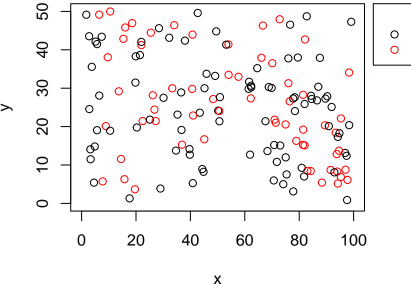
## Queensland rainforest

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.

The point configuration is plotted below.

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

## Points in the configuration



Types of the points

- Cleistanthus semiopacus
- Dinosperma erythrococcum

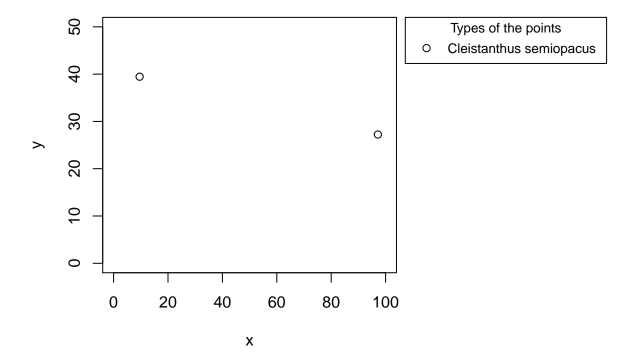
The function gibbsm fits a multivariate Gibbs point process to our dataset. For example,

```
fit <- ppjsdm::gibbsm(configuration, window = window, use_glmnet = FALSE)
                               alpha_1_1
#> log_lambda_1 log_lambda_2
                                             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$formula), family = binomial(),
      data = as.data.frame(qibbsm_data$data))
#>
#> Deviance Residuals:
               1Q
                    Median
#>
      Min
                                   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 ***
                             0.1377 -32.188
\#> log_lambda_2 -4.4312
                                              <2e-16 ***
#> alpha_1_1
                 -3.9364
                             2.3778 -1.656
                                              0.0978 .
                                     1.780
                                              0.0750 .
#> alpha_1_2
                8.0704
                             4.5335
#> 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.

## 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)</pre>
```

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:
                                alpha\_1\_1
#> log_lambda_1 log_lambda_2
                                             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.

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

## Points in the configuration

