

Queensland rainforest

```
library(ppjsdm)
library(spatstat)
#> Loading required package: spatstat.data
#> Loading required package: nlme
#> Loading required package: rpart
#>
#> spatstat 1.62-2      (nickname: 'Shape-shifting lizard')
#> For an introduction to spatstat, type 'beginner'
remove(list = ls())

source("../R/get_qld.R")

set.seed(1)
```

This vignette explains how to use the `ppjsdm` package with the Queensland rainforest dataset from CSIRO. We begin by loading the data with the most prevalent species.

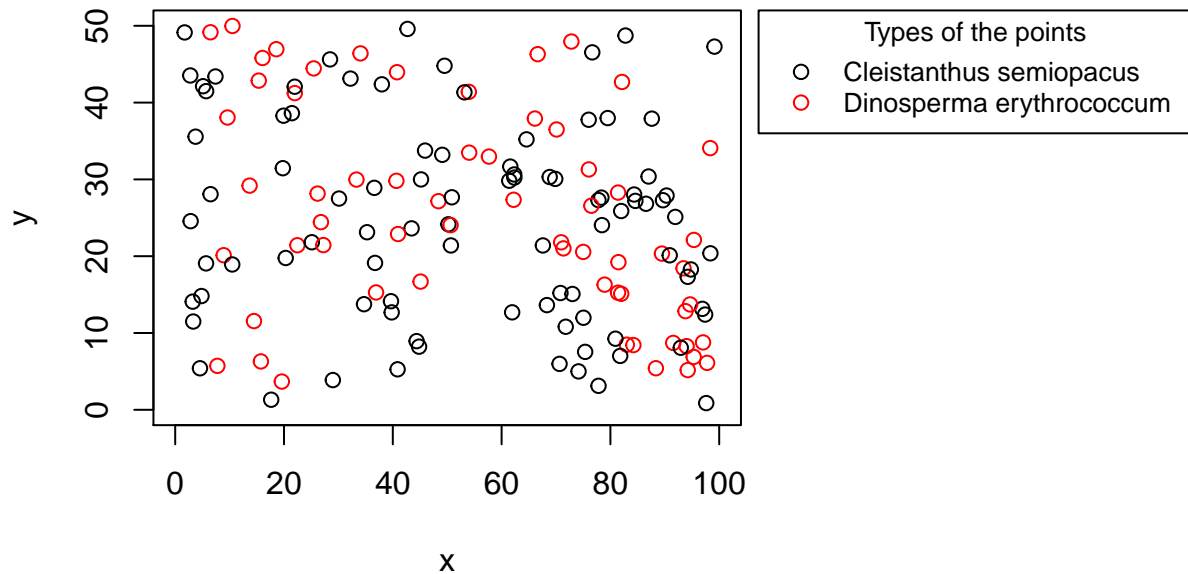
```
index_of_plot <- 3 # Between 1 and 20
year <- 2011 # Year of census
number_of_species <- 2 # Includes the most prevalent species from the plot

qld <- get_qld(index = index_of_plot,
               year = year,
               prevalent = number_of_species)
#> The chosen index corresponds to ep2.
configuration <- qld$configuration
window <- qld$window
```

The point configuration is plotted below.

```
par(mar = c(5, 4, 4, 13) + 0.1)
plot(configuration, window = window)
```

Points in the configuration



The function `gibbsm` fits a multivariate Gibbs point process to our dataset. For example,

```
fit <- ppjsdm::gibbsm(configuration, window = window, use_glmnet = FALSE)
#> log_lambda_1 log_lambda_2 alpha_1_1 alpha_1_2 alpha_2_2
#> -3.978784 -4.431216 -3.936413 8.070406 -3.636435
summary(fit)
#>
#> Call:
#> glm(formula = as.formula(gibbsm_data$formula), family = binomial(),
#> data = as.data.frame(gibbsm_data$data))
#>
#> Deviance Residuals:
#> Min 1Q Median 3Q Max
#> -0.6626 -0.5623 -0.5162 -0.4602 2.3202
#>
#> Coefficients:
#> Estimate Std. Error z value Pr(>|z|)
#> log_lambda_1 -3.9788 0.1161 -34.283 <2e-16 ***
#> log_lambda_2 -4.4312 0.1377 -32.188 <2e-16 ***
#> alpha_1_1 -3.9364 2.3778 -1.656 0.0978 .
#> alpha_1_2 8.0704 4.5335 1.780 0.0750 .
#> alpha_2_2 -3.6364 2.2191 -1.639 0.1013
#> ---
#> Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
#>
#> (Dispersion parameter for binomial family taken to be 1)
#>
#> Null deviance: 4824.57 on 1151 degrees of freedom
#> Residual deviance: 886.37 on 1146 degrees of freedom
#> AIC: 896.37
```

