z_{i,j} \sim \text{Bernoulli}(\psi_{i,j}),$$

 $\text{logit}(\psi_{i,j}) = \mathbf{x}'_{i} \cdot \boldsymbol{\beta}_{i},$ (5)

where $\psi_{i,j}$ is the probability of occurrence of species i at site j, which is a function of site-specific covariates X and a vector of species-specific regression coefficients (β_i). The regression coefficients in multispecies occupancy models are envisioned as random effects arising from a common community level distribution:

$$\beta_i \sim \text{Normal}(\mu_\beta, T_\beta),$$
 (6)

where μ_{β} is a vector of community level mean effects for each occurrence covariate effect (including the intercept) and T_{β} is a diagonal matrix with diagonal elements τ_{β}^2 that represent the variability of each occurrence covariate effect among species in the community.

We do not directly observe $z_{i,j}$ and rather we observe an imperfect representation of the latent occurrence process. Let $y_{i,j,k}$ be the observed detection (1) or nondetection (0) of a species i at site j during replicate k for each of $k = 1, \ldots, K_j$ replicates at each site j. We envision the detection-nondetection data as arising from a Bernoulli process conditional on the true latent occurrence process:

$$y_{i,j,k} \sim \operatorname{Bernoulli}(p_{i,j,k} \cdot z_{i,j}),$$

 $\operatorname{logit}(p_{i,j,k}) = \mathbf{v}'_{i,j,k} \cdot \mathbf{\alpha}_i,$ (7)

where $p_{i,j,k}$ is the probability of detecting species i at site j during replicate k (given it is present at site j), which is a function of site and replicate specific covariates V and a vector of species-specific regression coefficients (α_i). Similarly to the occurrence regression coefficients, the species specific detection coefficients are envisioned as random effects arising from a common community level distribution:

$$\alpha_i \sim \text{Normal}(\mu_\alpha, T_\alpha),$$
 (8)

where μ_{α} is a vector of community level mean effects for each detection covariate effect (including the intercept) and T_{α} is a diagonal matrix with diagonal elements τ_{α}^2 that represent the variability of each detection covariate effect among species in the community.

To complete the Bayesian specification of the model, we assign multivariate normal priors for the occurrence (μ_{β}) and detection (μ_{α}) community-level regression coefficient means and independent inverse-Gamma priors for each element of τ_{β}^2 and τ_{α}^2 . We again use Polya-Gamma data augmentation to yield an efficient implementation of the multispecies occupancy model, which is described in depth in the MCMC sampler vignette.

4.2 Fitting multispecies occupancy models with msPGOcc

spOccupancy uses nearly identical syntax for fitting multispecies models as it does for single species models and provides the same functionality for posterior predictive checks, GoF assessments using WAIC, and out of

sample prediction. The msPGOcc function fits nonspatial multispecies occupancy models using Polya-Gamma latent variables, which results in substantial increases in run time compared to standard implementations of logit link multispecies occupancy models. msPGOcc has exactly the same arguments as PGOcc:

We will again use the Hubbard Brook data in hbef2015 as an example data set, but we will now model occurrence for all 12 species in the community. Below we reload the hbef2015 data set to get a fresh copy.

```
data(hbef2015)
```

We will model occurrence for all species as a function of linear and quadratic elevation, and detection as a function of linear and quadratic day of survey as well as the time of day the survey occurred. These models are specified in occ.formula and det.formula as before, which reference variables stored in the data list. Random intercepts can be included in both the occurrence and detection portions of the occupancy model using lme4 syntax (Bates et al. 2015). For multispecies models, the multispecies detection-nondetection data y is now a three-dimensional array with dimensions corresponding to species, sites, and replicates. This is how the data are provided in the hbef2015 object, so we don't need to do any additional prep.

```
occ.ms.formula <- ~ Elevation + Elevation.2
det.ms.formula <- ~ day + tod + day.2
str(hbef2015)</pre>
```

```
List of 4
 $ y
           : num [1:12, 1:373, 1:3] 0 0 0 1 0 1 1 0 0 0 ...
  ..- attr(*, "dimnames")=List of 3
  ....$ : chr [1:12] "AMRE" "BAWW" "BHVI" "BLBW" ...
  ....$ : chr [1:373] "1" "2" "3" "4" ...
  .. ..$ : chr [1:3] "1" "2" "3"
 $ occ.covs: num [1:373, 1:2] -0.889 -0.765 -0.413 -0.14 -0.13 ...
  ..- attr(*, "dimnames")=List of 2
  .. ..$ : NULL
  ....$ : chr [1:2] "Elevation" "Elevation.2"
 $ det.covs:List of 3
  ..$ day : num [1:373, 1:3] -1.62 -1.62 -1.62 -1.62 -1.62 ...
  ..$ tod : num [1:373, 1:3] -1.565 -1.378 -1.084 -0.79 -0.549 ...
  ..$ day.2: num [1:373, 1:3] 2.61 2.61 2.61 2.61 2.61 ...
 $ coords : num [1:373, 1:2] 280000 280000 280000 280001 280000 ...
  ..- attr(*, "dimnames")=List of 2
  ....$ : chr [1:373] "1" "2" "3" "4" ...
  ....$ : chr [1:2] "Easting" "Northing"
```

Next we specify the starting values in starting. For multispecies occupancy models, we need to supply starting values for community-level and species-level parameters. In msPGOcc, we will supply starting values for the following parameters: alpha.comm (community level detection coefficients), beta.comm (community level occurrence coefficients), tau.sq.beta (species level occurrence variance parameters), tau.sq.alpha (community level detection variance parameters, z (latent occurrence values for all species). These are all specified in a single list. Starting values for community level parameters are vectors of length corresponding to the number of community-level detection or occurrence parameters in the model (including the intercepts), while starting values for species level parameters are matrices with the number of rows indicating the number of species, and each column corresponding to a different regression parameter. The starting values for the latent occurrence matrix are specified as a matrix with N rows corresponding to the number of species and J columns corresponding to the number of sites.

In multispecies models, we specify priors on the community-level coefficients rather than the species-level effects. For nonspatial models, these priors are specified with the following tags: beta.comm.normal (normal prior on the community level occurrence mean effects), alpha.comm.normal (normal prior on the community level detection mean effects), tau.sq.beta.ig (inverse-Gamma prior on the community level occurrence variance parameters), tau.sq.alpha.ig (inverse-Gamma prior on the community level detection variance parameters). Each tag consists of a list with elements corresponding to the mean and variance for normal priors and scale and shape for inverse-Gamma priors.

Below we specify normal priors to be relatively non-informative on the probability scale with a mean of 0 and variance of 2.72, and specify vague inverse gamma priors on the community level variance parameters setting both the shape and scale parameters to 0.1.

All that's left to do is specify the number of threads to use (n.omp.threads), the number of MCMC samples (n.samples), the amount of samples to discard as burn-in (n.burn), the thinning rate (n.thin), and arguments to control the display of sampler progress (verbose, n.report).

```
Preparing the data

------

Model description

Multi-species Occupancy Model with Polya-Gamma
```

Multi-species Occupancy Model with Polya-Gamma latent variable fit with 373 sites and 12 species.

Number of MCMC samples: 20000

Burn-in: 10000 Thinning Rate: 10

Total Posterior Samples: 1000

Source compiled with OpenMP support and model fit using 1 thread(s).

Sampling ...

Sampled: 5000 of 20000, 25.00%

Sampled: 10000 of 20000, 50.00%

Sampled: 15000 of 20000, 75.00%

Sampled: 20000 of 20000, 100.00%

out.ms\$run.time

```
user system elapsed 183.526 0.154 183.673
```

We see msPGOcc took less than 3 minutes to run the multispecies occupancy model with 373 sites and 12 species for a total of 20,000 iterations. The resulting object out.ms is a list of class msPGOcc consisting primarily of posterior samples of all community and species level parameters, as well as some additional objects that are used for summaries, prediction, and model fit evaluation. We can display a nice summary of these results using the summary function. For multispecies objects, when using summary we need to specify the level of parameters we want to summarize. We do this using the argument level, which takes values community, species, or both to print results for community-level parameters, species-level parameters, or all parameters. level is the second argument, so we can also avoid typing it out explicitly every time we want to call it

```
summary(out.ms, level = 'both')
```

```
Call:
```

```
msPGOcc(occ.formula = occ.ms.formula, det.formula = det.ms.formula,
  data = hbef2015, starting = ms.starting, priors = ms.priors,
  n.samples = 20000, n.omp.threads = 1, verbose = TRUE, n.report = 5000,
  n.burn = 10000, n.thin = 10)
```

Chain Information:

Total samples: 20000

Burn-in: 10000

Thin: 10

Total Posterior Samples: 1000

```
Community Level
```

Occurrence Means:

```
2.5% 25% 50% 75% 97.5% (Intercept) -1.5883 -0.2082 0.3206 0.8806 1.9983 Elevation -1.0832 -0.3174 0.0175 0.3377 1.0113 Elevation.2 -1.0032 -0.4738 -0.2413 0.0028 0.6059
```

Occurrence Variances:

2.5% 25% 50% 75% 97.5% (Intercept) 4.5074 7.8027 10.8072 15.3853 31.8971 Elevation 1.1178 1.9706 2.7677 3.8940 8.3826 Elevation.2 0.4290 0.8374 1.2677 2.0092 5.9462

Detection Means:

2.5% 25% 50% 75% 97.5% (Intercept) -1.3519 -0.7236 -0.4602 -0.1608 0.3992 day -0.2985 -0.1675 -0.1092 -0.0510 0.0567 tod -0.2991 -0.1649 -0.1170 -0.0668 0.0226 day.2 -0.2570 -0.1467 -0.0962 -0.0495 0.0453

Detection Variances:

2.5% 25% 50% 75% 97.5% (Intercept) 0.7079 1.4090 1.9250 2.7289 5.5469 day 0.0256 0.0474 0.0658 0.0956 0.1994 tod 0.0181 0.0307 0.0439 0.0630 0.1420 day.2 0.0163 0.0300 0.0419 0.0616 0.1306

Species Level

Occurrence:

