# Bayesian inference of an individual-based mutualistic network 12\_01

### Net 12\_01

```
library(BayesianNetworks)
library(network.tools)
library(tidyverse)
theme_set(theme_minimal())
options(mc.cores = 4)
```

#### Data

Load dataset and sampling effort per individual plant:

## dbl (3): crop, se\_obs\_h, seeds\_per\_fruit

```
web <- readr::read_csv(here::here("data/nets_raw", paste0(params$net, "_int.csv"))) |>
  arrange(ind)
## Rows: 8 Columns: 7
## -- Column specification -
## Delimiter: ","
## chr (1): ind
## dbl (6): Leptocoma_zeylonica, Pycnonotus_luteolus, Pycnonotus_cafer, Dicaeum_erythrorhynchos, Pycnon
## i Use 'spec()' to retrieve the full column specification for this data.
## i Specify the column types or set 'show_col_types = FALSE' to quiet this message.
mat <- as.matrix(web[, -1])</pre>
mat <- apply(mat, c(1,2), as.integer)</pre>
rownames(mat) <- web$ind</pre>
# create numeric vector of sampling effort for each plant with names = plant id
effort <- readr::read_csv(here::here("data/nets_attr", paste0(params$net, "_attr.csv"))) |>
  select(ind, starts_with("se_")) |>
  filter(ind %in% web$ind) |>
 arrange(ind)
## Rows: 8 Columns: 7
## -- Column specification -----
## Delimiter: ","
## chr (4): ind, neigh_inter, neigh_intra, fruit_color
```

```
##
## i Use 'spec()' to retrieve the full column specification for this data.
## i Specify the column types or set 'show_col_types = FALSE' to quiet this message.
## If there is only one column with sampling effort, use it:
if (!net %in% c("10 01", "15 01", "18 01", "18 02", "20 01", "21 01", "21 02")) {
  effort <- effort |>
    pull(starts_with("se_"), name = "ind")
}
# Otherwise, select sampling effort column in some specific nets:
if (net == "10_01") {
  effort <- effort |>
    mutate(se_cam_days = se_cam_h/24) |>
    pull(se_cam_days, name = "ind")
}
if (net == "15 01") {
  effort <- effort |>
    pull(se_cam_days, name = "ind")
}
if (net %in% c("18_01", "18_02", "20_01")) {
  effort <- effort |>
    mutate(se bc months = se bc days/30) |>
    pull(se_bc_months, name = "ind")
}
if (net %in% c("21_01", "21_02")) {
  effort <- effort |>
    pull(se_obs_h, name = "ind")
}
## Some nets may require adjusting of the count data or effort values
## Insert that here eg.
# if (params$net == "01_01") {
# mat <- round(mat/10)
  mat \leftarrow apply(mat, c(1,2), as.integer)
# }
stopifnot(identical(length(effort), nrow(mat)))
stopifnot(identical(names(effort), rownames(mat)))
summary(as.numeric(mat))
      Min. 1st Qu. Median
                              Mean 3rd Qu.
                                              Max.
## 0.0000 0.0000 0.0000 0.9583 1.0000 5.0000
if (max(mat) > 500) {
  stop("More than 500 counts in some cell(s)")
}
summary(effort)
```

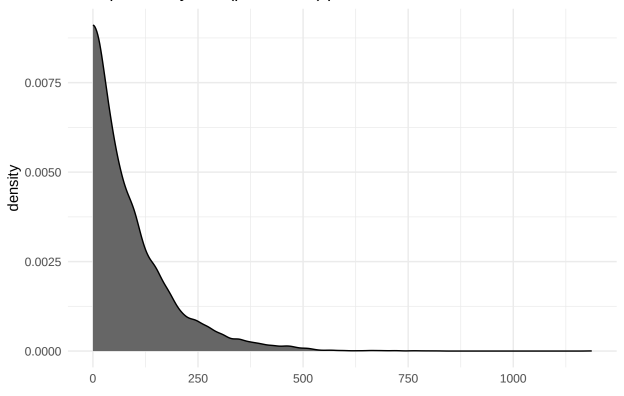
```
## Min. 1st Qu. Median Mean 3rd Qu. Max.
## 1.200 1.950 2.000 1.875 2.000 2.000

if (max(effort) > 500) {
   stop("Sampling effort > 500 for some plants")
}
```

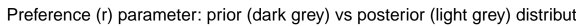
## Bayesian inference of network structure