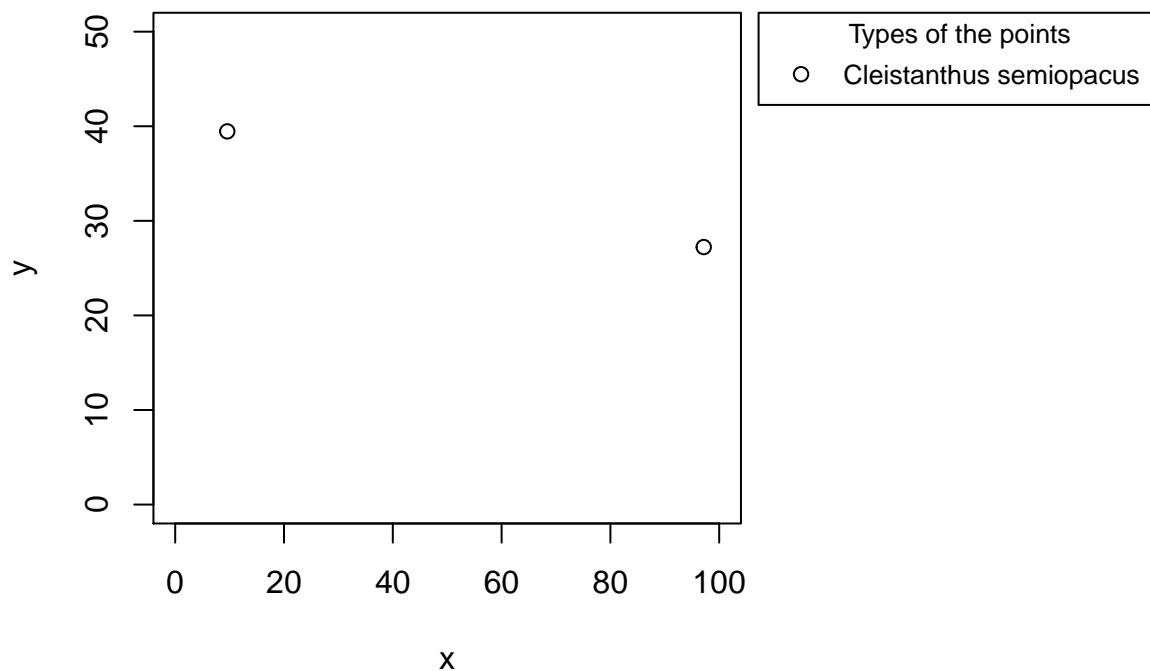
```
#>
#> Number of Fisher Scoring iterations: 4
```

By default, the function fits the model that was introduced in the ARC grant [TODO: Add reference]. This model has many drawbacks, the most important of which is that the model with the fitted values is degenerate, as can be seen by drawing from the model.

```
lambda <- exp(fit$coefficients[1:number_of_species])
alpha <- matrix(c(fit$coefficients[3],
                  fit$coefficients[4],
                  fit$coefficients[4],
                  fit$coefficients[5]),
                ncol = number_of_species,
                nrow = number_of_species)
draw <- ppjsdm::rgibbs(window = window,
                       alpha = alpha,
                       lambda = lambda,
                       types = levels(types(configuration)),
                       steps = 1000000)
```

```
par(mar = c(5, 4, 4, 13) + 0.1)
plot(draw, window = window)
```

Points in the configuration



The Geyer model is better suited to most situations, but the user needs to specify some additional parameters before the fitting may take place.

```
radii <- matrix(5, number_of_species, number_of_species)
```

The matrix `radii` models interaction radii within a species, and between species. An interaction radius of 5 gives good results in the fitting procedure.

```
fit <- ppjsdm::gibbsm(configuration,
                      window = window,
                      model = "Geyer",
                      radius = radii,
                      use_glmnet = FALSE)
#> log_lambda_1 log_lambda_2 alpha_1_1 alpha_1_2 alpha_2_2
#> -4.22694003 -4.85656953 0.16730695 0.02636396 0.49504414
print(fit)
#>
#> Call: glm(formula = as.formula(gibbsm_data$formula), family = binomial(),
#> data = as.data.frame(gibbsm_data$data))
#>
#> Coefficients:
#> log_lambda_1 log_lambda_2 alpha_1_1 alpha_1_2 alpha_2_2
#> -4.22694 -4.85657 0.16731 0.02636 0.49504
#>
#> Degrees of Freedom: 1151 Total (i.e. Null); 1146 Residual
#> Null Deviance: 4825
#> Residual Deviance: 880.5 AIC: 890.5
```

Note that in this case the model is not degenerate; indeed we can sample from it as follows.

```
lambda <- exp(fit$coefficients[1:number_of_species])
alpha <- matrix(c(fit$coefficients[3],
                  fit$coefficients[4],
                  fit$coefficients[4],
                  fit$coefficients[5]),
               ncol = number_of_species,
               nrow = number_of_species)
draw <- ppjsdm::rgibbs(window = window,
                       alpha = alpha,
                       lambda = lambda,
                       model = "Geyer",
                       radius = radii,
                       types = levels(types(configuration)),
                       steps = 1000000)
```

```
par(mar = c(5, 4, 4, 13) + 0.1)
plot(draw, window = window)
```

Points in the configuration