2.5% 25% 50% 75% 97.5% (Intercept)-AMRE -3.6705 -3.0544 -2.7023 -2.3786 -1.5224 (Intercept)-BAWW -1.0476 -0.1848 0.5865 1.6507 4.5329 (Intercept)-BHVI -0.7315 -0.1103 0.4062 1.3361 7.0292 (Intercept)-BLBW 2.5210 3.1313 3.5885 4.2179 5.9324 (Intercept)-BLPW -5.6866 -4.9212 -4.5147 -4.1461 -3.5110 (Intercept)-BTBW 3.6618 4.2728 4.6392 5.1618 6.3808 (Intercept)-BTNW 2.2191 2.8108 3.2468 4.2057 7.7523 (Intercept)-CAWA -1.8961 -1.4187 -1.1695 -0.8438 -0.0439 (Intercept)-MAWA -2.3572 -2.0647 -1.9074 -1.7539 -1.4761 (Intercept)-NAWA -5.3886 -4.1210 -3.4722 -2.7425 -0.8003 (Intercept)-OVEN 1.9925 2.3938 2.6217 2.8947 3.5793 (Intercept)-REVI 1.9386 2.2991 2.5409 2.7784 3.3804 Elevation-AMRE -3.4472 -2.2537 -1.7465 -1.2497 -0.5959 Elevation-BAWW -1.0811 -0.4945 -0.1637 0.1385 0.8521 Elevation-BHVI -1.4595 -0.4341 -0.0394 0.4771 2.1548 Elevation-BLBW -3.0485 -1.3710 -0.9715 -0.7603 -0.5111 1.5779 2.0313 2.3386 2.7157 3.6131 Elevation-BLPW -1.4440 -1.1214 -0.9881 -0.8446 -0.6267 Elevation-BTBW Elevation-BTNW -1.9934 -0.3195 0.5145 1.4879 3.2336 Elevation-CAWA 0.7959 1.3082 1.6299 2.0165 3.0128 1.5277 1.7719 1.9171 2.0831 2.4068 Elevation-MAWA Elevation-NAWA -0.1089 0.4370 0.7765 1.1705 2.2525 Elevation-OVEN -1.8144 -1.3754 -1.2173 -1.0841 -0.8687 Elevation-REVI -3.4426 -2.3516 -1.9570 -1.7023 -1.2899 Elevation.2-AMRE -2.2299 -1.4790 -1.0942 -0.7881 -0.2897 Elevation.2-BAWW -3.5712 -1.9982 -1.4928 -1.1028 -0.5939 Elevation.2-BHVI -0.4353 0.5453 1.1962 2.1129 4.8100 Elevation.2-BLBW -1.1322 -0.8424 -0.6806 -0.5373 -0.0256 Elevation.2-BLPW 0.0413 0.5590 0.7855 0.9801 1.3690

```
Elevation.2-BTBW -1.7909 -1.4237 -1.2749 -1.1193 -0.9131
Elevation.2-BTNW -1.1704 -0.3409 0.1520 0.6966 1.6915
Elevation.2-CAWA -2.1291 -1.4646 -1.1813 -0.9070 -0.4203
Elevation.2-MAWA 0.1422 0.4243 0.5625
                                        0.6936 0.9287
Elevation.2-NAWA 0.3712 0.7893 1.0972
                                        1.4545
                                                2.8903
Elevation.2-OVEN -1.1517 -0.8851 -0.7694 -0.6600 -0.3837
Elevation.2-REVI -0.7361 -0.4572 -0.2884 -0.0905 0.4637
Detection:
                   2.5%
                            25%
                                    50%
                                           75%
                                                 97.5%
(Intercept)-AMRE -2.1003 -1.2942 -0.8898 -0.5416 0.0952
(Intercept)-BAWW -3.2920 -2.7335 -2.3900 -1.9927 -1.1927
(Intercept)-BHVI -2.4948 -2.2079 -2.0369 -1.8812 -1.4776
(Intercept)-BLBW 0.0066 0.1457 0.2222 0.2873 0.4235
(Intercept)-BLPW -0.2996 0.0265 0.2037 0.3777 0.7031
(Intercept)-BTBW
                 0.6541
                         0.7862
                                0.8623
                                        0.9427
                                                1.0891
(Intercept)-BTNW 0.1226 0.2662 0.3532
                                       0.4222 0.5862
(Intercept)-CAWA -2.3256 -1.5345 -1.2355 -0.9747 -0.4967
(Intercept)-MAWA -0.0703 0.1705 0.3006 0.4428 0.6886
(Intercept)-NAWA -4.2937 -3.3436 -2.8068 -2.2510 -1.1097
(Intercept)-OVEN 0.3590 0.5271 0.6114 0.6882 0.8678
(Intercept)-REVI 0.4095 0.5756 0.6584 0.7488 0.9024
day-AMRE
                -0.8008 -0.4448 -0.3093 -0.1745 0.0663
day-BAWW
                -0.4589 -0.2593 -0.1528 -0.0554
                                                0.1454
                 0.0361 0.1749 0.2461 0.3153 0.4514
day-BHVI
day-BLBW
                -0.2373 -0.1557 -0.1116 -0.0750 -0.0008
day-BLPW
                -0.4669 -0.2799 -0.1737 -0.0747 0.1308
day-BTBW
                -0.0584 0.0222 0.0689 0.1139 0.2001
                -0.0291 0.0474 0.0863 0.1298 0.2069
day-BTNW
day-CAWA
                -0.5754 -0.3736 -0.2709 -0.1763 0.0037
day-MAWA
                -0.4635 -0.3253 -0.2534 -0.1786 -0.0351
day-NAWA
                -0.8219 -0.4260 -0.2723 -0.1167
                                                0.1187
day-OVEN
                -0.1557 -0.0707 -0.0307 0.0134 0.0958
day-REVI
                -0.3063 -0.2078 -0.1659 -0.1264 -0.0440
tod-AMRE
                -0.7179 -0.4037 -0.2625 -0.1419 0.0837
tod-BAWW
                -0.4347 -0.2213 -0.1170 -0.0232 0.1737
tod-BHVI
                tod-BLBW
                -0.1900 -0.1114 -0.0760 -0.0306 0.0454
tod-BLPW
                -0.5648 -0.3675 -0.2530 -0.1601
                                                0.0316
                -0.1878 -0.0978 -0.0538 -0.0058 0.0839
tod-BTBW
                -0.0634 0.0066 0.0486 0.0901 0.1647
tod-BTNW
tod-CAWA
                -0.4858 -0.2690 -0.1668 -0.0608 0.1126
                -0.4477 -0.2955 -0.2205 -0.1474 -0.0157
tod-MAWA
tod-NAWA
                -0.4974 -0.2367 -0.1132 0.0044 0.2725
                -0.2583 -0.1756 -0.1265 -0.0852 0.0011
tod-OVEN
                -0.1852 -0.1052 -0.0604 -0.0102
                                                0.0769
tod-REVI
day.2-AMRE
                -0.6218 -0.3291 -0.2037 -0.0859
                                                0.1431
                -0.5107 -0.3158 -0.2095 -0.1055
day.2-BAWW
                                                0.0791
day.2-BHVI
                -0.3720 -0.2212 -0.1388 -0.0675
                                                0.0593
day.2-BLBW
                -0.1006 -0.0220 0.0217
                                        0.0686
                                                0.1627
                -0.4219 -0.1937 -0.0970 0.0003 0.2214
day.2-BLPW
day.2-BTBW
                -0.1173 -0.0170 0.0359 0.0836 0.1713
day.2-BTNW
                -0.1895 -0.0986 -0.0557 -0.0097 0.0932
```

day.2-CAWA

-0.4760 -0.2578 -0.1573 -0.0432 0.1593

```
day.2-MAWA -0.2661 -0.1239 -0.0371 0.0403 0.2014
day.2-NAWA -0.5889 -0.2977 -0.1570 -0.0339 0.2160
day.2-OVEN -0.3025 -0.1969 -0.1506 -0.1007 0.0011
day.2-REVI -0.2063 -0.0934 -0.0418 0.0096 0.1043

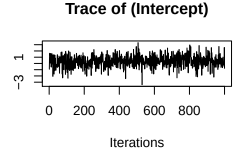
# Or
# summary(out.ms, 'both')
```

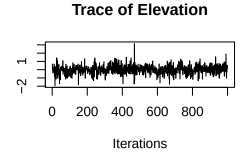
Looking at the community level variance parameters, we see large variability in the average occurrence (the intercept) for the twelve species, as well as substantial variability in the effect of elevation across the community. There appears to be less variability across species in the detection portion of the model. We can look directly at the species-specific effects to confirm this.

4.3 Convergence diagnostics

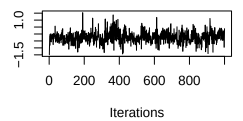
The resulting posterior samples in the msPGOcc object are coda::mcmc samples, and so convergence diagnostics can proceed as we saw with single species models.

```
plot(out.ms$beta.comm.samples, density = FALSE)
```



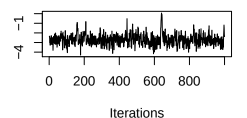


Trace of Elevation.2

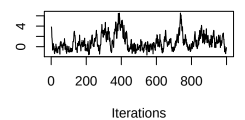


```
# Look at the first few species-specific occurrence intercepts
plot(out.ms$beta.samples[, 1:4], density = FALSE)
```

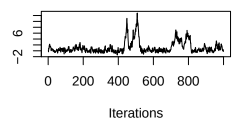
Trace of (Intercept)-AMRE



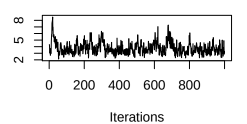
Trace of (Intercept)-BAWW



Trace of (Intercept)-BHVI



Trace of (Intercept)-BLBW



Looking at the species-specific intercepts, we would probably want to run the model a bit longer. Formal assessments of convergence using the Gelman-Rubin diagnostic can be accomplished following the steps shown for PGOcc using the gelman.diag function.

4.4 Posterior predictive checks

We can use the ppcOcc function to perform a posterior predictive check, and summarize the check with a Bayesian p-value using the summary function. The summary function again requires the level argument to specify if you want an overall Bayesian p-value for the entire community (level = 'community'), each individual species (level = 'species'), or both (level = 'both').

```
ppc.ms.out <- ppcOcc(out.ms, 'chi-square', group = 1)</pre>
[1] "Currently on species 1 out of 12"
[1] "Currently on species 2 out of 12"
[1] "Currently on species 3 out of 12"
[1] "Currently on species 4 out of 12"
[1] "Currently on species 5 out of 12"
[1] "Currently on species 6 out of 12"
[1] "Currently on species 7 out of 12"
[1] "Currently on species 8 out of 12"
[1] "Currently on species 9 out of 12"
[1] "Currently on species 10 out of 12"
[1] "Currently on species 11 out of 12"
[1] "Currently on species 12 out of 12"
summary(ppc.ms.out, level = 'both')
Call:
ppcOcc(object = out.ms, fit.stat = "chi-square", group = 1)
```

```
Chain Information:
Total samples: 20000
Burn-in: 10000
Thin: 10
Total Posterior Samples: 1000
   Community Level
Bayesian p-value: 0.3059167
    Species Level
AMRE Bayesian p-value: 0.323
BAWW Bayesian p-value: 0.561
BHVI Bayesian p-value: 0.392
BLBW Bayesian p-value: 0.107
BLPW Bayesian p-value: 0.325
BTBW Bayesian p-value: 0.099
BTNW Bayesian p-value: 0.109
CAWA Bayesian p-value: 0.45
MAWA Bayesian p-value: 0.397
NAWA Bayesian p-value: 0.676
OVEN Bayesian p-value: 0.219
REVI Bayesian p-value: 0.013
Fit statistic: chi-square
```

The Bayesian p-value for the overall community suggests an adequate model fit, but looking closer at each individual species reveals certain species the model may not be fitting well for all species. We should explore this further in a complete analysis (and also of course run the model longer to ensure convergence, as this is likely contributing to many of the extreme values).

4.5 Model selection using WAIC

We can compute the WAIC for comparison with alternative models using the waicOCC function.

```
waicOcc(out.ms)
```

```
elpd pD WAIC -4638.54370 72.18475 9421.45690
```

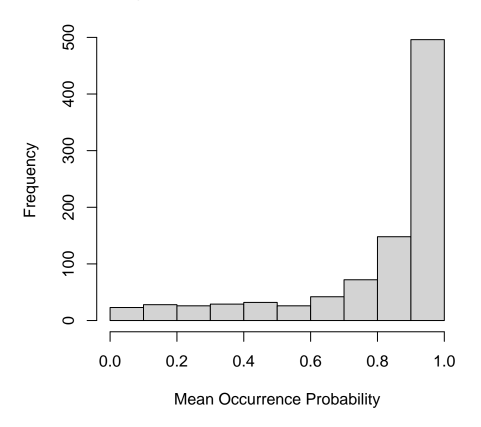
4.6 Prediction

Out-of-sample prediction with msPGOcc objects is exactly analogous to what we saw with PGOcc. We can use the predict function along with a data frame of covariates at new locations. We predict across the entire HBEF for all twelve species using the elevation data stored in hbefElev. Instead of predicting for every 30 x 30 m cell across the HBEF as we did with the single species model, below we predict values at every 50th cell. We predict only for OVEN for comparison to predictions using single species occupancy models.

```
X.0.ms <- cbind(1, hbefElev$val, hbefElev$val^2)
X.0.ms <- X.0.ms[seq(1, nrow(X.0.ms), by = 50), ]
out.ms.pred <- predict(out.ms, X.0.ms)</pre>
```

```
psi.pred.oven <- out.ms.pred$psi.0.samples[, which(sp.names == 'OVEN'), ]
hist(apply(psi.pred.oven, 2, mean), xlab = 'Mean Occurrence Probability')</pre>
```

Histogram of apply(psi.pred.oven, 2, mean)



The histogram supports our results from the single species model that indicated OVEN was common throughout most of the HBEF.

