Supplementary Text S6: Assessing synteny detection in Fabaceae

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```
library(cogeqc)
library(here)
library(tidyverse)
library(syntenet)
library(tidytext)
```

1 Overview

Here, we will assess synteny detection using a network-based approach. The anchor pairs from synteny identification will be interpreted as edges of an unweighted undirected graph (i.e., a synteny network), and the best synteny detection will be identified based on the graphs' clustering coefficients and node number.

We will demonstrate our network-based synteny assessment using genomic data on Fabaceae species available on PLAZA 5.0 (Van Bel et al. 2022).

2 Data acquisition

In this section, we will download whole-genome protein sequences and gene annotation from PLAZA 5.0, and then we will preprocess the data with syntenet::process_input().

```
species <- c("mtr", "tpr", "psa", "car", "lja", "gma", "vmu", "lal", "arhy")</pre>
```

```
base_url <- "https://ftp.psb.ugent.be/pub/plaza/plaza_public_dicots_05/"</pre>
# Get proteomes
seq_url <- paste0(</pre>
    base_url, "Fasta/proteome.selected_transcript.",
    species, ".fasta.gz"
)
## Import files and clean gene IDs
seq <- lapply(seq_url, function(x) {</pre>
    s <- Biostrings::readAAStringSet(x)</pre>
    names(s) <- gsub(".* | ", "", names(s))
    return(s)
})
names(seq) <- species
# Get gene annotation
annot_url <- paste0(</pre>
    base_url, "GFF/", species, "/annotation.selected_transcript.exon_features.",
    species, ".gff3.gz"
)
## Import files and keep only relevant fields
annot <- lapply(annot_url, function(x) {</pre>
    a <- rtracklayer::import(x)</pre>
```

```
a <- a[, c("type", "gene_id")]
  a <- a[a$type == "gene"]
  return(a)
})
names(annot) <- species

# Process data
pdata <- process_input(seq, annot)

# Remove unprocessed data to clean the working environment
rm(annot)
rm(seq)</pre>
```

3 Network-based synteny assessment

We will infer synteny networks using the Bioconductor package *syntenet*. This package detects synteny using the MCScanX algorithm (Wang et al. 2012), which can produce different results based on 2 main parameters:

- 1. anchors: minimum required number of genes to call a syntenic block. Default: 5.
- 2. **max_gaps:** number of upstream and downstream genes to search for anchors. Default: 25.

We will infer synteny networks with 5 combinations of parameters, similarly to Zhao and Schranz (2019), using two approaches:

- 1. A single Fabaceae synteny network;
- 2. Species-specific synteny networks for each Fabaceae species.

To start with, let's define the combinations of parameters we will use.

3.1 Assessing the Fabaceae synteny network

First, we will perform similarity searches with DIAMOND.

```
# Define wrapper function to run DIAMOND with different top_hits
out <- file.path(tempdir(), "diamond_all")
d5 <- run_diamond(seq = seq, top_hits = 5, outdir = out)</pre>
```

With the DIAMOND list, we can detect synteny.

```
# Define helper function to detect synteny with multiple combinations of params
synteny_wrapper <- function(diamond, annotation, params) {</pre>
    syn <- lapply(params, function(x) {</pre>
        anchors <- x[1]
        max_gaps <- x[2]
        outdir <- file.path(tempdir(), paste0("syn_a", anchors, "_m", max_gaps))</pre>
        s <- infer_syntenet(</pre>
            blast_list = diamond,
            annotation = pdata$annotation,
            outdir = outdir,
            anchors = anchors,
            max_gaps = max_gaps
        )
        return(s)
    })
    return(syn)
}
# Detect synteny
syn_fabaceae <- synteny_wrapper(d5, pdata$annotation, synteny_params)</pre>
names(syn_fabaceae) <- unlist(</pre>
    lapply(synteny_params, function(x) paste0("a", x[1], "_m", x[2]))
```

Now, let's use the network-based synteny assessment to see which combination of parameters is the best.

```
# Assess networks
fabaceae_scores <- assess_synnet_list(syn_fabaceae)

# Look at scores, ranked from highest to lowest
fabaceae_scores %>%
    arrange(-Score) |>
    knitr::kable(
        caption = "Scores for each synteny network."
    )
```

Table 1: Scores for each synteny network.

CC	Node_count	Rsquared	Score	Network
0.8253002	237723	0.6227916	122187.2	a3_m25
0.8290880	235290	0.6156847	120105.4	a5_m35
0.8392223	226657	0.6026291	114629.5	a5_m15
0.8412602	224325	0.5972865	112717.3	a7_m25
0.8347725	231820	0.5795957	112161.6	a5_m25

As we can see, the combination of parameters a = 3; m = 25 is the best for this data set.

Finally, let's visualize scores. To make visualization better, we will scale scores by the maximum value, so that values range from 0 to 1.

```
# Plot scores
synteny_scores_fabaceae <- fabaceae_scores %>%
    arrange(Score) %>%
    mutate(Score = Score / max(Score)) %>%
    mutate(Parameters = str_replace_all(Network, "_", ", ")) %>%
    mutate(Parameters = factor(Parameters, levels = unique(Parameters))) %>%
    ggplot(., aes(x = Parameters, y = Score)) +
    geom_col(fill = "grey60", color = "black") +
    theme_bw() +
    labs(
        title = "Assessment of the Fabaceae synteny network",
        subtitle = "a = minimum # of anchors; m = maximum # of gaps",
        y = "Scaled score"
    )
synteny_scores_fabaceae
```

Assessment of the Fabaceae synteny network a = minimum # of anchors; m = maximum # of gaps

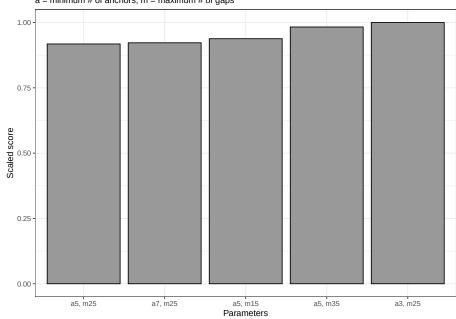


Figure 1: Scaled scores for each synteny network.

3.2 Assessing species-specific synteny networks

In this section, we will infer species-specific synteny networks and assess each of them with our network-based approach.

This time, as we already have synteny networks for the whole Fabaceae family, we don't need to infer them again; we will simply subset edges of the network that contain nodes from the same species.