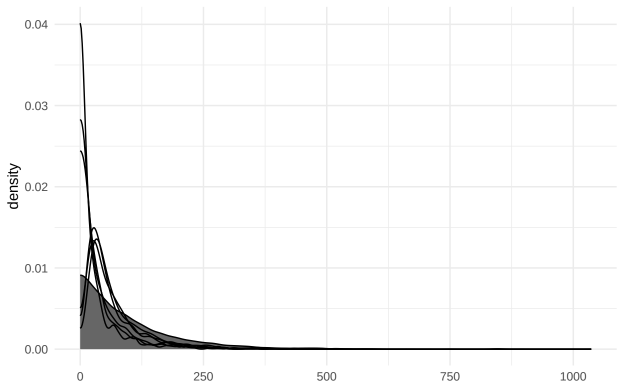
```
dt <- prepare_data(mat, sampl.eff = effort)
plot_prior(params$beta)</pre>
```

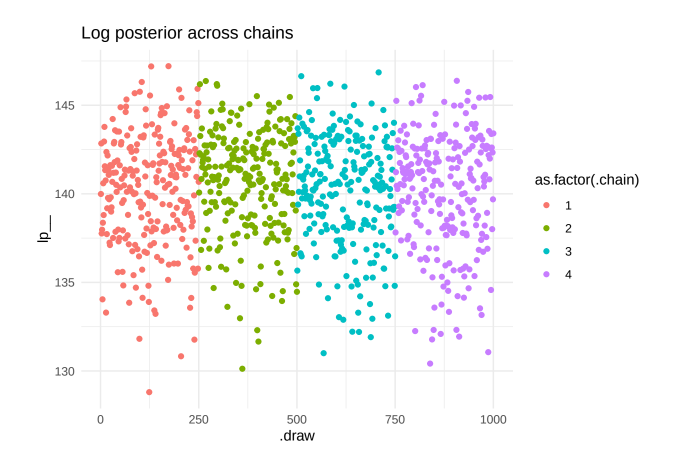
# Prior probability for r (preference) parameter with beta = 0.01



```
## Running MCMC with 4 parallel chains...
##
## Chain 4 finished in 3.2 seconds.
## Chain 1 finished in 3.7 seconds.
## Chain 3 finished in 3.7 seconds.
## Chain 2 finished in 4.1 seconds.
## All 4 chains finished successfully.
## Mean chain execution time: 3.7 seconds.
## Total execution time: 4.1 seconds.
get_seed(fit)
## [1] 563921464
check_model(fit, data = dt)
## Processing csv files: C:/Users/frodr/AppData/Local/Temp/RtmpkDNfRE/varying_preferences-202406241426-
## Checking sampler transitions treedepth.
## Treedepth satisfactory for all transitions.
## Checking sampler transitions for divergences.
## No divergent transitions found.
##
## Checking E-BFMI - sampler transitions HMC potential energy.
## E-BFMI satisfactory.
## Effective sample size satisfactory.
## Split R-hat values satisfactory all parameters.
##
## Processing complete, no problems detected.
```







#### Posteriors

Get posterior distributions:

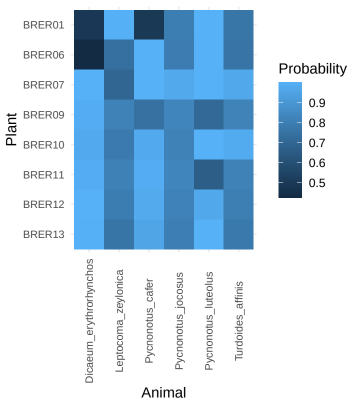
```
post <- get_posterior(fit, dt)
head(post)</pre>
```

```
## # A tibble: 6 x 11
## # Groups:
              Animal, Plant [6]
                     .chain .iteration .draw connectance preference plant.abund animal.abund int.pr
##
    Plant Animal
     <chr> <chr>
                         <int>
                                    <int> <int>
                                                       <dbl>
                                                                  <dbl>
                                                                              <dbl>
                                                                                           <dbl>
                                                                                                    <db
## 1 BRER01 Leptocoma_z~
                                                       0.849
                                                                   33.9
                                                                            0.156
                                                                                          0.0865
                                                                                                    1.0
                                         1
                             1
## 2 BRER06 Leptocoma_z~
                             1
                                         1
                                                       0.849
                                                                   33.9
                                                                            0.187
                                                                                          0.0865
                                                                                                    0.7
## 3 BRER07 Leptocoma_z~
                             1
                                         1
                                              1
                                                       0.849
                                                                   33.9
                                                                            0.221
                                                                                          0.0865
                                                                                                    0.6
## 4 BRER09 Leptocoma_z~
                             1
                                         1
                                                       0.849
                                                                   33.9
                                                                            0.00326
                                                                                          0.0865
                                                                                                    0.8
                                                                                                    0.7
## 5 BRER10 Leptocoma_z~
                                                                   33.9
                             1
                                         1
                                              1
                                                       0.849
                                                                            0.0663
                                                                                          0.0865
## 6 BRER11 Leptocoma_z~
                                                       0.849
                                                                   33.9
                                                                            0.152
                                                                                          0.0865
                                                                                                    0.6
```