5 Multispecies spatial occupancy models

5.1 Basic model description

Residual spatial autocorrelation may perhaps be more prominent in multispecies occupancy models compared to single species models, as a single set of covariates is used to explain occurrence probability across a region of interest for all species. Given the large variety individual species show in habitat requirements, this may result in important drivers of occurrence probability not being included for certain species, resulting in many species having high residual spatial autocorrelation. We extend the previous multispecies occupancy model to incorporate a distinct spatial Gaussian Process (GP) for each species that accounts for unexplained spatial variation in each individual species occurrence across a spatial region. Occurrence probability for species i at site j, $\psi_{i,j}$, now takes the form

$$logit(\psi_{i,j}) = \mathbf{x}_{i}'\boldsymbol{\beta}_{i} + w_{i,j}, \tag{9}$$

where the species-specific regression coefficients β_i follow the community level distribution in Equation (6), and $\mathbf{w}_{i,j}$ is a realization from a zero-mean spatial GP, i.e.,

$$\mathbf{w}_i \sim \text{Normal}(\mathbf{0}, \mathbf{\Sigma}_i(\mathbf{s}, \mathbf{s}', \boldsymbol{\theta}_i)).$$
 (10)

We define $\Sigma_i(s, s', \theta_i)$ as a $J \times J$ covariance matrix that is a function of the distances between any pair of site coordinates s and s' and a set of parameters (θ_i) that govern the spatial process. The vector θ_i is equal to $\theta_i = {\sigma_i^2, \phi_i, \nu_i}$, where σ_i^2 is a spatial variance parameter for species i, ϕ_i is a spatial decay parameter for species i, and ν_i is a spatial smoothness parameter for species i. ν_i is only specified when using a Matern correlation function.

The detection portion of the multispecies spatial occupancy model remains unchanged from the non-spatial multispecies occupancy model and follows Equations (7) and (8). We fit the model again using Polya-Gamma data augmentation to enable an efficient Gibbs sampler (see MCMC sampler vignette for details). Similar to our discussion on the single species spatial occupancy model, we also allow for specification of the spatial process using an NNGP instead fo a full GP. This leads to even larger computational gains over the full GP given that a separate covariance matrix is specified for each species in the model. See Datta et al. (2016), Finley, Datta, and Banerjee (2020), and the MCMC sampler vignette for additional details on NNGPs and their implementation in multispecies spatial occupancy models.

5.2 Fitting multispecies spatial occupancy models with spMsPGOcc

The function spMsPGOcc fits spatially explicit multispecies occupancy models. Similar to single species models using spPGOcc, models can be fit using either a full Gaussian Process (GP) or a Nearest Neighbor Gaussian Process (NNGP). spMsPGOcc fits a separate spatial process for each species. The syntax for spMsPGOcc is analogous to the syntax for single species spatially-explicit models using spPGOcc.

```
spMsPGOcc(occ.formula, det.formula, data, starting, n.batch,
    batch.length, accept.rate = 0.43, priors,
    cov.model = "exponential", tuning, n.omp.threads = 1,
    verbose = TRUE, NNGP = TRUE, n.neighbors = 15,
    search.type = "cb", n.report = 100,
    n.burn = round(.10 * n.batch * batch.length), n.thin = 1, ...)
```

We will again display the model using the HBEF foliage-gleaning bird data set, with the same predictors in our occurrence and detection models

```
occ.ms.sp.formula <- ~ Elevation + Elevation.2
det.ms.sp.formula <- ~ day + tod + day.2
```

Our starting values in the starting argument will look analogous to what we specified for the nonspatial multispecies occupancy model using msPGOcc, but we will also include additional starting values for the parameters controlling the spatial processes: sigma.sq is the species-specific spatial variance parameter, phi is the species specific spatial decay parameter, and w is the latent spatial process for each species at each site. We will use an exponential covariance model, but when using a Matern covariance model we must also specify starting values for nu, the species-specific spatial smoothness parameter. Note that all species-specific spatial parameters are independent of each other. We currently do not leverage any correlation between spatial processes of different species, although this is something we plan to incorporate for future spOccupancy development. Starting values for phi, sigma.sq, and nu (if applicable) are specified as vectors with N elements (the number of species being modeled), while the starting values for the latent spatial processes are specified as a matrix with N rows (i.e., species) and J columns (i.e., sites). Here we set the starting value for the spatial variances equal to 2 for all species and set the starting values for the spatial decay parameter to yield an effective range of the average distance between sites across the HBEF.

```
# Number of species
N \leftarrow dim(hbef2015\$y)[1]
# Distances between sites
dist.hbef <- dist(hbef2015$coords)</pre>
# Number of detection and occupancy regression parameters
p.det <- length(hbef2015$det.covs) + 1</pre>
p.occ <- ncol(hbef2015$occ.covs) + 1</pre>
# Exponential covariance model
cov.model <- "exponential"</pre>
ms.starting <- list(alpha.comm = rep(0, p.det),
                     beta.comm = rep(0, p.occ),
                     beta = matrix(0, N, p.occ),
                     alpha = matrix(0, N, p.det),
                     tau.sq.beta = rep(1, p.occ),
                     tau.sq.alpha = rep(1, p.det),
                     z = apply(hbef2015\$y, c(1, 2), max, na.rm = TRUE),
                     sigma.sq = rep(2, N),
                     phi = rep(3 / mean(dist.hbef), N),
                     w = matrix(0, N, dim(hbef2015\$y)[2]))
```

We next specify the priors in the priors argument. The priors are the same as those we specified for the non-spatial multispecies model, with the addition of priors for the parameters controlling the species-specific spatial processes. We assume independent priors for all spatial parameters across the different species. For each species, we assign an inverse gamma prior for the spatial varaince parameter sigma.sq (tag is sigma.sq.ig) and uniform priors for the spatial decay parameter phi and smoothness parameter nu (if cov.model = 'matern'), with the associated tags phi.unif and nu.unif. All priors are specified as lists with two elements. For the inverse-Gamma prior, the first element is a length N vector of shape parameters for each species, and the second element is a length N vector of scale parameters for each species. For the uniform priors, the first element is a length N vector of the lower bounds for each species, and the second element is a length N vector of upper bounds for each species. For the inverse-Gamma prior on the spatial variances, here we set the shape parameter to 2 and the scale parameter equal to 2. For a more formal analysis, we would likely want to do some exploratory data analysis to obtain a better guess for the spatial variance for each species, and then replace the scale parameter with this estimated guess for each species. For the spatial decay parameter, we determine the bounds of the uniform distribution by computing the smallest distance between sites and the largest distance between sites. We then set the lower bound of the uniform to 3/max and the upper bound to 3/min, where min and max correspond to the predetermined distances between sites.

We next set the parameters controlling the Adaptive MCMC algorithm (see spPGOcc section for details).

Notice our specification of the starting tuning values is exactly the same as for spPGOcc. We assume the same initial tuning value for all species. However, the adaptive algorithm will allow for species specific tuning parameters, so these will be adjusted in the algorithm as needed (and reported to the R console if verbose = TRUE).

```
batch.length <- 25
n.batch <- 400
n.burn <- 2000
n.thin <- 8
ms.tuning <- list(phi = 0.5)
n.omp.threads <- 1
# Values for reporting
verbose <- TRUE
n.report <- 50</pre>
```

Spatially explicit multispecies occupancy models are currently the most computationally intensive models fit by spOccupancy. Even for modest sized data sets, we encourage the use of NNGPs instead of full GPs when fitting spatially-explicit models to ease the computational burden of fitting these models. We fit the model with an NNGP below using 5 neighbors and summarize it using the summary function, where we specify that we want to summarize both species and community level parameters.

```
out.sp.ms <- spMsPGOcc(occ.formula = occ.ms.sp.formula,</pre>
               det.formula = det.ms.sp.formula,
               data = hbef2015,
               starting = ms.starting,
               n.batch = n.batch,
               batch.length = batch.length,
               accept.rate = 0.43,
               priors = ms.priors,
               cov.model = cov.model,
               tuning = ms.tuning,
               n.omp.threads = n.omp.threads,
               verbose = TRUE,
               NNGP = TRUE,
               n.neighbors = 5,
               n.report = n.report,
               n.burn = n.burn,
               n.thin = n.thin)
```

```
Preparing the data

Building the neighbor list

Building the neighbors of neighbors list

Model description

NNGP Multi-species Occupancy Model with Polya-Gamma latent variable fit with 373 sites and 12 species.

Number of MCMC samples 10000 (400 batches of length 25)
```

Burn-in: 2000

```
Thinning Rate: 8
```

Total Posterior Samples: 1000

Using the exponential spatial correlation model.

Using 5 nearest neighbors.

Source compiled with OpenMP support and model fit using 1 thread(s).

Adaptive Metropolis with target acceptance rate: 43.0

Sampling ...

Batch: 50 of 400, 12.50%

Batch: 50 of 4	±00, 12.50%	
parameter	acceptance	tuning
phi[0]	40.0	0.39727
phi[1]	72.0	0.65498
phi[2]	64.0	0.36672
phi[3]	80.0	0.64201
phi[4]	32.0	0.45697
phi[5]	80.0	0.50503
phi[6]	64.0	0.53625
phi[7]	48.0	0.41348
phi[8]	52.0	0.35946
phi[9]	32.0	0.37413
phi[10]	60.0	0.40529
phi[11]	28.0	0.36672
Batch: 100 of	400, 25.00%	
parameter	acceptance	tuning
phi[0]	40.0	0.38169
phi[1]	68.0	0.84947
phi[2]	80.0	0.48522

parameter	acceptance	cuning
phi[0]	40.0	0.38169
phi[1]	68.0	0.84947
phi[2]	80.0	0.48522
phi[3]	20.0	0.68171
phi[4]	20.0	0.38940
phi[5]	32.0	0.45697
phi[6]	52.0	0.84947
phi[7]	36.0	0.35946
phi[8]	44.0	0.38169
phi[9]	80.0	0.43905
phi[10]	24.0	0.49502
phi[11]	24.0	0.31881

atch: 150 of	400, 37.50%	
parameter	acceptance	tuning
phi[0]	36.0	0.31881
phi[1]	84.0	1.01700
phi[2]	28.0	0.70953
phi[3]	52.0	0.81616
phi[4]	40.0	0.37413
phi[5]	48.0	0.39727
phi[6]	56.0	1.03754
phi[7]	32.0	0.40529
phi[8]	28.0	0.31881
phi[9]	48.0	0.61684
phi[10]	68.0	0.54709
phi[11]	48.0	0.31250