```
# Create species-specific networks
species_ids <- substr(species, start = 1, stop = 3)</pre>
species_networks <- lapply(species_ids, function(x) {</pre>
    nets <- lapply(syn_fabaceae, function(y) {</pre>
        edges <- y[startsWith(y$Anchor1, x) \& startsWith(y$Anchor2, x), ]
        return(edges)
    })
    return(nets)
})
names(species_networks) <- species_ids</pre>
# Exploring data
names(species_networks)
## [1] "mtr" "tpr" "psa" "car" "lja" "gma" "vmu" "lal" "arh"
names(species_networks$mtr)
## [1] "a3_m25" "a5_m15" "a5_m25" "a5_m35" "a7_m25"
# Rename `species_networks` to keep full name
names(species_networks) <- c(</pre>
    "M. truncatula", "T. pratense", "P. sativum", "C. arietinum",
    "L. japonicus", "G. max", "V. mungo", "L. albus", "A. hypogaea"
)
```

For each species, we will assess the networks inferred with different combinations of parameters.

```
# Assess species-specific networks
scores_species_nets <- lapply(seq_along(species_networks), function(x) {</pre>
    species <- names(species_networks)[x]</pre>
    scores <- assess_synnet_list(species_networks[[species]])</pre>
    scores$Score[is.nan(scores$Score)] <- 0</pre>
    scores <- scores[order(scores$Score, decreasing = TRUE), ]</pre>
    scores$Species <- species</pre>
    scores$Score <- scores$Score / max(scores$Score)</pre>
    return(scores)
})
scores_species_nets <- Reduce(rbind, scores_species_nets)</pre>
# Plot data
synteny_scores_species <- scores_species_nets %>%
    mutate(
        Parameters = as.factor(str_replace_all(Network, "_", ", ")),
        Species = as.factor(Species)
    mutate(Network = reorder_within(Parameters, Score, Species)) %>%
    ggplot(., aes(x = Network, y = Score, fill = Parameters)) +
    geom_bar(stat = "identity", color = "grey90") +
    facet_wrap(~Species, ncol = 3, scales = "free") +
```

```
scale_x_reordered() +
ggsci::scale_fill_jama() +
theme_bw() +
theme(axis.text.x = element_blank()) +
labs(
    title = "Assessment of species-specific synteny networks",
    subtitle = "a = minimum # of anchors; m = maximum # of gaps",
    y = "Scaled score (by species)", x = ""
)
synteny_scores_species
```

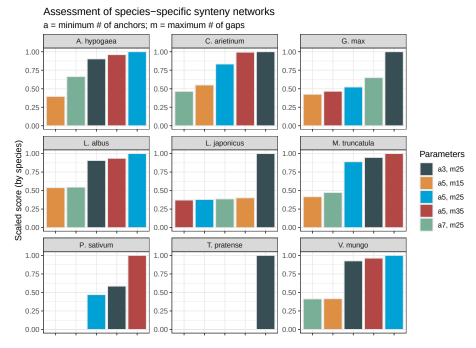


Figure 2: Scores for species-specific synteny networks.

The figure demonstrates that the best combination of parameters depends on the species, so there is no "universally" best combination. However, some patterns emerge. The combinations a=7; m=25 and a=5; m=15 are typically the worst. In some cases, they even lead to zero scores due to clustering coefficients of zero. Thus, if users want to test multiple combinations of parameters for their own data set, they should only test the combinations a=3; m=25, a=5; m=25, and a=5; m=35, which lead to the best score in 45%, 33%, and 22% of the species-specific networks, respectively. Interestingly, the combination that leads to the best score in most networks (a=3; a=25) is also the best when considering the whole Fabaceae synteny network (see previous section).

Session info

This document was created under the following conditions:

```
## setting value
## version R version 4.3.0 (2023-04-21)
## os Ubuntu 20.04.5 LTS
## system x86_64, linux-qnu
## ui X11
## language (EN)
## collate en_US.UTF-8
## ctype en_US.UTF-8
              Europe/Brussels
## tz
## date 2023-10-06
## pandoc 3.1.1 @ /usr/lib/rstudio/resources/app/bin/quarto/bin/tools/ (via rmarkdown)
## - Packages -------
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2.68.0 2023-04-25 [1] Bioconductor

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* 1.4.0 2023-04-25 [1] Bioconductor

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## codetools
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## colorspace
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## DelayedArray 0.26.1 2023-05-01 [1] Bioconductor digest 0.6.33 2023-07-07 [1] CRAN (R 4.3.0) ## dplyr * 1.1.2 2023-04-20 [1] CRAN (R 4.3.0) ## evaluate 0.21 2023-05-05 [1] CRAN (R 4.3.0) ## fansi 1.0.4 2023-01-22 [1] CRAN (R 4.3.0)
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   [2] /usr/local/lib/R/site-library
## [3] /usr/lib/R/site-library
## [4] /usr/lib/R/library
##
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```

References

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