canaper: Categorical analysis of neo- and paleo-endemism in R

Joel H. Nitta^{1, iD}

¹ University of Tokyo, Department of Biological Sciences

- joelnitta@gmail.com
- https://joelnitta.com

https://github.com/joelnitta/canaper



1. Background

- CANAPE (Categorical Analysis of Neo- and Paleo-endemism) is a recently developed method that provides insight into the evolutionary processes underlying endemism [1]
- CANAPE integrates a phylogenetic tree with a community (species × sites) matrix to infer if endemic areas are so because of recent speciation (neoendemism), or because they harbor old lineages that have mostly gone extinct in other areas (paleoendemism), or if they are a mixture of both
- CANAPE is currently only implemented in Biodiverse [2], a program written in perl that is used via a GUI or custom scripts.
- canaper is a new R package to conduct CANAPE entirely in R

2. Features

- Sparse matrix encoding of community matrices to increase computing efficiency via phyloregion [3]
- Simple implementation of parallel computing to increase speed via future

3. Installation and loading

canaper is currently available on GitHub

```
devtools::install github("joelnitta/canaper")
# Load packages
library(canaper)
library(ape) # For handling phylogenies
library(future) # For parallel computing
library(tidyverse) # For data wrangling and visualization
```

4. Example: Australian Acacia

This demonstrates the package using the dataset of the paper where CANAPE was first published [1]: 506 species of Acacia in Australia distributed over 3037 sites:

```
# Phylogenetic tree
acacia$phy
#> Phylogenetic tree with 510 tips and 509 internal nodes.
#> Tip labels:
#> Pararchidendron_pruinosum, Paraserianthes_lophantha, adinophylla,
        semicircinalis, aphanoclada, inaequilatera, ...
#> Rooted; includes branch lengths.
# Community data matrix (in part).
# Rownames correspond to geographical coordinates
acacia$comm[1:4, 1:4]
                       abbreviata acanthaster acanthoclada acinacea
#> '-1025000:-1825000'
#> '-1025000:-1875000'
#> '-1025000:-1925000'
#> '-1025000:-1975000'
```

4.1. Run randomization test

The first step of CANAPE is to compare the observed values of phylogenetic endemism (PE) and alternative PE (PE measured on a modified tree where all branch lengths are set equal) with those from a set of random communities. The cpr rand test() conducts the randomization, using parallel computing to increase speed. The picante package [4] is used to generate the random communities.

```
# Set a parallel back-end, with 4 CPUs running simultaneously
plan(multisession, workers = 4)
# Run randomization test
acacia rand res <- cpr rand test(
 acacia$comm, acacia$phy,
 n reps = 100, n iterations = 100000)
#> [1] "Dropping tips from the tree because they are not present in
        the community data:"
#> [1] "Pararchidendron_pruinosum" "Paraserianthes_lophantha"
#> [3] "saligna"
                                  "clunies-rossiae"
# Check some of the results
acacia rand res %>%
 slice(1:3) %>%
 select(pe_obs, pe_rand_mean, pe_obs_p_upper, pe_obs_p_lower) %>%
 as tibble()
#> # A tibble: 3 x 4
        pe_obs pe_rand_mean pe_obs_p_upper pe_obs_p_lower
                      <db1>
#> 1 0.0000248
                   0.000130
                   0.000297
#> 2 0.000145
                  0.000232
#> 3 0.000172
                                                    0.42
```

Output summary (in part):

- pe obs = observed PE
- pe rand mean = mean PE of the randomizations
- pe obs p upper = percent of randomizations where observed PE was greater than random values
- pe obs p lower = percent of randomizations where observed PE was lower than random values

4.2. Classify significance

The next step of CANAPE is to classify the endemism type of each site. The cpr_classify_endem() function does this automatically given output of cpr rand test().

```
# Classify endemism type
acacia canape <- cpr classify endem(acacia rand res)</pre>
# Count the results
count(acacia_canape, endem_type)
         endem_type
#> 1
               mixed 101
#> 2
#> 3 not significant 2760
              paleo 43
              super 122
#> 5
```