phi[11] 48.0 Batch: 200 of 400, 50.00%

parameter	acceptance	tuning
phi[0]	32.0	0.40529
phi[1]	76.0	0.81616
phi[2]	84.0	0.76863
phi[3]	68.0	0.83265
phi[4]	40.0	0.33853
phi[5]	40.0	0.34537
phi[6]	40.0	1.10170
phi[7]	36.0	0.37413
phi[8]	56.0	0.30025
phi[9]	28.0	0.51523
phi[10]	24.0	0.48522
phi[11]	44.0	0.31250
Batch: 250 of	400, 62.50%	
parameter	acceptance	tuning
phi[0]	68.0	0.40529
phi[1]	48.0	0.92022
phi[2]	28.0	1.05850
phi[3]	44.0	0.83265
phi[4]	48.0	0.35946
phi[5]	48.0	0.38940
phi[6]	80.0	1.07988
phi[7]	76.0	0.33183
phi[8]	44.0	0.37413
phi[9]	76.0	0.53625
phi[3]	52.0	0.37413
phi[10] phi[11]	32.0	0.28847
PHT [TI]	02.0	0.20041
Batch: 300 of	400 75 00%	
Batch: 300 of	•	tuning
parameter	acceptance	tuning 0 52564
parameter phi[0]	acceptance 68.0	0.52564
parameter phi[0] phi[1]	acceptance 68.0 32.0	0.52564 0.97712
<pre>parameter phi[0] phi[1] phi[2]</pre>	acceptance 68.0 32.0 8.0 0.8	0.52564 0.97712 66663
parameter phi[0] phi[1] phi[2] phi[3]	acceptance 68.0 32.0 8.0 0.8 48.0	0.52564 0.97712 66663 0.66821
parameter phi[0] phi[1] phi[2] phi[3] phi[4]	acceptance 68.0 32.0 8.0 0.8 48.0 44.0	0.52564 0.97712 66663 0.66821 0.32525
parameter phi[0] phi[1] phi[2] phi[3] phi[4] phi[5]	acceptance 68.0 32.0 8.0 0.8 48.0 44.0 28.0	0.52564 0.97712 6663 0.66821 0.32525 0.53625
parameter phi[0] phi[1] phi[2] phi[3] phi[4] phi[5] phi[6]	acceptance 68.0 32.0 8.0 0.8 48.0 44.0 28.0 24.0	0.52564 0.97712 6663 0.66821 0.32525 0.53625 0.99686
parameter phi[0] phi[1] phi[2] phi[3] phi[4] phi[5] phi[6] phi[7]	acceptance 68.0 32.0 8.0 0.8 48.0 44.0 28.0 24.0	0.52564 0.97712 66663 0.66821 0.32525 0.53625 0.99686 0.47561
parameter phi[0] phi[1] phi[2] phi[3] phi[4] phi[5] phi[6] phi[7] phi[8]	acceptance 68.0 32.0 8.0 0.8 48.0 44.0 28.0 24.0 24.0 52.0	0.52564 0.97712 66663 0.66821 0.32525 0.53625 0.99686 0.47561 0.35234
parameter phi[0] phi[1] phi[2] phi[3] phi[4] phi[5] phi[6] phi[6] phi[7] phi[8] phi[9]	acceptance 68.0 32.0 8.0 0.8 48.0 44.0 28.0 24.0 24.0 52.0 88.0	0.52564 0.97712 66663 0.66821 0.32525 0.53625 0.99686 0.47561 0.35234 0.76863
parameter phi[0] phi[1] phi[2] phi[3] phi[4] phi[5] phi[6] phi[6] phi[7] phi[8] phi[9]	acceptance 68.0 32.0 8.0 48.0 44.0 28.0 24.0 24.0 52.0 88.0 36.0	0.52564 0.97712 66663 0.66821 0.32525 0.53625 0.99686 0.47561 0.35234 0.76863 0.34537
parameter phi[0] phi[1] phi[2] phi[3] phi[4] phi[5] phi[6] phi[6] phi[7] phi[8] phi[9] phi[10] phi[11]	acceptance 68.0 32.0 8.0 48.0 44.0 28.0 24.0 24.0 52.0 88.0 36.0 60.0	0.52564 0.97712 66663 0.66821 0.32525 0.53625 0.99686 0.47561 0.35234 0.76863
parameter phi[0] phi[1] phi[2] phi[3] phi[4] phi[5] phi[6] phi[7] phi[8] phi[9] phi[10] phi[11] Batch: 350 of	acceptance 68.0 32.0 8.0 48.0 44.0 28.0 24.0 52.0 88.0 36.0 60.0 400, 87.50%	0.52564 0.97712 66663 0.66821 0.32525 0.53625 0.99686 0.47561 0.35234 0.76863 0.34537 0.29430
parameter phi[0] phi[1] phi[2] phi[3] phi[4] phi[5] phi[6] phi[7] phi[8] phi[9] phi[10] phi[11] Batch: 350 of	acceptance 68.0 32.0 8.0 0.8 48.0 44.0 28.0 24.0 52.0 88.0 36.0 60.0 400, 87.50% acceptance	0.52564 0.97712 66663 0.66821 0.32525 0.53625 0.99686 0.47561 0.35234 0.76863 0.34537 0.29430 tuning
parameter phi[0] phi[1] phi[2] phi[3] phi[4] phi[5] phi[6] phi[7] phi[8] phi[9] phi[10] phi[11] Batch: 350 of parameter phi[0]	acceptance 68.0 32.0 8.0 0.8 48.0 44.0 28.0 24.0 52.0 88.0 36.0 60.0 400, 87.50% acceptance 44.0	0.52564 0.97712 66663 0.66821 0.32525 0.53625 0.99686 0.47561 0.35234 0.76863 0.34537 0.29430 tuning 0.43905
parameter phi[0] phi[1] phi[2] phi[3] phi[4] phi[5] phi[6] phi[7] phi[8] phi[9] phi[10] phi[11] Batch: 350 of parameter phi[0] phi[1]	acceptance 68.0 32.0 8.0 0.8 48.0 44.0 24.0 24.0 52.0 88.0 36.0 60.0 400, 87.50% acceptance 44.0 16.0	0.52564 0.97712 66663 0.66821 0.32525 0.53625 0.99686 0.47561 0.35234 0.76863 0.34537 0.29430 tuning 0.43905 0.81616
parameter phi[0] phi[1] phi[2] phi[3] phi[4] phi[5] phi[6] phi[7] phi[8] phi[9] phi[10] phi[11] Batch: 350 of parameter phi[0] phi[1] phi[2]	acceptance 68.0 32.0 8.0 0.8 48.0 44.0 28.0 24.0 24.0 52.0 88.0 36.0 60.0 400, 87.50% acceptance 44.0 16.0 40.0	0.52564 0.97712 66663 0.66821 0.32525 0.53625 0.99686 0.47561 0.35234 0.76863 0.34537 0.29430 tuning 0.43905 0.81616 0.78416
parameter phi[0] phi[1] phi[2] phi[3] phi[4] phi[5] phi[6] phi[7] phi[8] phi[9] phi[10] phi[11] Batch: 350 of parameter phi[0] phi[1] phi[2] phi[3]	acceptance 68.0 32.0 8.0 0.8 48.0 44.0 28.0 24.0 52.0 88.0 36.0 60.0 400, 87.50% acceptance 44.0 16.0 40.0 48.0	0.52564 0.97712 6663 0.66821 0.32525 0.53625 0.99686 0.47561 0.35234 0.76863 0.34537 0.29430 tuning 0.43905 0.81616 0.78416 0.62930
parameter phi[0] phi[1] phi[2] phi[3] phi[4] phi[5] phi[6] phi[7] phi[8] phi[9] phi[10] phi[11] Batch: 350 of parameter phi[0] phi[1] phi[2] phi[3] phi[4]	acceptance 68.0 32.0 8.0 0.8 48.0 44.0 28.0 24.0 52.0 88.0 36.0 60.0 400, 87.50% acceptance 44.0 16.0 40.0 48.0 48.0	0.52564 0.97712 6663 0.66821 0.32525 0.53625 0.99686 0.47561 0.35234 0.76863 0.34537 0.29430 tuning 0.43905 0.81616 0.78416 0.62930 0.30025
parameter phi[0] phi[1] phi[2] phi[3] phi[4] phi[5] phi[6] phi[7] phi[8] phi[9] phi[10] phi[11] Batch: 350 of parameter phi[0] phi[1] phi[2] phi[3] phi[4] phi[5]	acceptance 68.0 32.0 8.0 0.8 48.0 44.0 28.0 24.0 52.0 88.0 36.0 60.0 400, 87.50% acceptance 44.0 16.0 40.0 48.0 48.0 44.0	0.52564 0.97712 66663 0.66821 0.32525 0.53625 0.99686 0.47561 0.35234 0.76863 0.34537 0.29430 tuning 0.43905 0.81616 0.78416 0.62930 0.30025 0.43905
parameter phi[0] phi[1] phi[2] phi[3] phi[4] phi[5] phi[6] phi[7] phi[8] phi[9] phi[10] phi[11] Batch: 350 of parameter phi[0] phi[1] phi[2] phi[3] phi[4] phi[5] phi[6]	acceptance 68.0 32.0 8.0 0.8 48.0 44.0 28.0 24.0 52.0 88.0 36.0 60.0 400, 87.50% acceptance 44.0 16.0 40.0 48.0 48.0 44.0 16.0	0.52564 0.97712 6663 0.66821 0.32525 0.53625 0.99686 0.47561 0.35234 0.76863 0.34537 0.29430 tuning 0.43905 0.81616 0.78416 0.62930 0.30025 0.43905 1.03754
parameter phi[0] phi[1] phi[2] phi[3] phi[4] phi[5] phi[6] phi[7] phi[8] phi[9] phi[10] phi[11] Batch: 350 of parameter phi[0] phi[1] phi[2] phi[3] phi[4] phi[5] phi[6] phi[7]	acceptance 68.0 32.0 8.0 0.8 48.0 44.0 28.0 24.0 24.0 52.0 88.0 36.0 60.0 400, 87.50% acceptance 44.0 16.0 40.0 48.0 44.0 16.0 20.0	0.52564 0.97712 6663 0.66821 0.32525 0.53625 0.99686 0.47561 0.35234 0.76863 0.34537 0.29430 tuning 0.43905 0.81616 0.78416 0.62930 0.30025 0.43905 1.03754 0.38940
parameter phi[0] phi[1] phi[2] phi[3] phi[4] phi[5] phi[6] phi[7] phi[8] phi[9] phi[10] phi[11] Batch: 350 of parameter phi[0] phi[1] phi[2] phi[3] phi[4] phi[5] phi[6] phi[7] phi[8]	acceptance 68.0 32.0 8.0 0.8 48.0 44.0 28.0 24.0 24.0 52.0 88.0 36.0 60.0 400, 87.50% acceptance 44.0 16.0 40.0 48.0 44.0 16.0 20.0 52.0	0.52564 0.97712 66663 0.66821 0.32525 0.53625 0.99686 0.47561 0.35234 0.76863 0.34537 0.29430 tuning 0.43905 0.81616 0.78416 0.62930 0.30025 0.43905 1.03754 0.38940 0.31250
parameter phi[0] phi[1] phi[2] phi[3] phi[4] phi[5] phi[6] phi[7] phi[8] phi[9] phi[10] phi[11] Batch: 350 of parameter phi[0] phi[1] phi[2] phi[3] phi[4] phi[5] phi[6] phi[7]	acceptance 68.0 32.0 8.0 0.8 48.0 44.0 28.0 24.0 24.0 52.0 88.0 36.0 60.0 400, 87.50% acceptance 44.0 16.0 40.0 48.0 44.0 16.0 20.0	0.52564 0.97712 6663 0.66821 0.32525 0.53625 0.99686 0.47561 0.35234 0.76863 0.34537 0.29430 tuning 0.43905 0.81616 0.78416 0.62930 0.30025 0.43905 1.03754 0.38940

phi[11] 44.0 0.28847 Batch: 400 of 400, 100.00% summary(out.sp.ms, level = 'both') Call: spMsPGOcc(occ.formula = occ.ms.sp.formula, det.formula = det.ms.sp.formula, data = hbef2015, starting = ms.starting, priors = ms.priors, tuning = ms.tuning, cov.model = cov.model, NNGP = TRUE, n.neighbors = 5, n.batch = n.batch, batch.length = batch.length, accept.rate = 0.43, n.omp.threads = n.omp.threads, verbose = TRUE, n.report = n.report, n.burn = n.burn, n.thin = n.thin) Chain Information: Total samples: 10000 Burn-in: 2000 Thin: 8 Total Posterior Samples: 1000 Community Level Occurrence Means: 2.5% 25% 50% 75% 97.5% (Intercept) -1.8936 -0.6268 0.2256 0.9063 2.4591 Elevation -1.3568 -0.3698 0.0922 0.5554 1.5368 Elevation.2 -1.3757 -0.6148 -0.2402 0.1657 1.0368 Occurrence Variances: 2.5% 25% 50% 75% 97.5% (Intercept) 9.6901 17.6464 24.4021 33.0838 71.4737 Elevation 2.3249 4.4001 6.2989 9.0780 20.0931 Elevation.2 0.9753 2.0873 3.0545 4.7092 10.6003 Detection Means: 25% 50% 2.5% 75% 97.5% (Intercept) -1.2421 -0.6388 -0.3866 -0.1460 0.3318 -0.3118 -0.1773 -0.1157 -0.0610 0.0501 day -0.2838 -0.1728 -0.1185 -0.0720 0.0397 tod -0.2633 -0.1543 -0.1022 -0.0533 0.0467 dav.2 Detection Variances: 2.5% 25% 50% 75% 97.5% (Intercept) 0.6572 1.2363 1.7107 2.4942 5.6476 0.0265 0.0469 0.0667 0.0965 0.2150 0.0168 0.0314 0.0452 0.0634 0.1371 tod 0.0165 0.0297 0.0423 0.0598 0.1198 Species Level Occurrence: 2.5% 25% 50% 75% 97.5%