Mean edge probability:

```
plot_interaction_prob(post)
```



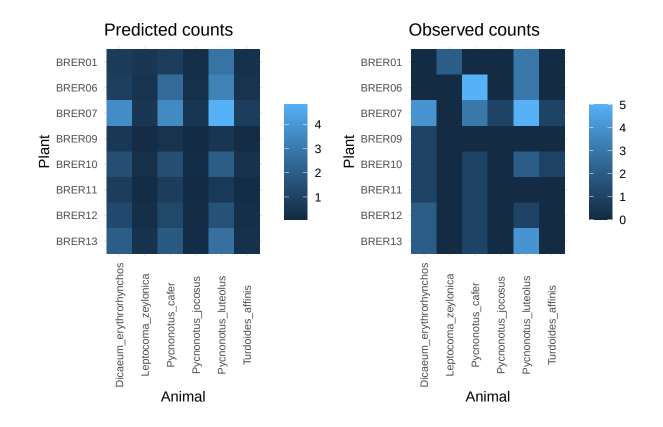


# Generate predicted visits for each pairwise interaction

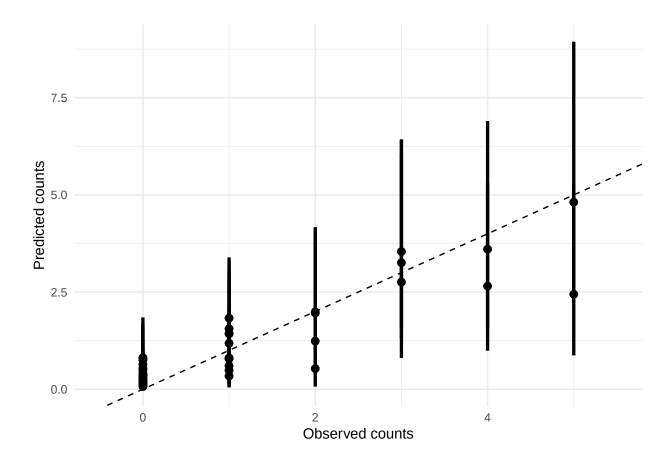
```
post.counts <- predict_counts(fit, dt)</pre>
```

Compare observed and predicted visits by the model:

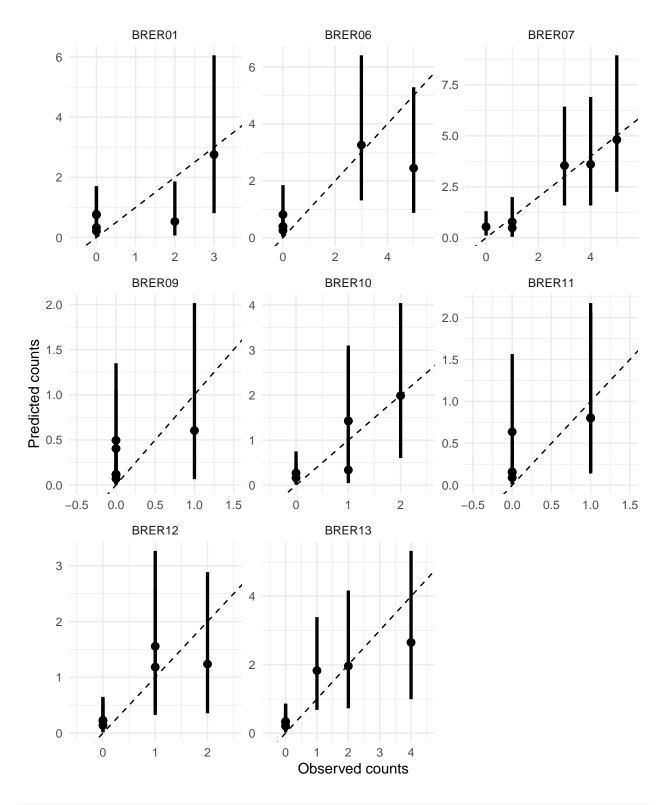
```
p <- plot_counts_pred(post.counts, sort = FALSE)
o <- plot_counts_obs(mat, sort = FALSE, zero.na = FALSE)
library(patchwork)
p + o</pre>
```



plot\_counts\_pred\_obs(post.counts, dt)



plot\_counts\_pred\_obs(post.counts, dt, byplant = TRUE, scales = "free")



saveRDS(post.counts, here::here(paste0("data/nets\_post/", params\$net, "\_post\_counts.rds")))