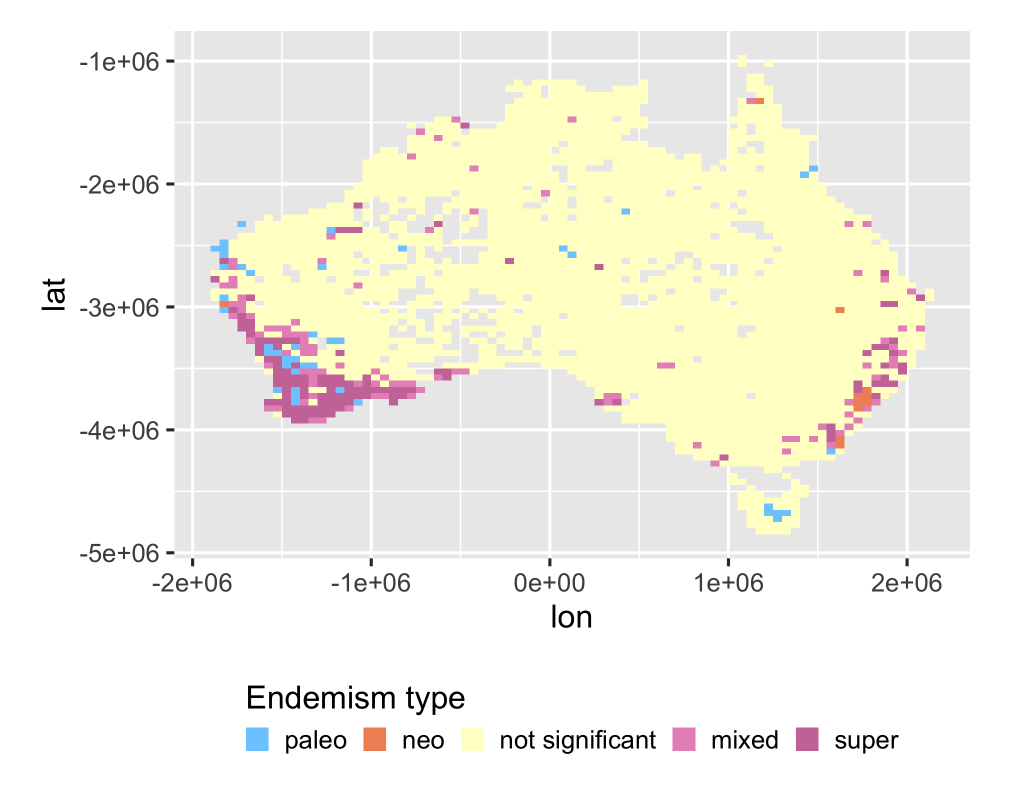
Endemism codes:

- neo = Neo-endemic
- paleo = Paleo-endemic
- mixed = Mix of neo and paleo
- super = Mixed, and highly significant (p < 0.01)

4.3. Visualize results

We can visualize the results of CANAPE using ggplot2. The cpr_endem cols palette that is accessible regardless of color vision deficiency is provided.

```
# Fist do some data wrangling to make
# the results easier to plot (add lat/long columns)
acacia_canape <- as_tibble(acacia_canape, rownames = "site") %>%
 separate(site, c("lon", "lat"), sep = ":") %>%
 mutate(dplyr::across(c(lon, lat), parse number))
# Plot the results
ggplot(acacia\_canape, aes(x = lon, y = lat, fill = endem type)) +
 geom tile() +
  scale fill manual(values = cpr endem cols, name = "Endemism type")
 guides(fill = guide legend(title.position = "top")) +
 theme_gray(base_size = 24) +
 theme(legend.position = "bottom")
```



5. Next steps

- Implement rand structured randomization algorithm of Biodiverse (should speed up randomizations ~10×)
- Submit to CRAN

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References

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Source code: https://github.com/joelnitta/botany_poster_2021