(Intercept)-AMRE -7.5341 -4.8335 -3.9663 -3.2496 -1.9354 (Intercept)-BAWW -1.6241 -0.2222 0.7847 2.4441 6.1424

```
(Intercept)-BHVI
                 -2.3836 -0.6380 -0.1069 0.4615 2.5974
                  3.0234 3.8141 4.4871 5.3877
(Intercept)-BLBW
                                                 8.0272
(Intercept)-BLPW -10.1521 -7.4784 -6.3515 -5.6558 -4.6875
(Intercept)-BTBW
                  4.2340 5.2069 5.9487
                                          6.7446 8.7794
(Intercept)-BTNW
                  2.8283 3.7540 4.8977
                                         7.8188 13.5993
(Intercept)-CAWA
                -4.1384 -2.5753 -1.9395 -1.4893 -0.1514
(Intercept)-MAWA
                 -8.4829 -5.5054 -4.3507 -3.4704 -1.0921
(Intercept)-NAWA
                 -7.2746 -5.5081 -4.5714 -3.6583 -1.7902
(Intercept)-OVEN
                  2.4538 2.9607 3.3607
                                          3.9089 5.9036
(Intercept)-REVI
                  2.6592 3.6782 4.6272 5.8671 8.2952
Elevation-AMRE
                 -4.0059 -2.6045 -1.9130 -1.2972 -0.2319
Elevation-BAWW
                 -1.6670 -0.6301 -0.2092 0.2588 1.2130
Elevation-BHVI
                 -2.0168 -0.6830 -0.1142 0.4187
                                                 1.9519
Elevation-BLBW
                 -3.8295 -1.6491 -1.1323 -0.8767 -0.5253
Elevation-BLPW
                  1.8741 2.8632 3.5032 4.2353 5.5870
Elevation-BTBW
                 -2.2433 -1.5604 -1.3083 -1.0958 -0.7315
                 -4.4135 -0.8930 0.9836 2.6784
Elevation-BTNW
                                                 4.9338
Elevation-CAWA
                  0.8424 1.7144 2.1893
                                         2.7588
                                                 3.8344
                  2.4591 3.3506 4.0089 4.8042 6.6808
Elevation-MAWA
Elevation-NAWA
                 -0.1207 0.4653 0.8383 1.3316
                                                 2.7605
Elevation-OVEN
                 -3.2187 -1.9384 -1.6172 -1.3709 -1.0418
Elevation-REVI
                 -5.1981 -3.5744 -2.9429 -2.4636 -1.7294
Elevation.2-AMRE -3.1741 -1.9739 -1.4570 -1.0380 -0.3160
                 -4.9748 -3.0020 -2.1751 -1.5625 -0.7909
Elevation.2-BAWW
Elevation.2-BHVI
                  0.1177 1.1252 2.0689 3.1742 6.0868
Elevation.2-BLBW
                -1.6170 -1.0867 -0.8723 -0.6768 -0.0040
Elevation.2-BLPW
                 -0.3258  0.6641  1.1072  1.5259  2.3327
Elevation.2-BTBW
                 -2.5905 -1.8943 -1.6211 -1.4194 -1.0879
Elevation.2-BTNW
                -1.7693 -0.4102 0.7390 1.5440 4.6694
Elevation.2-CAWA -3.4075 -2.1249 -1.6715 -1.3056 -0.6300
                  0.6070 1.1881 1.5764
Elevation.2-MAWA
                                          2.0163
                                                 3.2086
Elevation.2-NAWA
                  0.4627 0.9756 1.3432 1.7688
                                                 2.9452
Elevation.2-OVEN -1.7966 -1.2151 -1.0039 -0.8232 -0.4539
Elevation.2-REVI -1.8753 -1.0687 -0.7504 -0.4363 0.2572
```

Detection:

2.5% 25% 50% 75% 97.5% (Intercept)-AMRE -2.0383 -1.1635 -0.7852 -0.4459 0.2289 (Intercept)-BAWW -3.2351 -2.7199 -2.3770 -2.0068 -1.3221 (Intercept)-BHVI -2.3780 -2.0987 -1.9313 -1.7550 -1.1960 (Intercept)-BLBW 0.0026 0.1460 0.2197 0.2937 0.4361 (Intercept)-BLPW -0.3298 0.0544 0.2546 0.4340 0.7793 (Intercept)-BTBW 0.6389 0.7916 0.8636 0.9402 1.0754 (Intercept)-BTNW 0.1133 0.2406 0.3234 0.3992 0.5464 (Intercept)-CAWA -1.8901 -1.3927 -1.1167 -0.8650 -0.4394 (Intercept)-MAWA 0.0145 0.2342 0.3689 0.4875 0.6833 (Intercept)-NAWA -4.0310 -3.1931 -2.6705 -2.0211 -0.8679 (Intercept)-OVEN 0.3685 0.5222 0.6103 0.6912 0.8515 (Intercept)-REVI 0.4672 0.6189 0.7017 0.7840 0.9525 day-AMRE -0.8249 -0.4747 -0.3325 -0.1956 0.0652 day-BAWW -0.4609 -0.2471 -0.1487 -0.0490 0.1616 day-BHVI 0.0166 0.1702 0.2510 0.3231 0.4840 day-BLBW -0.2338 -0.1592 -0.1200 -0.0748 -0.0004 day-BLPW -0.4706 -0.2790 -0.1815 -0.0786 0.1125

```
day-BTBW
                 -0.0566 0.0222 0.0669 0.1142 0.1938
day-BTNW
                 -0.0293 0.0474 0.0933 0.1304 0.2040
day-CAWA
                 -0.6292 -0.4053 -0.3004 -0.1951 -0.0200
day-MAWA
                 -0.4534 -0.3245 -0.2562 -0.1826 -0.0653
day-NAWA
                 -0.7505 -0.4226 -0.2529 -0.1053 0.1879
day-OVEN
                 -0.1471 -0.0723 -0.0326 0.0122 0.0953
dav-REVI
                 -0.2926 -0.2126 -0.1675 -0.1248 -0.0417
tod-AMRE
                 -0.7498 -0.3879 -0.2657 -0.1474 0.0795
tod-BAWW
                 -0.4187 -0.2239 -0.1183 -0.0286
                                                 0.1737
tod-BHVI
                 -0.1758 -0.0564 0.0118 0.0707
                                                 0.2089
tod-BLBW
                 -0.1907 -0.1104 -0.0687 -0.0301
                                                 0.0454
                 -0.5662 -0.3605 -0.2634 -0.1602
tod-BLPW
                                                 0.0330
tod-BTBW
                 -0.1831 -0.0990 -0.0529 -0.0081
                                                 0.0819
tod-BTNW
                 -0.0700 0.0072 0.0436 0.0838 0.1650
                 -0.5194 -0.2859 -0.1675 -0.0629 0.1258
tod-CAWA
tod-MAWA
                 -0.4401 -0.2800 -0.2069 -0.1373 -0.0115
                 -0.5294 -0.2550 -0.1328 -0.0106 0.2436
tod-NAWA
tod-OVEN
                 -0.2636 -0.1807 -0.1342 -0.0847 -0.0039
                 -0.1873 -0.0976 -0.0507 -0.0065 0.0818
tod-REVI
day.2-AMRE
                 -0.6344 -0.3205 -0.2030 -0.0855
                                                0.1614
day.2-BAWW
                 -0.5504 -0.3156 -0.2017 -0.1005 0.0660
day.2-BHVI
                 -0.3637 -0.2183 -0.1424 -0.0630
day.2-BLBW
                 -0.1172 -0.0230 0.0256 0.0671
                                                 0.1538
day.2-BLPW
                 -0.4165 -0.2182 -0.1110 -0.0099
                                                 0.2125
                 -0.1055 -0.0162 0.0340 0.0826 0.1936
day.2-BTBW
                 -0.1890 -0.1018 -0.0532 -0.0074 0.0686
day.2-BTNW
day.2-CAWA
                 -0.4802 -0.2665 -0.1496 -0.0536
                                                 0.1558
                 -0.2723 -0.1235 -0.0519 0.0256
day.2-MAWA
                                                 0.1775
day.2-NAWA
                 -0.6085 -0.2866 -0.1510 -0.0229
                                                 0.2102
day.2-OVEN
                 -0.2981 -0.2010 -0.1509 -0.1050
                                                 0.0024
                 -0.1921 -0.0962 -0.0433 0.0039 0.0909
day.2-REVI
```

Covariance:

2.5% 25% 50% 75% 97.5% sigma.sq-AMRE 0.4491 2.0707 4.0096 6.8227 21.0515 sigma.sq-BAWW 0.4090 0.9258 1.7390 3.6436 27.9956 sigma.sq-BHVI 0.4314 1.1152 1.8177 3.4330 53.9690 sigma.sq-BLBW 0.3798 0.9177 1.4518 2.3688 7.7654 sigma.sq-BLPW 0.9954 2.3613 3.9039 6.8801 13.5029 sigma.sq-BTBW 0.5513 1.2168 2.1031 3.7292 12.0288 sigma.sq-BTNW 0.4181 0.8714 1.3969 2.3182 7.2021 sigma.sq-CAWA 0.6885 2.2422 3.9887 6.5925 27.4594 sigma.sq-MAWA 4.2975 8.9865 14.1951 20.5783 38.7941 1.2314 sigma.sq-NAWA 0.4431 0.8400 1.9489 6.5867 sigma.sq-OVEN 0.5338 1.0928 1.6036 2.7541 13.2737 sigma.sq-REVI 1.9314 5.0939 8.8422 15.1200 29.1024 phi-AMRE 0.0011 0.0024 0.0037 0.0081 0.0262 0.0010 0.0054 0.0135 0.0218 0.0290 phi-BAWW 0.0286 phi-BHVI 0.0024 0.0070 0.0126 0.0207 phi-BLBW 0.0023 0.0060 0.0109 0.0187 0.0283 0.0007 0.0014 0.0020 0.0031 0.0190 phi-BLPW phi-BTBW 0.0005 0.0011 0.0024 0.0066 0.0286 0.0046 0.0089 phi-BTNW 0.0151 0.0220 0.0290 phi-CAWA 0.0012 0.0028 0.0046 0.0093 0.0251

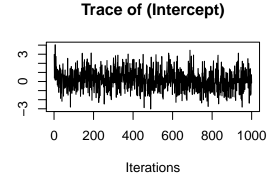
phi-MAWA	0.0004 0.0009	0.0013	0.0018	0.0027
phi-NAWA	0.0012 0.0076	0.0139	0.0215	0.0286
phi-OVEN	0.0011 0.0024	0.0043	0.0088	0.0285
phi-REVI	0.0010 0.0016	0.0020	0.0028	0.0050

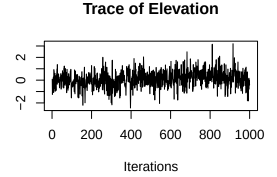
The resulting object out.sp.ms is a list of class spMsPGOcc consisting primarily of posterior samples of all community and species-level parameters, as well as some additional objects that are used for summaries, predictions, and model fit evaluation.

5.3 Convergence diagnostics

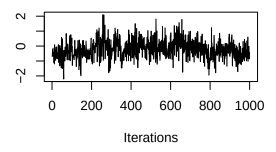
Convergence diagnostics proceed as we have seen with all previous spOccupancy model objects. Posterior samples are returned as coda::mcmc objects, so we can use functions like plot and gelman.diag to assess convergence.

plot(out.sp.ms\$beta.comm.samples, density = FALSE)



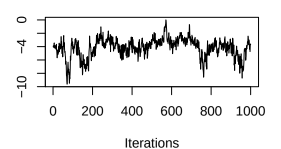


Trace of Elevation.2

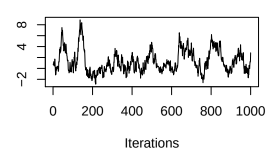


```
# Species-specific effects have yet to converge
plot(out.sp.ms$beta.samples[, 1:4], density = FALSE)
```

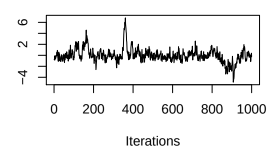
Trace of (Intercept)-AMRE



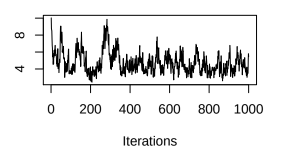
Trace of (Intercept)-BAWW



Trace of (Intercept)-BHVI



Trace of (Intercept)-BLBW



5.4 Posterior predictive checks

We perform posterior predictive checks to assess Goodness of Fit using ppcOcc just as we have previously seen.

```
ppc.sp.ms.out <- ppcOcc(out.sp.ms, 'freeman-tukey', group = 2)

[1] "Currently on species 1 out of 12"
[1] "Currently on species 2 out of 12"
[1] "Currently on species 3 out of 12"
[1] "Currently on species 4 out of 12"
[1] "Currently on species 5 out of 12"
[1] "Currently on species 6 out of 12"
[1] "Currently on species 7 out of 12"
[1] "Currently on species 8 out of 12"
[1] "Currently on species 9 out of 12"
[1] "Currently on species 10 out of 12"
[1] "Currently on species 11 out of 12"
[1] "Currently on species 12 out of 12"
[1] "Currently on species 12 out of 12"
[1] "Currently on species 12 out of 12"</pre>
```

```
Call:
```

ppcOcc(object = out.sp.ms, fit.stat = "freeman-tukey", group = 2)

Chain Information:

Total samples: 10000

Burn-in: 2000 Thin: 8

Total Posterior Samples: 1000

Community Level

Bayesian p-value: 0.4835

Species Level

AMRE Bayesian p-value: 0.395
BAWW Bayesian p-value: 0.405
BHVI Bayesian p-value: 0.52
BLBW Bayesian p-value: 0.511
BLPW Bayesian p-value: 0.546
BTBW Bayesian p-value: 0.516
BTNW Bayesian p-value: 0.49
CAWA Bayesian p-value: 0.277
MAWA Bayesian p-value: 0.514
NAWA Bayesian p-value: 0.593
OVEN Bayesian p-value: 0.554
REVI Bayesian p-value: 0.481
Fit statistic: freeman-tukey

We see all Bayesian p-values are quite large, which is probably at least partly due to the fact that the chains have yet to converge.

5.5 Model selection using WAIC

Below we compute the WAIC using waicOcc and compare it to the WAIC for the non-spatial multispecies occupancy model.

```
waicOcc(out.sp.ms)
```

```
elpd pD WAIC -4288.5133 306.2453 9189.5171
```

waicOcc(out.ms)

```
elpd pD WAIC -4638.54370 72.18475 9421.45690
```

The WAIC for the spatial model is much larger than that for the nonspatial model, potentially indicating we don't need the additional complexities brought in by the species-specific spatial processes. However, we should not place a large emphasis on this since neither of the models has completely converged.

5.6 Prediction

Out-of-sample prediction with spMsPGOcc objects again uses the predict function given a set of covariates and spatial coordinates of unobserved locations. Here we predict values for all 12 species at every 50th cell of the total cells. Results are very similar to the nonspatial multispecies model, so we do not execute the following code.

```
X.O.ms <- cbind(1, hbefElev$val, hbefElev$val^2)
X.O.ms <- X.O.ms[seq(1, nrow(X.O.ms), by = 50), ]
coords.0 <- hbefElev[seq(1, nrow(hbefElev), by = 50), 2:3]
out.sp.ms.pred <- predict(out.sp.ms, X.O.ms, coords.0)
psi.pred.oven <- out.sp.ms.pred$psi.O.samples[, which(sp.names == 'OVEN'), ]
hist(apply(psi.pred.oven, 2, mean), xlab = 'Mean Occurrence Probability')</pre>
```

6 Single species integrated occupancy models

Data integration is a model-based approach that leverages multiple data sources to provide inference and prediction on some latent process of interest. Data integration is particularly relevant in ecology as many data sources are often collected to study a single ecological phenomenon, with each data source having pros and cons. Often, multiple detection-nondetection data sources are available to study the occurrence and distribution of some species of interest. For example, both human point count surveys and autonomous recording units could be used to monitor a bird species of conservation concern. Different types of data have different sources of observation error, which should be explicitly incorporated into a model to avoid attributing any variation in detection probability to the true ecological process. Here we describe single species integrated occupancy models, which combine multiple sources of detection-nondetection data (which may or may not be replicated) in a single hierarchical modeling framework.

6.1 Basic model description

The integrated occupancy model has an identical process model to the single species occupancy model, and has a distinct detection model for each data source that are all conditional on the same shared ecological process (species occurrence).

Let z_j be the presence or absence of a species at site j, with j = 1, ..., J. We assume this latent occurrence process arises from a Bernoulli process following

$$z_j \sim \text{Bernoulli}(\psi_j),$$

 $\text{logit}(\psi_j) = \mathbf{x}'_j \boldsymbol{\beta},$ (11)

where ψ_j is the probability of occurrence at site j, which is a function of site-specific covariates X and a vector of regression coefficients (β) .

We do not directly observe z_j and rather we observe an imperfect representation of the latent occurrence process. In integrated models, we have $r=1,\ldots,R$ distinct sources of data that are all imperfect representations of a single, shared occurrence process. Let $y_{r,a,k}$ be the observed detection (1) or nondetection (0) of a species of interest in data set r at site a during replicate k. Because different data sources have different variables influencing the observation process, we envision a separate detection model for each data source that is conditional on a single, shared ecological process described by Equation (11). We envision the detection-nondetection data from source r as arising from a Bernoulli process conditional on the true latent occurrence process:

$$y_{r,a,k} \sim \text{Bernoulli}(p_{r,a,k}z_{j[a]}),$$

 $\text{logit}(p_{r,a,k}) = \mathbf{v}'_{r,a,k}\alpha_r,$ (12)

where $p_{r,a,k}$ is the probability of detecting a species at site a during replicate k (given it is present at site a) for data source r, which is a function of site, replicate, and data source specific covariates V_r and a vector of

regression coefficients specific to each data source (α_r) . Note that $z_{j[a]}$ is the true occurrence status at site j corresponding to the ath data source site in the given data set r. Each data source may be available at all J sites in the region of interest or at a subset of the J sites. Additionally, data sources can overlap in the sites they sample, or they can be obtained at distinct sites within all J sites of interest in the overall region.

We assume multivariate normal priors for the occurrence (β) and data-set specific detection (α) regression coefficients to complete the Bayesian specification of a single species occupancy model. Polya-Gamma data augmentation is implemented analogous to previous models to yield an efficient implementation of integrated occupancy models.

6.2 Simulating data using simIntOcc

Here we will simulate data to fit with intPGOcc using the function simIntOcc. Given input values for certain parameters, simIntOcc simulates single-species occupancy data from multiple data sources. spOccupancy also includes functions for simulation of single species (simOcc) and multispecies (simMsOcc) data sets that we do not describe in this vignette. simIntOcc takes the following form:

```
simIntOcc(n.data, J.x, J.y, J.obs, n.rep, beta, alpha, sigma.sq = 2,
    phi = 3/0.5, sp = FALSE)
```

simIntOcc simulates data assumed to come from a set of sites distributed in a rectangular sampling design. We next briefly detail the purpose of each of the arguments to simIntOcc.

- n.data: the number of detection-nondetection data sources to simulate.
- J.x: number of sites along the horizontal axis.
- J.y: number of sites along the vertical axis. The total number of sites across the simulated region of interest is J.x x J.y.
- J.obs: a numeric vector with n.data values indicating the number of sites to simulate each data source from.
- n.rep: a list of n.rep elements, where each element is a numeric vector indicating the number of replicates of the detection-nondetection data at each site of a given data source.
- beta: a numeric vector of the simulated regression coefficients for the occurrence portion of the model (including an intercept parameter).
- alpha: a list of n.data elements, where each element is a numeric vector containing the intercept and regression parameters simulated for each data source.

The remaining parameters are used for single species spatial integrated occupancy models, and so we will leave them as their default values for now (which simulates nonspatial data). We simulate data across a region of 225 sites with four data sources.

```
set.seed(101)
J.x < -15
J.y < -15
J.all \leftarrow J.x * J.y
# Number of data sources.
n.data < -4
# Sites for each data source.
J.obs <- sample(ceiling(0.2 * J.all):ceiling(0.5 * J.all), n.data, replace = TRUE)</pre>
# Replicates for each data source.
n.rep <- list()</pre>
for (i in 1:n.data) {
  n.rep[[i]] <- sample(1:4, size = J.obs[i], replace = TRUE)</pre>
}
# Occupancy covariates
beta \leftarrow c(0.5, 1, -2)
p.occ <- length(beta)</pre>
```

The return object dat is a list comprised of a series of objects related to the simulated data that we can use for fitting an integrated occupancy model, verifying the model works as we would expect, and assessing predictive performance of the model at non-sampled locations. For initial model fitting, we are primarily concerned with the objects X.obs, X.p, sites, and y. X.obs is a numeric matrix for the occurrence portion of the occupancy model. X.p is a list of design matrices for the detection portion of the occupancy model, where each element of the list corresponds to a different data source. sites is a list where each element is a vector of site indices for each data source. In an integrated model, data sources can be obtained at the same locations, completely different locations within a single region of interest, or a mix of the two. A key task when working with integrated models is ensuring each data point in a given data source is linked to the correct site in the occurrence portion of the model. For example, consider the simulated sites list below

str(dat\$sites)

```
List of 4
$: num [1:101] 1 3 5 6 9 10 11 13 15 16 ...
$: num [1:90] 3 6 8 12 17 18 26 28 29 30 ...
$: num [1:102] 1 2 3 4 7 9 12 14 21 24 ...
$: num [1:105] 1 6 7 8 9 10 13 14 18 19 ...
```

The indices in the first vector indicate that the first site in the first data source corresponds to overall site 1, the second site in the first data source corresponds to the overall site 3, the third element in the first data source corresponds to the overall site 5, and so on. Finally, the element y consists of the detection-nondetection data matrices for each data source, stored in a list.

```
y <- dat$y
X.p <- dat$X.p
X <- dat$X.obs
sites <- dat$sites</pre>
```

6.3 Fitting single species integrated occupancy models with intPGOcc

The function intPGOcc fits single species integrated occupancy models in spOccupancy. Syntax is very similar to single data source models, and specifically takes the following form:

```
intPGOcc(occ.formula, det.formula, data, starting, n.samples, priors,
    n.omp.threads = 1, verbose = TRUE, n.report = 1000,
    n.burn = round(.10 * n.samples), n.thin = 1, ...)
```

The data argument contains the list of data elements necessary for fitting an integrated occupancy model. For nonspatial integrated occupancy model, data should be a list comprised of the following objects: y (list of detection-nondetection data matrices for each data source), occ.covs (data frame or matrix of covariates for occurrence model), det.covs (a list of lists where each element of the list corresponds to the detection-nondetection data for the given dat asource), sites (a list where each element consists of the site

indices for the given data source. To get started, we'll first package up the simulated data into a a list in the necessary format for the data argument

```
# Occurrence covariates
occ.covs \leftarrow X[, -1, drop = FALSE]
colnames(occ.covs) <- c('occ.cov.1', 'occ.cov.2')</pre>
# Detection covariates
det.covs <- list()</pre>
# Number of covariates on detection for each data source
lapply(alpha, function(a) length(a) - 1)
[[1]]
[1] 2
[[2]]
[1] 3
[[3]]
[1] 1
[[4]]
Γ1 1
# Add detection covariates one by one
det.covs[[1]] <- list(det.cov.1.1 = X.p[[1]][, , 2],</pre>
              det.cov.1.2 = X.p[[1]][, , 3])
det.covs[[2]] <- list(det.cov.2.1 = X.p[[2]][, , 2],</pre>
              det.cov.2.2 = X.p[[2]][, , 3],
              det.cov.2.3 = X.p[[2]][, , 4])
det.covs[[3]] \leftarrow list(det.cov.3.1 = X.p[[3]][, , 2])
det.covs[[4]] <- list(det.cov.4.1 = X.p[[4]][, , 2])</pre>
data.list \leftarrow list(y = y,
          occ.covs = occ.covs,
          det.covs = det.covs,
          sites = sites)
str(data.list)
List of 4
 $ y
           :List of 4
  ..$: int [1:101, 1:4] 0 0 0 1 0 0 0 1 1 1 ...
  ..$: int [1:90, 1:4] 0 1 0 0 1 1 0 0 0 1 ...
  ..$: int [1:102, 1:4] 0 1 0 1 0 0 0 0 0 0 ...
  ..$: int [1:105, 1:4] 0 0 1 1 0 0 1 0 1 1 ...
 $ occ.covs: num [1:202, 1:2] -2.1677 0.5984 0.0431 1.295 0.7063 ...
  ..- attr(*, "dimnames")=List of 2
  .. ..$ : NULL
 ....$ : chr [1:2] "occ.cov.1" "occ.cov.2"
 $ det.covs:List of 4
  ..$ :List of 2
  ....$ det.cov.1.1: num [1:101, 1:4] 2.1575 -0.0408 -1.8213 -0.5695 0.4083 ...
  ....$ det.cov.1.2: num [1:101, 1:4] -0.163 -1.549 -1.703 0.68 -1.078 ...
  ....$ det.cov.2.1: num [1:90, 1:4] 0.519 -1.884 -0.333 0.591 0.468 ...
  ....$ det.cov.2.2: num [1:90, 1:4] -2.033049 0.669742 -2.371423 -0.000267 1.435769 ...
  ....$ det.cov.2.3: num [1:90, 1:4] 0.39 -0.309 0.073 -1.692 -0.218 ...
  ..$ :List of 1
```

```
....$ det.cov.3.1: num [1:102, 1:4] -0.702 -1.253 0.91 -1.371 1.348 ...
..$ :List of 1
....$ det.cov.4.1: num [1:105, 1:4] 0.1934 -0.1415 2.8139 1.0457 -0.0637 ...
$ sites :List of 4
..$ : num [1:101] 1 3 5 6 9 10 11 13 15 16 ...
..$ : num [1:90] 3 6 8 12 17 18 26 28 29 30 ...
..$ : num [1:102] 1 2 3 4 7 9 12 14 21 24 ...
..$ : num [1:105] 1 6 7 8 9 10 13 14 18 19 ...
```

An important thing to note for integrated occupancy models in spOccupancy is that the order of sites should be based on the ordering of the covariates specified in occ.covs. That is, for the site indices for each data source stored in sites, site 1 should correspond to the first row in occ.covs, site 2 should correspond to the second row, and so on.

We specify the occurrence and detection model formulas using the occ.formula, and det.formula arguments. The occ.formula remains unchanged from previous models. For our simulation here, we model occurrence as a function of a two covariates.

```
sim.occ.formula <- ~ occ.cov.1 + occ.cov.2
```

For the detection models, we need to specify a different detection model for each data source. We do this by sending in a list to the det.formula argument, where each element of the list is the model formula for that given data set. Here we specify the four formulas in a list, where the data sources are modeled with 2, 3, 1, and 1 covariates, respectively. The name of the individual list elements is not important. Rather, the position of the model formula should be consistent with the position of the data source in the detection-nondetection data list (y) specified in the data argument.

Next we specify the starting values. Starting values are specified in a list with the following tabs: z (latent occurrence values), alpha (detection regression coefficients), and beta (occurrence regression coefficients. This aligns with fitting single species occupancy models using PGOcc. However, since we now have multiple detection models with different coefficients for each data source, starting values for alpha are now sent passed to intPGOcc as a list, with each element of the list corresponding to the starting detection parameter values for a given data source.

We next specify the priors for all parameters in the integrated occupancy model in a list that is passed into the priors argument. We specify normal priors for both the occurrence and detection regression coefficients, using tags beta.normal and alpha.normal, respectively.

```
alpha.mu <- list()
alpha.var <- list()
for (q in 1:n.data) {
   alpha.mu[[q]] <- rep(0, p.det.long[q])
   alpha.var[[q]] <- rep(2.72, p.det.long[q])
}</pre>
```

Priors for the occurrence regression coefficients are specified as we have seen in previous models. Because we have multiple detection-nondetection data sets each with distinct detection parameters, we specify the hypermeans and hypervariances in individual lists, where each element of the list corresponds to a specific data source. Again, the ordering of the data sources in the lists must align with the order the data sources are saved in the detection-nondetection data supplied to the data argument.

Finally, we specify the number of samples, burn-in, and thinning rate using the same approach we have used for previous models.

```
n.samples <- 5000
n.burn <- 1000
n.thin <- 1
```

We can now run the integrated occupancy model. Below we set the number of threads used to 1 and print out sampler progress after every 1000th iteration.

 Sampled: 3000 of 5000, 60.00% _____ Sampled: 4000 of 5000, 80.00% Sampled: 5000 of 5000, 100.00% We again consult the summary function for a concise description of the model results. summary(out.int) Call: intPGOcc(occ.formula = sim.occ.formula, det.formula = sim.det.formula, data = data.list, starting = starting.list, priors = prior.list, n.samples = n.samples, n.omp.threads = 1, verbose = TRUE, n.report = 1000, n.burn = n.burn, n.thin = n.thin) Chain Information: Total samples: 5000 Burn-in: 1000 Thin: 1 Total Posterior Samples: 4000 Occurrence. 2.5% 25% 50% (Intercept) 0.0514 0.3178 0.4563 0.5988 0.8784 0.8308 1.1409 1.3145 1.4969 1.8788 occ.cov.1 -3.0320 -2.5006 -2.2431 -1.9965 -1.6012 occ.cov.2 Data source 1 Detection: 25% 2.5% 50% 75% 97.5% (Intercept) 0.5393 0.7922 0.9350 1.0759 1.3547 det.cov.1.1 -1.1305 -0.8228 -0.6764 -0.5189 -0.2423 det.cov.1.2 0.2396 0.5356 0.6778 0.8288 1.1198 Data source 2 Detection: 2.5% 25% 50% 75% 97.5% (Intercept) -0.6192 -0.3523 -0.2026 -0.0625 det.cov.2.1 -1.3877 -1.0319 -0.8686 -0.7076 -0.4300 det.cov.2.2 -0.1577 0.0902 0.2186 0.3540 0.6073 det.cov.2.3 0.6245 0.9673 1.1593 1.3588 1.7550 Data source 3 Detection: 2.5% 50% 75% 25% (Intercept) -0.722 -0.4981 -0.3853 -0.2688 -0.0427 det.cov.3.1 -0.771 -0.5242 -0.4067 -0.2876 -0.0636 Data source 4 Detection: 25% 2.5% 50% 75% 97.5%

(Intercept) 0.3428 0.5469 0.6618 0.7805 1.0147 det.cov.4.1 0.4121 0.6572 0.7860 0.9252 1.2178

The summary function for integrated models returns the detection parameters separately for each detection covariate. We can compare these results to the values we used to simulate the data. It looks like our model does a pretty good job of recovering the true parameter values.

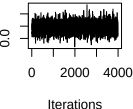
6.4 Convergence diagnostics

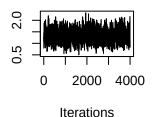
Posterior samples are returned as coda::mcmc objects, so as with all spOccupancy model objects, we use standard coda functions like plot and gelman.diag to assess convergence.

```
# Occurrence effects
plot(out.int$beta.samples, density = FALSE)
```

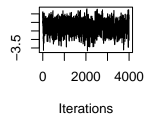


Trace of occ.cov.1





Trace of occ.cov.2



6.5 Posterior predictive checks

We perform posterior predictive checks using ppcocc as before. GoF assessment for integrated models is an active area of research. In sp0ccupancy, we compute posterior predictive checks separately for each dataset in the integrated model.

```
ppc.int.out <- ppcOcc(out.int, 'freeman-tukey', group = 2)
summary(ppc.int.out)

Call:
ppcOcc(object = out.int, fit.stat = "freeman-tukey", group = 2)

Chain Information:
Total samples: 5000
Burn-in: 1000
Thin: 1
Total Posterior Samples: 4000

Data Source 1

Bayesian p-value: 0.517</pre>
```

```
Fit statistic: freeman-tukey

Data Source 2

Bayesian p-value: 0.78475

Fit statistic: freeman-tukey

Data Source 3

Bayesian p-value: 0.35975

Fit statistic: freeman-tukey

Data Source 4

Bayesian p-value: 0.9115

Fit statistic: freeman-tukey
```

6.6 Model selection using WAIC

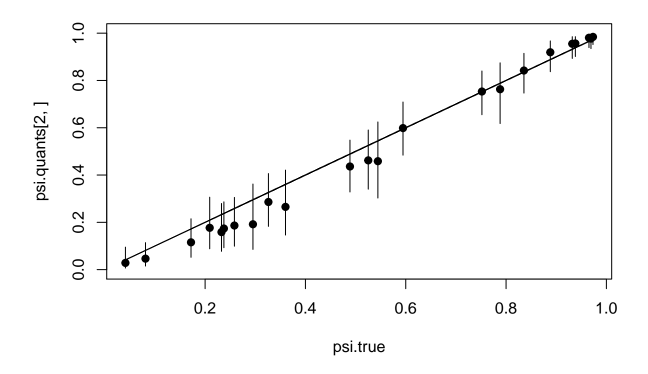
We use waicOcc to compute the WAIC for integrated occupancy models. Similar to the posterior predictive check, individual WAIC values are reported for each data set. These can be summed across all data sources for an overall WAIC value if desired.

```
waicOcc(out.int)

    elpd    pD    WAIC
1 -86.91207 5.899967 185.62407
2 -35.29849 1.497667 73.59231
3 -48.35002 1.374544 99.44913
4 -74.30623 2.487703 153.58786
```

6.7 Prediction

Prediction for integrated occupancy models proceeds exactly as before using predict. The simIntOcc function automatically splits the data into a set of sampled sites and nonsampled sites. We can use the simulated data from the non-sampled sites to assess the ability of our model to make accurrate out of sample predictions.



7 Single species spatial integrated occupancy models

7.1 Basic model description

Single species spatial integrated occupancy models are identical to integrated occupancy models except the ecological process model now incorporates a spatially-structured random effect following the discussion in Section 3. All details for the single species integrated spatial occupancy model have already been presented in previous model descriptions.

7.2 Simulating data using simIntOcc

Here we again use simIntOcc to simulate a data set to display a single species spatial integrated occupancy model. We use the simIntOcc function just as we did in Section ??, except we now specify the data to be simulated with spatially structured random effects following an exponential correlation function. Notice we now specify the spatial variance (sigma.sq) and spatial range (phi) parameters that we need to simulate the spatial random effects. We also set sp = TRUE to include the spatial random effects in the data simulation.

```
set.seed(405)
J.x <- 15
J.y <- 15
J.all <- J.x * J.y
# Number of data sources.
n.data <- 4</pre>
```

```
# Sites for each data source.
J.obs <- sample(ceiling(0.2 * J.all):ceiling(0.5 * J.all), n.data, replace = TRUE)
# Replicates for each data source.
n.rep <- list()</pre>
for (i in 1:n.data) {
  n.rep[[i]] <- sample(1:4, size = J.obs[i], replace = TRUE)</pre>
# Occupancy covariates
beta <-c(0.5, 0.5)
p.occ <- length(beta)</pre>
# Detection covariates
alpha <- list()</pre>
for (i in 1:n.data) {
  # A single detection covariate effect for each data set.
  alpha[[i]] <- runif(2, -1, 1)
p.det.long <- sapply(alpha, length)</pre>
p.det <- sum(p.det.long)</pre>
sigma.sq <- 3
phi <-3 / .5
sp <- TRUE
# Simulate occupancy data.
dat <- simIntOcc(n.data = n.data, J.x = J.x, J.y = J.y, J.obs = J.obs,
         n.rep = n.rep, beta = beta, alpha = alpha, sigma.sq = sigma.sq,
         phi = phi, sp = sp)
y <- dat$y
X.p <- dat$X.p</pre>
X <- dat$X.obs</pre>
sites <- dat$sites
coords <- as.matrix(dat$coords.obs)</pre>
```

7.3 Fitting single speces spatial integrated occupancy models using spIntPGOcc

The function spIntPGOcc fits single species spatial integrated occupancy models in spOccupancy. Syntax is very similar to single data source models and specifically takes the following form:

```
spIntPGOcc(occ.formula, det.formula, data, starting, n.batch,
    batch.length, accept.rate = 0.43, priors,
    cov.model = "exponential", tuning, n.omp.threads = 1,
    verbose = TRUE, NNGP = TRUE, n.neighbors = 15,
    search.type = 'cb', n.report = 100,
    n.burn = round(.10 * n.batch * batch.length),
    n.thin = 1, ...)
```

The occ.formula, det.formula, and data arguments are analogous to what we saw with the nonspatial integrated occupancy model. However, as for all spatial models in spOccupancy, the data list must also contain the spatial coordinates.

```
# Occurrence covariates
occ.covs <- X[, -1, drop = FALSE]
colnames(occ.covs) <- c('occ.cov.1')
# Detection covariates
det.covs <- list()</pre>
```

```
# Number of covariates on detection for each data source
lapply(alpha, function(a) length(a) - 1)
[[1]]
[1] 1
[[2]]
[1] 1
[[3]]
[1] 1
[[4]]
[1] 1
# Add detection covariates one by one
det.covs[[1]] <- list(det.cov.1.1 = X.p[[1]][, , 2])</pre>
det.covs[[2]] \leftarrow list(det.cov.2.1 = X.p[[2]][, , 2])
det.covs[[3]] \leftarrow list(det.cov.3.1 = X.p[[3]][, , 2])
det.covs[[4]] <- list(det.cov.4.1 = X.p[[4]][, , 2])</pre>
data.list <- list(y = y,</pre>
          occ.covs = occ.covs,
          det.covs = det.covs,
          sites = sites,
          coords = coords)
str(data.list)
List of 5
 $у
           :List of 4
  ..$: int [1:101, 1:4] 0 0 0 1 0 0 0 0 1 1 ...
  ..$: int [1:93, 1:4] 0 1 0 0 0 0 0 0 0 0 ...
  ..$: int [1:74, 1:4] 1 1 0 1 0 1 1 1 0 1 ...
  ..$: int [1:109, 1:4] 0 1 0 0 0 0 1 1 1 1 ...
 $ occ.covs: num [1:200, 1] -0.9567 0.0462 -1.1998 -1.7878 0.6352 ...
  ..- attr(*, "dimnames")=List of 2
  .. ..$ : NULL
  .. ..$ : chr "occ.cov.1"
 $ det.covs:List of 4
  ..$ :List of 1
  ....$ det.cov.1.1: num [1:101, 1:4] 0.6613 -0.217 -0.0912 1.6326 -0.0668 ...
  ..$ :List of 1
  ....$ det.cov.2.1: num [1:93, 1:4] -0.318 -0.849 -1.132 -0.457 0.742 ...
  ..$ :List of 1
  .. ..$ det.cov.3.1: num [1:74, 1:4] -0.673 1.258 -0.511 -0.587 -1.143 ...
  ..$ :List of 1
 .. ..$ det.cov.4.1: num [1:109, 1:4] 1.152 -0.948 1.657 1.154 0.789 ...
 $ sites :List of 4
  ..$ : num [1:101] 1 3 5 7 8 9 10 13 14 15 ...
  ..$: num [1:93] 3 6 7 10 11 15 16 17 20 21 ...
  ..$: num [1:74] 1 2 5 10 11 12 13 15 16 20 ...
  ..$: num [1:109] 1 2 4 6 7 9 13 14 15 17 ...
 $ coords : num [1:200, 1:2] 0 0.0714 0.2143 0.3571 0.4286 ...
  ..- attr(*, "dimnames")=List of 2
  ....$ : chr [1:200] "1" "2" "4" "6" ...
  ....$ : chr [1:2] "Var1" "Var2"
```

Starting values specified in starting and priors in priors are specified in the same form as for intPGOcc with the additional values for spatial parameters. Analogous to all other spatial models in spOccupancy, the spatial variance parameter takes an inverse-Gamma prior and the spatial range parameter (and the spatial smoothness parameter if cov.model = 'matern') takes a uniform prior.

```
J <- nrow(occ.covs)</pre>
alpha.start <- list()</pre>
for (q in 1:n.data) {
  alpha.start[[q]] <- rep(0, p.det.long[q])
starting.list <- list(alpha = alpha.start,</pre>
               beta = rep(0, p.occ),
               z = rep(1, J),
               phi = 3 / .4,
               sigma.sq = 1,
               w = rep(0, J)
alpha.mu <- list()</pre>
alpha.var <- list()</pre>
for (q in 1:n.data) {
  alpha.mu[[q]] <- rep(0, p.det.long[q])</pre>
  alpha.var[[q]] \leftarrow rep(2.72, p.det.long[q])
prior.list <- list(beta.normal = list(mean = rep(0, p.occ),</pre>
                        var = rep(2.72, p.occ)),
            alpha.normal = list(mean = alpha.mu,
                             var = alpha.var),
            sigma.sq.ig = c(2, 2),
            phi.unif = c(3/1, 3/.1))
```

Finally, we specify the remaining parameters regarding the NNGP specifications, tuning parameters, and the length of the MCMC sampler we will run. We are then all set to run the model.

```
tuning <- list(phi = 1)</pre>
batch.length <- 25
n.batch <- 400
cov.model <- "exponential"</pre>
out.sp.int <- spIntPGOcc(occ.formula = sim.occ.formula,</pre>
             det.formula = sim.det.formula,
             data = data.list,
             starting = starting.list,
             n.batch = n.batch,
             batch.length = batch.length,
             priors = prior.list,
             cov.model = cov.model,
             NNGP = TRUE,
             n.neighbors = 5,
             n.report = 100,
             n.burn = 1000,
             n.thin = 5)
```

```
Preparing the data
_____
   Building the neighbor list
_____
Building the neighbors of neighbors list
_____
  Model description
_____
NNGP Integrated Occupancy Model with Polya-Gamma latent
variable fit with 200 sites.
Integrating 4 occupancy data sets.
Number of MCMC samples: 10000 (400 batches of length 25)
Burn-in: 1000
Thinning Rate: 5
Total Posterior Samples: 1800
Using the exponential spatial correlation model.
Using 5 nearest neighbors.
Source compiled with OpenMP support and model fit using 1 thread(s).
Adaptive Metropolis with target acceptance rate: 43.0
Sampling ...
Batch: 100 of 400, 25.00%
   parameter acceptance tuning
   phi 24.0 0.81058
Batch: 200 of 400, 50.00%
   parameter acceptance tuning
   phi 60.0 0.70469
  _____
Batch: 300 of 400, 75.00%
   parameter acceptance tuning
   phi 44.0 0.69073
Batch: 400 of 400, 100.00%
   parameter acceptance tuning
   phi 0.0 0.69768
summary(out.sp.int)
spIntPGOcc(occ.formula = sim.occ.formula, det.formula = sim.det.formula,
   data = data.list, starting = starting.list, priors = prior.list,
```

cov.model = cov.model, NNGP = TRUE, n.neighbors = 5, n.batch = n.batch,

batch.length = batch.length, n.report = 100, n.burn = 1000,

```
n.thin = 5)
```

Chain Information: Total samples: 10000

Burn-in: 1000

Thin: 5

Total Posterior Samples: 1800

Occurrence:

2.5% 25% 50% 75% 97.5% (Intercept) 0.2502 0.8066 1.1237 1.4328 2.2670 occ.cov.1 -0.5009 -0.1323 0.0619 0.2382 0.5967

Data source 1 Detection:

2.5% 25% 50% 75% 97.5% (Intercept) -0.2356 -0.0162 0.1067 0.2229 0.4635 det.cov.1.1 -1.3791 -1.0930 -0.9356 -0.7997 -0.5595

Data source 2 Detection:

2.5% 25% 50% 75% 97.5% (Intercept) -0.9854 -0.7445 -0.6169 -0.4931 -0.2873 det.cov.2.1 -0.8979 -0.6490 -0.5179 -0.3989 -0.1694

Data source 3 Detection:

2.5% 25% 50% 75% 97.5% (Intercept) 0.4250 0.6596 0.7976 0.9301 1.2136 det.cov.3.1 -1.2075 -0.8878 -0.7242 -0.5817 -0.3024

Data source 4 Detection:

2.5% 25% 50% 75% 97.5% (Intercept) 0.1425 0.3348 0.4335 0.5439 0.758 det.cov.4.1 -0.7901 -0.5740 -0.4569 -0.3537 -0.152

Covariance:

2.5% 25% 50% 75% 97.5% sigma.sq 0.8024 1.8483 3.0164 4.9325 14.7629 phi 3.7089 6.6381 8.9619 11.8340 21.0222

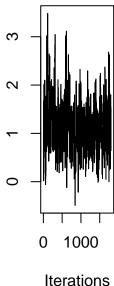
7.4 Convergence diagnostics

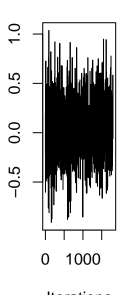
We use the coda package to explore the trace plots. The trace plot below suggests we may want to run the model for longer to ensure convergence.

plot(out.sp.int\$beta.samples, density = FALSE)

Trace of (Intercept

Trace of occ.cov.1





iterations

Iterations

7.5 Posterior predictive checks

Data Source 3

Below we perform a poseterior predictive check for each of the data sets included in the occupancy model using ppcOcc.

```
ppc.sp.int.out <- ppcOcc(out.sp.int, 'freeman-tukey', group = 2)
summary(ppc.sp.int.out)

Call:
ppcOcc(object = out.sp.int, fit.stat = "freeman-tukey", group = 2)

Chain Information:
Total samples: 10000
Burn-in: 1000
Thin: 5
Total Posterior Samples: 1800

Data Source 1

Bayesian p-value: 0.8122222
Fit statistic: freeman-tukey

Data Source 2

Bayesian p-value: 0.3633333
Fit statistic: freeman-tukey</pre>
```

```
Bayesian p-value: 0.73
Fit statistic: freeman-tukey
Data Source 4
Bayesian p-value: 0.8494444
Fit statistic: freeman-tukey
```

According to the Bayesian p-values, there is no lack of fit for any of the three data sets.

7.6 Model selection using WAIC

We can perform model selection using WAIC with the waicOcc function as we have seen previously. Below, we reanalyze the simulated data using the intPGOcc function and compare the WAIC for the spatial and nonspatial integrated occupancy models

```
out.no.sp.int <- intPGOcc(occ.formula = sim.occ.formula,
              det.formul = sim.det.formula,
              data = data.list,
              starting = starting.list,
              priors = prior.list,
              n.samples = n.batch * batch.length,
              n.report = 2000,
              n.burn = 1000,
              n.thin = 5,
              verbose = FALSE)
waicOcc(out.sp.int)
       elpd
                           WAIC
                   рD
1 -66.96099 14.544463 163.01092
2 -35.01689 6.417414 82.86861
3 -36.78481 8.221961 90.01355
4 -77.13772 13.436789 181.14901
```

```
elpd pD WAIC
1 -71.03359 1.8575341 145.78224
2 -38.50946 0.6521299 78.32317
3 -39.92932 1.1152679 82.08918
4 -85.13409 1.5408829 173.34995
```

waicOcc(out.no.sp.int)

We see for this specific set of simulated data the WAIC is smaller for all four data sources in the spatial model compared to the nonspatial model.

7.7 Prediction

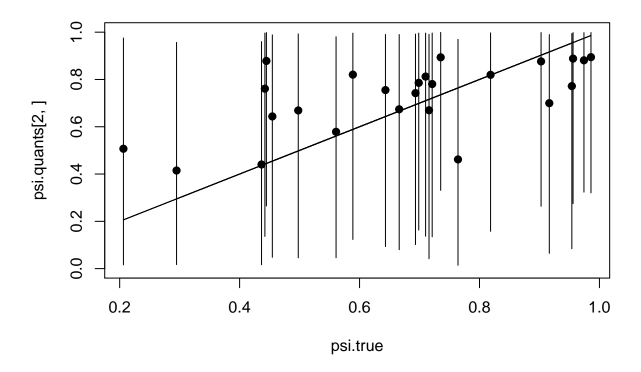
Prediction for spatial integrated occupancy models proceeds exactly analogous to our approach using nonspatial integrated occupancy models. The only difference is that now we must also provide the coordinates of the nonsampled locations.

```
X.0 <- dat$X.pred
coords.0 <- dat$coords.pred
out.sp.int.pred <- predict(out.sp.int, X.0, coords.0)</pre>
```

```
Prediction description
_____
NNGP Occupancy model with Polya-Gamma latent
variable fit with 200 observations.
Number of covariates 2 (including intercept if specified).
Using the exponential spatial correlation model.
Using 5 nearest neighbors.
Number of MCMC samples 1800.
Predicting at 25 locations.
Source compiled with OpenMP support and model fit using 1 threads.
_____
      Predicting
_____
Location: 25 of 25, 100.00%
# Compare predictions to true simulated values
psi.true <- dat$psi.pred</pre>
psi.quants <- apply(out.sp.int.pred$psi.0.samples, 2, quantile,</pre>
          probs = c(0.025, 0.5, 0.975))
plot(psi.true, psi.quants[2, ], pch = 19, ylim = c(0, 1))
```

segments(psi.true, psi.quants[1,], y1 = psi.quants[3,])

lines(psi.true, psi.true)



For this data set, the predictions seem to capture the trend in the non-sampled data, but the predictions have large amounts of uncertainty.

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